
saemix

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Chapter 1

Introduction

`saemix` is a package for the R software [33] to perform parameter estimation in non-linear mixed effect models. It has been hosted on the CRAN since version 0.95 in June 2011.

1.1 The objectives

The objectives of `saemix` are to perform:

1. parameter estimation for non-linear mixed effects models
 - computing the maximum likelihood estimator of the population parameters, without any approximation of the model (linearization, quadrature approximation, ...), using the Stochastic Approximation Expectation Maximization (SAEM) algorithm,
 - computing standard errors for the maximum likelihood estimator
 - computing the conditional modes, the conditional means and the conditional standard deviations of the individual parameters, using the Hastings-Metropolis algorithm
2. goodness of fit plots
3. model selection
 - comparing several models using some information criteria (AIC, BIC)
 - testing hypotheses using the Likelihood Ratio Test
 - testing parameters using the Wald Test

The R package `saemix` is an implementation of the Stochastic Approximation Expectation Maximization (SAEM) algorithm in R [33], developed by Kühn and Lavielle [19], and implemented in the MONOLIX software available in Matlab and as a standalone software for Windows and Linux [20].

The current version of the R version of `saemix` handles only analytical functions. The following features have not yet been implemented in the R package `saemix`, but are available in the MONOLIX software:

- categorical covariates with more than 2 categories
- models defined with differential equations
- multi-response model
- left censored data
- interoccasion variability
- prior distribution for the random effects
- complex variables, including discrete data or repeated time to events
- hidden Markov models
- mixture models
- autocorrelation of the residuals

Theoretical analysis of the algorithms used in this software can be found in [11, 13, 18, 19]. Several application of SAEM in agronomy [28], animal breeding [17] and PKPD analysis [5, 23, 37, 39, 2] have been published by several members of the Monolix group. Several applications to PKPD analysis were also proposed during the last PAGE (Population Approach Group in Europe) meetings ([29, 22, 21, 36, 38, 40] as well as a comparison of estimation algorithms [15], (<http://www.page-meeting.org>).

The present document describes the non-linear mixed effects models (section 1.3) and the algorithms used in this package (section 2). The library's inputs and outputs are described in section 3. Section 4 shows some examples made available in the library (section).

1.2 Installation and legalese

1.2.1 Installation

1.2.2 Citing saemix

If you use this program in a scientific publication, we would like you to cite the following reference:

Comets E, Lavenu A, Lavielle M (2017). SAEMIX, an R version of the SAEM algorithm. *Journal of Statistical Software*, 80:1-41.

A BibTeX entry for \LaTeX users is:

```
@Article{,
author = {Emmanuelle Comets and Audrey Lavenu and Marc Lavielle},
title = {Parameter estimation in nonlinear mixed effect models using saemix, an {R} implemen
volume = {80},
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1.3 The non-linear mixed effects model

1.3.1 Model for the observations

Longitudinal outcome

Detailed and complete presentations of the non-linear mixed effects model can be found in [6, 7, 32]. See also the many references therein.

We consider the following general non-linear mixed effects model for continuous outputs:

$$y_{ij} = f(x_{ij}, \psi_i) + g(x_{ij}, \psi_i, \xi) \varepsilon_{ij} \quad , \quad 1 \leq i \leq N \quad , \quad 1 \leq j \leq n_i \quad (1.1)$$

Here,

- $y_{ij} \in \mathbb{R}$ is the j th observation of subject i ,
- N is the number of subjects,
- n_i is the number of observations of subject i ,
- the regression variables, or design variables, (x_{ij}) are assumed to be known, $x_{ij} \in \mathbb{R}^{n_x}$,
- for subject i , the vector $\psi_i = (\psi_{i,\ell}; 1 \leq \ell \leq n_\psi) \in \mathbb{R}^{n_\psi}$ is a vector of n_ψ individual parameters:

$$\psi_i = H(\mu, c_i, \eta_i) \quad (1.2)$$

where

- $c_i = (c_{im}; 1 \leq m \leq M)$ is a known vector of M covariates,
- μ is an unknown vector of fixed effects of size n_μ ,
- η_i is an unknown vector of normally distributed random effects of size n_η :

$$\eta_i \sim_{i.i.d.} \mathcal{N}(0, \Omega)$$

- the residual errors (ε_{ij}) are random variables with mean zero and variance 1,
- the residual error model is defined by the function g and some parameters ξ .

The residual error model

The within-group errors (ε_{ij}) are supposed to be Gaussian random variables with mean zero and variance 1. Furthermore, we suppose that the ε_{ij} and the η_i are mutually independent.

Different error models can be used in saemix 3.0:

- the constant error model assumes that $g = a$ and $\xi = a$,
- the proportional error model assumes that $g = b f$ and $\xi = b$,
- a combined error model assumes that $g = a + b f$ and $\xi = (a, b)$,

Furthermore, all these error models can be applied to some transformation of the data:

$$t(y_{ij}) = t(f(x_{ij}, \psi_i)) + g(x_{ij}, \psi_i, \xi)\varepsilon_{ij} \quad (1.3)$$

In the current version of **saemix**, the exponential error model is also available: it assumes that $y > 0$ and that:

$$\begin{aligned} t(y) &= \log(y) \\ y &= f e^{g\varepsilon} \end{aligned}$$

Discrete outcome

Categorical responses can take a finite number of possibly values, and we define the distribution \mathcal{D}_y as the set of probabilities for each value, summing up to 1, so that the j^{th} observation y_{ij} in subject i follows $\mathcal{D}_y(x_{ij}; \psi_i, \xi)$.

Binary data: An example is binary response, where we model the probability of the response being 1, for exemple through the logistic model:

$$p(y_{ij}|\psi_i) = \frac{e^{f(x_{ij}, \psi_i)}}{1 + e^{f(x_{ij}, \psi_i)}}$$

where $f(x_{ij}, \psi_i)$ is a function of the individual parameters and the design variables. f is most often a linear function, and the model is equivalently written through the logit of p , where $\text{logit}(p) = \ln \frac{p}{1-p}$, for example:

$$\text{logit}(p(y_{ij}|\psi_i)) = \alpha_i + \beta_i x_{ij}$$

Categorical data: Assume now that the observed data takes its values in a fixed and finite set of nominal categories $\{c_1, c_2, \dots, c_K\}$. Considering the observations $(y_{ij}, 1 \leq j \leq n_i)$ for any individual i as a sequence of conditionally independent random variables, the model is completely defined by the probability mass functions $\mathbb{P}(y_{ij} = c_k|\psi_i)$ for $k = 1, \dots, K$ and $1 \leq j \leq n_i$. For a given (i, j) , the sum of the K probabilities is 1, so in fact only $K - 1$ of them need to be defined. In the most general way possible, any model can be considered so long as it defines a probability

distribution, i.e., for each k , $\mathbb{P}(y_{ij} = c_k | \psi_i) \in [0, 1]$, and $\sum_{k=1}^K \mathbb{P}(y_{ij} = c_k | \psi_i) = 1$. Ordinal data further assume that the categories are ordered, i.e., there exists an order \prec such that

$$c_1 \prec c_2, \prec \dots \prec c_K$$

We can think, for instance, of levels of pain (*low* \prec *moderate* \prec *severe*) or scores on a discrete scale, e.g., from 1 to 10. Instead of defining the probabilities of each category, it may be convenient to define the cumulative probabilities $\mathbb{P}(y_{ij} \preceq c_k | \psi_i)$ for $k = 1, \dots, K - 1$, or in the other direction: $\mathbb{P}(y_{ij} \succeq c_k | \psi_i)$ for $k = 2, \dots, K$. Any model is possible as long as it defines a probability distribution, i.e., it satisfies

$$0 \leq \mathbb{P}(y_{ij} \prec c_1 | \psi_i) \leq \mathbb{P}(y_{ij} \prec c_2 | \psi_i) \leq \dots \leq \mathbb{P}(y_{ij} \prec c_K | \psi_i) = 1$$

Note: It is possible to introduce dependence between observations from the same individual by assuming that $(y_{ij}, j = 1, 2, \dots, n_i)$ forms a Markov chain. For instance, a Markov chain with memory 1 assumes that all that is required from the past to determine the distribution of y_{ij} is the value of the previous observation $y_{i,j-1}$, i.e., for all $k = 1, 2, \dots, K$,

$$\mathbb{P}(y_{ij} = c_k | y_{i,j-1}, y_{i,j-2}, y_{i,j-3}, \dots, \psi_i) = \mathbb{P}(y_{ij} = c_k | y_{i,j-1}, \psi_i)$$

Such a model is currently not within the scope of `saemix`.

Count data: Count data can take a number of possible values, possibly infinite, and again we model the probability of each count. A common model is the Poisson model with parameter λ , where for subject i at time x_{ij} the probability of observing a count equal to k is given by:

$$p(y_{ij} = k | \lambda_i) = \frac{e^{-\lambda_i(x_{ij})}}{k!} \lambda_i(x_{ij})^k$$

Time-to-event outcome

Although technically a continuous outcome, time-to-event data are modelled like discrete outcomes by considering their likelihood. In a time-to-event data model, the observations are the times at which events occur. An event may be one-off (e.g., death, hardware failure) or repeated (e.g., epileptic seizures, mechanical incidents). To begin with, we consider a model for

a one-off event. The survival function $S(t)$ gives the probability that the event happens after time t :

$$S(t) \triangleq \mathbb{P}(T > t) = \exp \left\{ - \int_0^t h(u) du \right\} , \quad (1.4)$$

where h is called the hazard function. In a population approach, we consider a parametric and individual hazard function $h(\cdot, \psi_i)$.

The random variable representing the time-to-event for individual i is typically written T_i and may possibly be right-censored. Then, the observation y_i for individual i is

$$y_i = \begin{cases} T_i & \text{if } T_i \leq \tau_c \\ "T_i > \tau_c" & \text{otherwise ,} \end{cases} \quad (1.5)$$

where τ_c is the censoring time and " $T_i > \tau_c$ " is the information that the event occurred after the censoring time.

For repeated event models, times when events occur for individual i are random times $(T_{ij}, 1 \leq j \leq n_i)$ for which conditional survival functions can be defined:

$$\mathbb{P}(T_{ij} > t | T_{i(j-1)} = t_{i(j-1)}) = \exp \left\{ - \int_{t_{i(j-1)}}^t h(u, \psi_i) du \right\} . \quad (1.6)$$

Here, t_{ij} is the observed value of the random time T_{ij} . If the last event is right censored, then the last observation y_{i,n_i} for individual i is the information that the censoring time has been reached " $T_{i,n_i} > \tau_c$ ". The conditional pdf of $y_i = (y_{ij}, 1 \leq n_i)$ reads (see [20] for more details)

$$p(y_i | \psi_i) = \exp \left\{ - \int_0^{\tau_c} h(u, \psi_i) du \right\} \prod_{j=1}^{n_i-1} h(t_{ij}, \psi_i) . \quad (1.7)$$

1.3.2 The statistical model for the individual parameters

We assume that ψ_i is a transformation of a Gaussian random vector ϕ_i :

$$\psi_i = h(\phi_i) \quad (1.8)$$

where, by rearranging the covariates (c_{im}) into a matrix C_i , ϕ_i can be written as:

$$\phi_i = C_i \mu + \eta_i \quad (1.9)$$

Examples of transformations

Here, different transformations (h_ℓ) can be used for the different components of $\psi_i = (\psi_{i,\ell})$ where $\psi_{i,\ell} = h_\ell(\phi_{i,\ell})$ for $\ell = 1, 2, \dots, \ell$.

- $\psi_{i,\ell}$ has a normal distribution if $h_\ell(u) = u$
- $\psi_{i,\ell}$ has a log-normal distribution if $h_\ell(u) = e^u$
- assuming that $\psi_{i,\ell}$ takes its values in $(0, 1)$, we can use a logit transformation $h_\ell(u) = 1/(1 + e^{-u})$, or a probit transformation $h_\ell(u) = \mathbb{P}(\mathcal{N}(0, 1) \leq u)$.

In the following, we will use either the parameters ψ_i or the Gaussian transformed parameters $\phi_i = h^{-1}(\psi_i)$.

The model can address continuous and/or binary covariates.

Example of continuous covariate model

Consider a PK model that depends on volume and clearance and consider the following covariate model for these two parameters:

$$\begin{aligned} CL_i &= CL_{\text{pop}} \left(\frac{W_i}{W_{\text{pop}}} \right)^{\beta_{CL,W}} \left(\frac{A_i}{A_{\text{pop}}} \right)^{\beta_{CL,A}} e^{\eta_{i,1}} \\ V_i &= V_{\text{pop}} \left(\frac{W_i}{W_{\text{pop}}} \right)^{\beta_{V,W}} e^{\eta_{i,2}} \end{aligned}$$

Where W_i and A_i are the weight and the age of subject i and where W_{pop} and A_{pop} are some “typical” values of these two covariates in the population. Here, ψ_i will denote the PK parameters (clearance and volume) of subject i and ϕ_i its log-clearance and log-volume. Let

$$W_i^* = \log \left(\frac{W_i}{W_{\text{pop}}} \right) \quad ; \quad A_i^* = \log \left(\frac{A_i}{A_{\text{pop}}} \right)$$

Then,

$$\begin{aligned} \phi_i &= \begin{pmatrix} \log(CL_i) \\ \log(V_i) \end{pmatrix} \\ &= \begin{pmatrix} 1 & 0 & W_i^* & W_i^* & 0 \\ 0 & 1 & 0 & 0 & W_i^* \end{pmatrix} \begin{pmatrix} \log(CL_{\text{pop}}) \\ \log(V_{\text{pop}}) \\ \beta_{CL,W} \\ \beta_{CL,A} \\ \beta_{V,W} \end{pmatrix} + \begin{pmatrix} \eta_{i,1} \\ \eta_{i,2} \end{pmatrix} \\ &= C_i \mu + \eta_i \end{aligned}$$

Example of categorical covariate model

Assume that some categorical covariate G_i takes the values $1, 2, \dots, K$. Assume that if patient i belongs to group k , *i.e.* $G_i = k$, then

$$\log(CL_i) = \log(CL_{\text{pop},k}) + \eta_i$$

where $CL_{\text{pop},k}$ is the population clearance in group k .

Let k^* be the reference group. Then, for any group k , we will decompose the population clearance $CL_{\text{pop},k}$ as

$$\log(CL_{\text{pop},k}) = \log(CL_{\text{pop},k^*}) + \beta_k$$

where $\beta_{k^*} = 0$.

The variance of the random effects can also depend on this categorical covariate:

$$\eta_i \sim \mathcal{N}(0, \Omega_k) \quad \text{if } G_i = k$$

Remark: It is assumed in *saemix* 3.0 that the categorical covariate has only 2 categories (binary covariate). It is also assumed that the variance remains the same for both groups. Covariates with more than 2 categories can always be recoded into (N-1) binary covariates using dummy variables but this is for the moment up to the user.

1.3.3 General form of the non-linear mixed effect model (NLMEM)

A general form of the non-linear mixed effect model regrouping the different types of responses is to define the model fully in terms of the probabilities:

$$\begin{cases} y_{ij} \sim \mathcal{D}_y(x_{ij}; \psi_i, \xi) \\ \psi_i \sim \mathcal{D}_\psi(c_i; \mu, \Omega) \end{cases}$$

The parameters of the model are $\theta = (\mu, \Omega, \xi)$, with ξ denoting the additional parameters of the residual error model for continuous response models.

We will denote $\ell(y; \theta)$ the likelihood of the observations $y = (y_{ij}; 1 \leq i \leq n, 1 \leq j \leq n_i)$ and $p(y, \psi; \theta)$ the likelihood of the complete data $(y, \psi) = (y_{ij}, \psi_i; 1 \leq i \leq n, 1 \leq j \leq n_i)$. Thus,

$$\ell(y; \theta) = \int p(y, \psi; \theta) d\psi.$$

Chapter 2

Methodology and algorithms

2.1 Estimation of the parameters

2.1.1 The SAEM algorithm

We are in a classical framework of incomplete data: the observed data is $y = (y_{ij}; 1 \leq i \leq N, 1 \leq j \leq n_i)$, whereas the random parameters $(\psi = \psi_i; 1 \leq i \leq N)$ are the non observed data. Then, the complete data of the model is (y, ψ) . Our purpose is to compute the maximum likelihood estimator of the unknown set of parameters $\theta = (\mu, \Omega, a, b, c)$, by maximizing the likelihood of the observations $\ell(y; \theta)$.

In the case of a linear model, the estimation of the unknown parameters can be treated with the usual EM algorithm. At iteration k of EM, the E-step consists in computing the conditional expectation of the complete log-likelihood $Q_k(\theta) = \mathbb{E}(\log p(y, \psi; \theta) | y, \theta_{k-1})$ and the M-step consists in computing the value θ_k that maximises $Q_k(\theta)$.

Following [12, 47], the EM sequence (θ_k) converges to a stationary point of the observed likelihood (*i.e* a point where the derivative of ℓ is 0) under general regularity conditions. In cases where the regression function f does not linearly depend on the random effects, the E-step cannot be performed in a closed-form.

The stochastic approximation version of the standard EM algorithm, proposed by [11] consists in replacing the usual E-step of EM by a stochastic procedure. At iteration k of SAEM:

- *Simulation-step* : draw $\psi^{(k)}$ from the conditional distribution $p(\cdot | y; \theta_k)$.

- *Stochastic approximation* : update $Q_k(\theta)$ according to

$$Q_k(\theta) = Q_{k-1}(\theta) + \gamma_k(\log p(y, \psi^{(k)}; \theta) - Q_{k-1}(\theta)) \quad (2.1)$$

where (γ_k) is a decreasing sequence of positive numbers with $\gamma_1 = 1$.

- *Maximization-step* : update θ_k according to

$$\theta_{k+1} = \text{Arg max}_{\theta} Q_k(\theta).$$

It is shown in [11] that SAEM converges to a maximum (local or global) of the likelihood of the observations under very general conditions.

Here, the complete log-likelihood can be written

$$\begin{aligned} \log p(y, \psi; \theta) &= \log p(y, h(\phi); \theta) \\ &= - \sum_{i,j} \log(g(x_{ij}, \psi_i, \xi)) - \frac{1}{2} \sum_{i,j} \left(\frac{y_{ij} - f(x_{ij}, \psi_i)}{g(x_{ij}, \psi_i, \xi)} \right)^2 \\ &\quad - \frac{N}{2} \log(|\Omega|) - \frac{1}{2} \sum_{i=1}^N (\phi_i - C_i \mu)' \Omega^{-1} (\phi_i - C_i \mu) - \frac{N_{tot} + Nd}{2} \log(2\pi) \end{aligned}$$

where $N_{tot} = \sum_{i=1}^N n_i$ is the total number of observations.

First, consider a constant residual error model ($g = a$). The set of parameters to estimate is $\theta = (\mu, \Omega, a)$. Then, the complete model belongs to the exponential family and the approximation step reduces to only updating the sufficient statistics of the complete model:

$$\begin{aligned} s_{1,i,k} &= s_{1,i,k-1} + \gamma_k (\phi_{i,k} - s_{1,i,k-1}), \quad i = 1, \dots, N \\ s_{2,k} &= s_{2,k-1} + \gamma_k \left(\sum_{i=1}^N \phi_{i,k} \phi'_{i,k} - s_{2,k-1} \right) \\ s_{3,k} &= s_{3,k-1} + \gamma_k \left(\sum_{i,j} \left(y_{ij} - f(x_{ij}, \psi_i^{(k)}) \right)^2 - s_{3,k-1} \right). \end{aligned}$$

Then, θ_{k+1} is obtained in the maximization step as follows:

$$\mu_{k+1} = \left(\sum_{i=1}^N C_i' \Omega_k^{-1} C_i \right)^{-1} \sum_{i=1}^N C_i' \Omega_k^{-1} s_{1,i,k} \quad (2.2)$$

$$\Omega_{k+1} = \frac{1}{N} \left(s_{2,k} - \sum_{i=1}^N (C_i \mu_{k+1}) s'_{1,i,k} - \sum_{i=1}^N s_{1,i,k} (C_i \mu_{k+1})' + \sum_{i=1}^N (C_i \mu_{k+1}) (C_i \mu_{k+1})' \right) \quad (2.3)$$

$$a_{k+1} = \sqrt{\frac{s_{3,k}}{N_{tot}}} \quad (2.4)$$

Remark 1: The sequence of step sizes used in `saemix` decreases as k^{-a} . More precisely, for any sequence of integers K_1, K_2, \dots, K_J and any sequence a_1, a_2, \dots, a_J of real numbers such that $0 \leq a_1 < a_2 < \dots < a_J \leq 1$, we define the sequence of step sizes (γ_k) as follows:

$$\gamma_k = \frac{1}{k^{a_1}} \quad \text{for any } 1 \leq k \leq K_1 \quad (2.5)$$

and for $2 \leq j \leq J$,

$$\gamma_k = \frac{1}{\left(k - K_{j-1} + \gamma_{K_{j-1}}^{-1/a_j}\right)^{a_j}} \quad \text{for any } \sum_{i=1}^{j-1} K_i + 1 \leq k \leq \sum_{i=1}^j K_i \quad (2.6)$$

Here, $K = \sum_{j=1}^J K_j$ is the total number of iterations.

We recommend to use $a_1 = 0$ (that is $\gamma_k = 1$) during the first iterations, and $a_J = 1$ during the last iterations. Indeed, the initial guess θ_0 may be far from the maximum likelihood value we are looking for and the first iterations with $\gamma_k = 1$ allow to converge quickly to a neighborhood of the maximum likelihood estimator. Then, smaller step sizes ensure the almost sure convergence of the algorithm to the maximum likelihood estimator.

In the case where $J = 2$ with $a_1 = 0$ and $a_2 = 1$, the sequence of step sizes is

$$\begin{aligned} \gamma_k &= 1 && \text{for } 1 \leq k \leq K_1 \\ &= \frac{1}{k - K_1 + 1} && \text{for } K_1 + 1 \leq k \leq K_1 + K_2 \end{aligned}$$

Remark 2: The estimated covariance matrix Ω_{k+1} defined in (2.3) is a full covariance matrix. However, the covariance matrix Ω of the random effects can have any covariance structure. If we assume, for example, that there is no correlation between the random effects, we will set to 0 the non diagonal elements of Ω_{k+1} defined in (2.3).

We can also assume that a random effect has no variance. If the ℓ th random effect has a variance equal to 0, then the ℓ th individual parameter is no longer random and the simulation step of SAEM needs some modification. During the first K_0 iterations, we use SAEM as it was described above, considering that all the effects are random and assuming that there is no correlation between the ℓ th random effect and the other ones ($\omega_{\ell\ell'}^2 = 0$ for any $\ell \neq \ell'$). Then, during the next iterations, we use again SAEM, but the variance of this random effect is no longer estimated: it is forced to decrease at each iteration by setting

$$\omega_{\ell\ell,k+1}^2 = \alpha \omega_{\ell\ell,k}^2, \quad K_0 \leq k \leq K \quad (2.7)$$

where α is chosen between 0 and 1 such that $\omega_{\ell\ell,K}^2 = 10^{-6} \omega_{\ell\ell,K_0}^2$.

Remark 3: - For a residual variance model of the form $g = b f^c$, where c is fixed, the complete model also belongs to the exponential family and the estimation of b is straightforward: the

sufficient statistics sequence $(s_{3,k})$ is defined by

$$s_{3,k} = s_{3,k-1} + \gamma_k \left(\sum_{i,j} \left(\frac{y_{ij} - f(x_{ij}, \psi_i^{(k)})}{f^c(x_{ij}, \psi_i^{(k)})} \right)^2 - s_{3,k-1} \right)$$

and $b_{k+1} = \sqrt{s_{3,k}/N_{tot}}$.

- For a general residual variance model $g = a + b f^c$, the complete model does not belong to the exponential family and the estimates of the residual variance parameters (a, b, c) cannot be expressed as a function of some sufficient statistics. Then, let (A_k, B_k, C_k) that minimise the complete log-likelihood:

$$(A_k, B_k, C_k) = \text{Arg min}_{(a,b,c)} \left\{ \sum_{i,j} \log(a + b f^c(x_{ij}, \psi_i^{(k)})) + \frac{1}{2} \sum_{i,j} \left(\frac{y_{ij} - f(x_{ij}, \psi_i^{(k)})}{a + b f^c(x_{ij}, \psi_i^{(k)})} \right)^2 \right\}$$

We update the residual variance parameters as follows:

$$a_{k+1} = a_k + \gamma_k (A_k - a_k) \quad (2.8)$$

$$b_{k+1} = b_k + \gamma_k (B_k - b_k) \quad (2.9)$$

$$c_{k+1} = c_k + \gamma_k (C_k - c_k) \quad (2.10)$$

The estimation of μ and Ω remains unchanged.

2.1.2 The MCMC-SAEM algorithm

For model (1.1), the simulation step cannot be directly performed. Kuhn and Lavielle [18] propose to combine the SAEM algorithm with a MCMC (Markov Chain Monte Carlo) procedure. This procedure consists in replacing the Simulation-step at iteration k by m iterations of the Hastings-Metropolis algorithm.

Here, we will consider the Gaussian parameters (ϕ_i) . For $i = 1, 2, \dots, N$

- let $\phi_{i,0} = \phi_i^{(k-1)}$
- for $p = 1, 2, \dots, m$,
 1. draw $\tilde{\phi}_{i,p}$ using the proposal kernel $q_{\theta_k}(\phi_{i,p-1}, \cdot)$
 2. set $\phi_{i,p} = \tilde{\phi}_{i,p}$ with probability

$$\alpha(\phi_{i,p-1}, \tilde{\phi}_{i,p}) = \min \left(1, \frac{p(\tilde{\phi}_{i,p}|y_i; \theta_k) q_{\theta_k}(\phi_{i,p-1}, \tilde{\phi}_{i,p})}{p(\phi_{i,p-1}|y_i; \theta_k) q_{\theta_k}(\phi_{i,p-1}, \tilde{\phi}_{i,p})} \right)$$

and $\phi_{i,p} = \phi_{i,p-1}$ with probability $1 - \alpha(\phi_{i,p-1}, \tilde{\phi}_{i,p})$.

- let $\phi_i^{(k)} = \phi_{i,m}$.

Several transition kernels, associated to different proposals can be successively used. We use the four following proposal kernels:

1. $q_{\theta_k}^{(1)}$ is the prior distribution of ϕ_i at iteration k , that is the Gaussian distribution $\mathcal{N}(C_i \mu_k, \Omega_k)$ and then

$$\alpha(\phi_{i,p-1}, \tilde{\phi}_{i,p}) = \min \left(1, \frac{p(y_i | \tilde{\phi}_{i,p}; \theta_k)}{p(y_i | \phi_{i,p-1}; \theta_k)} \right)$$

2. $q_{\theta_k}^{(2)}$ is a random permutation of the ϕ_i : generate a random permutation σ of $\{1, 2, \dots, N\}$ and set $\tilde{\phi}_{i,p} = \phi_{\sigma(i),p-1}$.

3. $q_{\theta_k}^{(3)}$ is the multidimensional random walk $\mathcal{N}(\phi_{i,p-1}, \kappa \Omega_k)$. This kernel is symmetric and then

$$\alpha(\phi_{i,p-1}, \tilde{\phi}_{i,p}) = \min \left(1, \frac{p(y_i, \tilde{\phi}_{i,p}; \theta_k)}{p(y_i, \phi_{i,p-1}; \theta_k)} \right)$$

4. $q_{\theta_k}^{(4)}$ is a succession of d unidimensional Gaussian random walks: each component of ϕ_i are successively updated.

Then, the simulation-step at iteration k consists in running m_1 iterations of the Hasting-Metropolis with proposal $q_{\theta_k}^{(1)}$, m_2 iterations with proposal $q_{\theta_k}^{(2)}$, m_3 iterations with proposal $q_{\theta_k}^{(3)}$ and m_4 iterations with proposal $q_{\theta_k}^{(4)}$.

Remark 1 : During the first K_b iterations ("burning" iterations) of SAEM, we only run the MCMC algorithm but the parameters are not updated.

Remark 2 : When the number N of subjects is small, convergence of the algorithm can be improved by running L Markov Chain instead of only one. The simulation step requires to draw L sequences $\phi^{(k,1)}, \dots, \phi^{(k,L)}$ at iteration k and to combine stochastic approximation and Monte Carlo in the approximation step:

$$Q_k(\theta) = Q_{k-1}(\theta) + \gamma_k \left(\frac{1}{L} \sum_{\ell=1}^L \log p(y, \phi^{(k,\ell)}; \theta) - Q_{k-1}(\theta) \right) \quad (2.11)$$

2.1.3 The Simulated Annealing SAEM algorithm

Convergence of SAEM can strongly depend on the initial guess if the likelihood ℓ possesses several local maxima. The Simulated Annealing version of SAEM improves the convergence of the algorithm toward the global maximum of ℓ .

For the sake of simplicity, we will consider here a constant residual error model $g = a$. Let

$$U(y, \phi; \theta) = \frac{1}{2a^2} \sum_{i,j} (y_{ij} - f(x_{ij}, h(\phi_i)))^2 + \frac{1}{2} \sum_{i=1}^N (\phi_i - C_i \mu)' \Omega^{-1} (\phi_i - C_i \mu)$$

Then, we can write the complete likelihood:

$$p(y, \phi; \theta) = C(\theta) e^{-U(y, \phi; \theta)}$$

where $C(\theta)$ is a normalizing constant that only depends on θ .

For any *temperature* $T \geq 0$, we consider the complete model

$$p_T(y, \phi; \theta) = C_T(\theta) e^{-\frac{1}{T} U(y, \phi; \theta)}$$

where $C_T(\theta)$ is a normalizing constant. This model consists in replacing the variance matrix Ω by $T\Omega$ and the residual variance a^2 by Ta^2 . In other words, a model “with a large temperature” is a model with large variances.

We introduce a decreasing temperature sequence $(T_k, 1 \leq k \leq K)$ and use the MCMC-SAEM algorithm considering the complete model $p_{T_k}(y, \phi; \theta)$ at iteration k (while the usual version of MCMC-SAEM uses $T_k = 1$ at each iteration). The sequence (T_k) is large during the first iterations and decreases to 1 with exponential rate. This is done by choosing large initial variances Ω_0 and a_0^2 and setting

$$\tilde{\Omega}_{k+1} = \frac{1}{N} \left(s_{2,k} - \sum_{i=1}^N (C_i \mu_{k+1}) s'_{1,i,k} - \sum_{i=1}^N s_{1,i,k} (C_i \mu_{k+1})' + \sum_{i=1}^N (C_i \mu_{k+1}) (C_i \mu_{k+1})' \right) \quad (2.12)$$

$$a_{k+1} = \sqrt{\frac{s_{3,k}}{N_{tot}}} \quad (2.13)$$

$$\Omega_{k+1} = \max \left(\tau \Omega_k, \tilde{\Omega}_{k+1} \right) \quad (2.14)$$

$$a_{k+1}^2 = \max \left(\tau a_k^2, \frac{s_{3,k}}{N} \right) \quad (2.15)$$

during the first iterations of the algorithm and where $0 \leq \tau \leq 1$.

These large values of the variances make the conditional distribution $p(\phi|y; \theta)$ less concentrated around its mode. This procedure allows the sequence (θ_k) to escape from the local maxima

of the likelihood and to converge to a neighborhood of the global maximum of ℓ . After that, the usual MCMC-SAEM algorithm is used, estimating the variances at each iteration.

Remark 1: The Simulated Annealing version of SAEM is performed during the first K_{sa} iterations. Of course, SAEM without any simulated annealing can be run by setting $\tau = 0$. On the other hand, simulated annealing is obtained with τ close to 1.

Remark 2: We can use two different coefficients τ_1 and τ_2 for Ω and a^2 in `saemix`. It is possible, for example, to choose $\tau_1 < 1$ and $\tau_2 > 1$, with a small initial residual variance and large initial inter-subject variances. In this case, SAEM tries to obtain the best possible fit during the first iterations, allowing a large inter-subject variability. During the next iterations, this variability is reduced and the residual variance increases until reaching the best possible trade-off between these two criteria.

2.2 Estimation of the Fisher Information matrix

Let θ^* be the true unknown value of θ , and let $\hat{\theta}$ be the maximum likelihood estimate of θ . If the observed likelihood function ℓ is sufficiently smooth, asymptotic theory for maximum-likelihood estimation holds and

$$\sqrt{N}(\hat{\theta} - \theta^*) \xrightarrow[N \rightarrow \infty]{} \mathcal{N}(0, I(\theta^*)^{-1}) \quad (2.16)$$

where $I(\theta^*) = -\partial_{\theta}^2 \log \ell(y; \theta^*)$ is the true Fisher information matrix. Thus, an estimate of the asymptotic covariance of $\hat{\theta}$ is the inverse of the Fisher information matrix $I(\hat{\theta}) = -\partial_{\theta}^2 \log \ell(y; \hat{\theta})$.

2.2.1 Linearization of the model

The Fisher information matrix of the non-linear mixed effects model defined in (1) cannot be computed in a closed-form.

An alternative is to approximate this information matrix by the Fisher information matrix of the Gaussian model deduced from the non-linear mixed effects model after linearization of the function f around the conditional expectation of the individual Gaussian parameters $(\mathbb{E}(\phi_i | y; \hat{\theta}), 1 \leq i \leq N)$. The Fisher information matrix of this Gaussian model is a block matrix (no correlations between the estimated fixed effects and the estimated variances). The gradient of f is numerically computed.

Remark 1: We do not recommend the linearization of the model to estimate the parameters of the model, as it is done with the FO and FOCE algorithms. On the other hand, many numerical

experiments have shown that this approach can be used to estimate the Fisher information matrix.

Remark 2: Obviously, this approach cannot be used with discrete data models...

2.2.2 A stochastic approximation of the Fisher Information Matrix

It is possible to obtain an estimation of the Fisher information matrix using the Louis's missing information principle [27]:

$$\partial_{\theta}^2 \log \ell(y; \theta) = E(\partial_{\theta}^2 \log p(y, \phi; \theta) | y; \theta) + \text{Cov}(\partial_{\theta} \log p(y, \phi; \theta) | y; \theta) \quad (2.17)$$

where

$$\begin{aligned} \text{Cov}(\partial_{\theta} \log p(y, \phi; \theta) | y; \theta) &= E(\partial_{\theta} \log p(y, \phi; \theta) \partial_{\theta} \log p(y, \phi; \theta)' | y; \theta) \\ &\quad - E(\partial_{\theta} \log p(y, \phi; \theta) | y; \theta) E(\partial_{\theta} \log p(y, \phi; \theta) | y; \theta)' \end{aligned}$$

and

$$\partial_{\theta} \log g(y; \theta) = E(\partial_{\theta} \log p(y, \phi; \theta) | y; \theta)$$

Here, $\partial_{\theta} u$ is the gradient of u (*i.e.* the vector of first derivatives of u with respect to θ) and $\partial_{\theta}^2 u$ is the hessian of u (*i.e.* the matrix of second derivatives of u with respect to θ).

Then, using SAEM, the matrix $\partial_{\theta}^2 \log \ell(y; \hat{\theta})$ can be approximated by the sequence (H_k) defined as follows:

$$\begin{aligned} \Delta_k &= \Delta_{k-1} + \gamma_k (\partial_{\theta} \log f(y, \phi_k; \theta_k) - \Delta_{k-1}) \\ D_k &= D_{k-1} + \gamma_k (\partial_{\theta}^2 \log f(y, \phi_k; \theta_k) - D_{k-1}) \\ G_k &= G_{k-1} + \gamma_k (\partial_{\theta} \log f(y, \phi_k; \theta_k) \partial_{\theta} \log f(y, \phi_k; \theta_k)^t - G_{k-1}) \\ H_k &= D_k + G_k - \Delta_k \Delta_k^t \end{aligned}$$

In the current version of `saemix`, only the linearisation approach has been implemented.

2.3 Estimation of the individual parameters

When the parameters of the model have been estimated, we can estimate the individual parameters (ψ_i) . To do that, we will estimate the individual normally distributed parameters (ϕ_i) and derive the estimates of (ψ_i) using the transformation $\psi_i = h(\psi_i)$.

Let $\hat{\theta}$ be the estimated value of θ computed with the SAEM algorithm and let $p(\phi_i|y_i; \hat{\theta})$ be the conditional distribution of ϕ_i for $1 \leq i \leq N$.

We use the MCMC procedure used in the SAEM algorithm to estimate these conditional distributions. More precisely, for $1 \leq i \leq N$, we empirically estimate:

- the conditional mode (or Maximum A Posteriori) $m(\phi_i|y_i; \hat{\theta}) = \text{Arg max}_{\phi_i} p(\phi_i|y_i; \hat{\theta})$,
- the conditional mean $E(\phi_i|y_i; \hat{\theta})$,
- the conditional standard deviation $sd(\phi_i|y_i; \hat{\theta})$.

Remarks:

1. The prior distribution of ϕ_i is a normal distribution, but not the conditional distribution $p(\phi_i|y_i; \hat{\theta})$ (remember that the structural model is not a linear function of $\phi_i \dots$). Then, the conditional mode $m(\phi_i|y_i; \hat{\theta})$ and the conditional expectation $E(\phi_i|y_i; \hat{\theta})$ are two different predictors of ϕ_i .
2. If the transformation h is not linear,

$$\begin{aligned} \mathbb{E}(\psi_i|y_i; \hat{\theta}) &= \mathbb{E}(h(\phi_i|y_i; \hat{\theta})) \\ &\neq h(\mathbb{E}(\phi_i|y_i; \hat{\theta})) \end{aligned}$$

In **saemix**, we estimate $\mathbb{E}(\phi_i|y_i; \hat{\theta})$ and $\mathbb{E}(\psi_i|y_i; \hat{\theta})$.

The number of iterations of the MCMC algorithm used to estimate the conditional mean and standard deviation is adaptively chosen as follows:

1. the (ϕ_i) are initialised with the last value obtained in SAEM
2. we run the Hastings-Metropolis with kernel $q^{(1)}$, $q^{(3)}$ and $q^{(4)}$ and compute at each iteration the empirical conditional mean and s.d. of ϕ_i :

$$e_{i,K} = \frac{1}{K} \sum_{k=1}^K \phi_{i,k} \tag{2.18}$$

$$sd_{i,K} = \sqrt{\frac{1}{K} \sum_{k=1}^K \phi_{i,k}^2 - e_{i,K}^2} \tag{2.19}$$

where $\phi_{i,k}$ is the value of ϕ_i at iteration k of the MCMC algorithm.

3. we stop the algorithm at iteration K and use $e_{i,K}$ and $sd_{i,K}$ to estimate the conditional mean and s.d. of ϕ_i if, for any $K - L_{mcmc} + 1 \leq k \leq K$,

$$\begin{aligned} (1 - \rho_{mcmc})\bar{e}_K &\leq \bar{e}_k \leq (1 + \rho_{mcmc})\bar{e}_K \\ (1 - \rho_{mcmc})\bar{sd}_K &\leq \bar{sd}_k \leq (1 + \rho_{mcmc})\bar{sd}_K \end{aligned} \quad (2.20)$$

where $0 < \rho_{mcmc} < 1$. That means that the sequence of empirical means and s.d. must stay in a ρ_{mcmc} -confidence interval during L_{mcmc} iterations.

2.4 Estimation of the likelihood

2.4.1 Linearization of the model

The likelihood of the non-linear mixed effects model defined in (1) cannot be computed in a closed-form.

An alternative is to approximate this likelihood by the likelihood of the Gaussian model deduced from the non-linear mixed effects model after linearization of the function f around the predictions of the individual parameters $(\phi_i, 1 \leq i \leq N)$.

2.4.2 Estimation using importance sampling

The likelihood of the observations can be estimated without any approximation using a Monte-Carlo approach. The likelihood ℓ of the observations can be decomposed as follows:

$$\begin{aligned} \ell(y; \theta) &= \int p(y, \phi; \theta) d\phi \\ &= \int h(y|\phi; \theta) \pi(\phi; \theta) d\phi \end{aligned}$$

where π is the so-called *prior distribution* of ϕ . According to (1.2), π is a Gaussian distribution.

For any distribution $\tilde{\pi}$ absolutely continuous with respect to the prior distribution π , we can write

$$\ell(y; \theta) = \int h(y|\phi; \theta) \frac{\pi(\phi; \theta)}{\tilde{\pi}(\phi; \theta)} \tilde{\pi}(\phi; \theta) d\phi$$

Then, $\ell(y; \theta)$ can be approximated via an *Importance Sampling* integration method:

1. draw $\phi^{(1)}, \phi^{(2)}, \dots, \phi^{(M)}$ with the distribution $\tilde{\pi}(\cdot; \theta)$,

2. let

$$\ell_M(y; \theta) = \frac{1}{M} \sum_{j=1}^M h(y|\phi^{(j)}; \theta) \frac{\pi(\phi^{(j)}; \theta)}{\tilde{\pi}(\phi^{(j)}; \theta)} \quad (2.21)$$

The statistical properties of the estimator $\ell_M(y; \theta)$ of the likelihood $\ell(y; \theta)$ strongly depend on the sampling distribution $\tilde{\pi}$. First, note that

$$\begin{aligned} \mathbb{E}(\ell_M(y; \theta)) &= \ell(y; \theta), \\ \text{Var}(\ell_M(y; \theta)) &= \mathcal{O}(1/M). \end{aligned}$$

Furthermore, if $\tilde{\pi}$ is the conditional distribution $p(\phi|y; \theta)$, the variance of the estimator is null and $\hat{\ell}_M(y; \theta) = \ell(y; \theta)$ for any value of M . That means that an accurate estimation of $\ell(y; \theta)$ can be obtained with a small value of M if the sampling distribution is close to the conditional distribution $p(\phi|y; \theta)$.

In **saemix**, for $i = 1, 2, \dots, N$, we empirically estimate the conditional mean $\mathbb{E}(\phi_i|y_i; \hat{\theta})$ and the conditional variance $\text{Var}(\phi_i|y_i; \hat{\theta})$ of ϕ_i as described above. Then, the $\phi_i^{(j)}$ are drawn with the sampling distribution $\tilde{\pi}$ as follows:

$$\phi_i^{(j)} = \mathbb{E}(\phi_i|y_i; \hat{\theta}) + \text{Var}(\phi_i|y_i; \hat{\theta})^{\frac{1}{2}} \times T_{ij}$$

where (T_{ij}) is a sequence of *i.i.d.* random variables distributed with a t -distribution with ν degrees of freedom. In the current version of **saemix**, the default value is $\nu = 5$.

The quality of the approximation depends on the estimates of the conditional mean and variances of the individual distributions.

2.4.3 Estimation using Gaussian Quadrature

Gauss-Hermite quadrature methods use a fixed set of K_{GQ} ordinates (called nodes) and weights $(x_k, w_k)_{k=1, \dots, K_{GQ}}$ to approximate the likelihood function.

As for importance sampling, the quality of the approximation depends on the estimates of $\mathbb{E}(\phi_i|y_i; \hat{\theta})$ and $\text{Var}(\phi_i|y_i; \hat{\theta})$.

2.5 Model predictions

2.5.1 Population predictions

Population predictions represent the predictions from the model in the absence of data, and they only take into account individual design variables (eg dose regimen) and covariates.

Two types of population predictions are available in **saemix**:

1. the predictions using the population parameters: $f(x_{ij}; h(\mathbb{E}_{\hat{\theta}}(\phi_i))) = f(x_{ij}; h(C_i \hat{\mu}))$. These are provided in the output under the name **ppred**
2. the population mean predictions: $\mathbb{E}_{\hat{\theta}}(f(x_{ij}; \psi_i)) = \mathbb{E}_{\hat{\theta}}(f(x_{ij}; h(\phi_i)))$. These are provided in the output under the name **ypred**

2.5.2 Individual predictions

Individual predictions take into account not only covariates and individual design variables such as dose regimen, but also use the observations in that individual to obtain the parameters providing the best fit for that particular subject, given the population parameters.

In section 2.3, we described how the conditional distribution of the parameters for each individual is obtained in **saemix**. Two types of individual parameters are reported in the output:

1. the conditional mode (or Maximum A Posteriori): $m(\phi_i|y_i; \hat{\theta}) = \text{Arg max}_{\phi_i} p(\phi_i|y_i; \hat{\theta})$. These are reported in the output as **map.psi**
2. the conditional mean: $E(\phi_i|y_i; \hat{\theta})$. These are reported in the output as **cond.mean.psi**

Correspondingly, two types of individual predictions can be obtained in **saemix**:

1. the predictions obtained using the conditional mode are reported in the output as **ipred**
2. the predictions obtained using the conditional mean are reported in the output as **icpred**

2.6 Estimation of the weighted residuals

2.6.1 Population Weighted Residuals

The vector of Population Weighted Residuals are evaluated as:

$$PWRES_i = Var_{\hat{\theta}}(y_i)^{-1/2} (y_i - \hat{y}_i^{pop})$$

where \hat{y}_{ij}^{pop} is the population prediction of y_{ij} and $Var_{\hat{\theta}}(y_{ij})$ is the variance-covariance matrix of y_i .

In **saemix**, weighted residuals are computed using the population mean predictions **ypred** $\mathbb{E}_{\hat{\theta}}(f(x_{ij}; \psi_i)) = \mathbb{E}_{\hat{\theta}}(f(x_{ij}; h(\phi_i)))$ for \hat{y}_{ij}^{pop} . $\mathbb{E}_{\hat{\theta}}(f(x_{ij}; h(\phi_i)))$ and $Var_{\hat{\theta}}(y_{ij})$ are estimated with a Monte-Carlo procedure.

Remark: This computation is performed during the computation of **pd/npde**, so that a basic **saemix** object does not include these elements.

2.6.2 Individual Weighted Residuals

The Individual Weighted Residuals are evaluated as

$$IWRES_{ij} = \frac{y_{ij} - \hat{y}_{ij}^{ind}}{\hat{\sigma}_{ij}^{ind}}$$

where $\hat{y}_{ij}^{ind} = f(x_{ij}; \hat{\psi}_i)$ is the individual prediction of y_{ij} and $(\hat{\sigma}_{ij}^{ind})^2 = g(x_{ij}; \hat{\phi}_i, \hat{\xi})^2$ is the residual variance of y_{ij} .

The two types of individual parameters described in section 2.5 yield two types of individual weighted residuals in the **saemix** output:

1. the individual weighted residuals obtained using the conditional mode are reported in the output as **iwres**
2. the individual weighted residuals obtained using the conditional mean are reported in the output as **icwres**

Remark: When a transformed residual error model is used (an exponential error model for instance), the weighted residuals are computed using $t(y)$ instead of y .

2.6.3 Normalised Prediction Distribution Errors

The Normalised Prediction Distribution Errors are defined as follow

$$\text{npde}_{ij} = \Phi^{-1}(\hat{p}_{ij})$$

where Φ is the $\mathcal{N}(0, 1)$ cumulative distribution function and where \hat{p}_{ij} is an empirical estimator of

$$p_{ij} = \mathbb{P}(Y_{ij} < y_{ij})$$

obtained by Monte-Carlo.

In more details, prediction discrepancies (pd) are first obtained, as the percentile of the observation in the cumulative distribution function F_{ij} of the predictive distribution of Y_{ij} under the model being evaluated. F_{ij} is obtained by simulating K datasets under the model, and the corresponding prediction discrepancies is given by:

$$\text{pd}_{ij} = F_{ij}(y_{ij}) \approx \frac{1}{K} \sum_{k=1}^K \delta_{ijk} \quad (2.22)$$

where $\delta_{ijk} = 1$ if $y_{ij}^{\text{sim}(k)} < y_{ij}$ and 0 otherwise, $y_{ij}^{\text{sim}(k)}$ denoting the value of y_{ij} simulated in the k^{th} replication.

To handle correlations within the observations obtained in the same individual, we first compute the empirical mean $\hat{\mathbf{E}}(\mathbf{y}_i)$ and empirical variance-covariance matrix $\text{Var}((\mathbf{y}_i))$ over the K simulations. Decorrelation is performed simultaneously for simulated data:

$$\mathbf{y}_i^{\text{sim}(k)*} = \hat{\mathbf{V}}_i^{-1/2}(\mathbf{y}_i^{\text{sim}(k)} - \hat{\mathbf{E}}(\mathbf{y}_i)) \quad (2.23)$$

and for observed data:

$$\mathbf{y}_i^* = \hat{\mathbf{V}}_i^{-1/2}(\mathbf{y}_i - \hat{\mathbf{E}}(\mathbf{y}_i)) \quad (2.24)$$

Decorrelated pd are then obtained using the same formula as in (2.22) but with the decorrelated data, and we call the resulting variables prediction distribution errors (pde):

$$\text{pde}_{ij} = F_{ij}^*(y_{ij}^*) \approx \frac{1}{K} \sum_{k=1}^K \delta_{ijk}^* \quad (2.25)$$

where $\delta_{ijk}^* = 1$ if $y_{ij}^{\text{sim}(k)*} < y_{ij}^*$ and 0 otherwise.

Normalised prediction distribution errors (npde) are then obtained as:

$$\text{npde}_{ij} = \Phi^{-1}(\text{pde}_{ij}) \quad (2.26)$$

Remark: the empirical mean and covariance-matrix computed here are also used for the decorrelation step in the computation of the population weighted residuals, *WRES*. The *WRES* in *saemix* are thus computed in conjunction with the more advanced metric npde.

Chapter 3

The saemix package

This chapter presents the input and output of the `saemix` package. You will find the details of the settings and options in here for reference, but readers who wish to apply the methods quickly can directly skip to chapter 4 to find detailed examples of running the package.

Section 3.1 explains how to use the `saemix` library and the many settings that can be tuned, while section 3.2 contains some technical details on the structure of the package and the architecture of the main S4 classes and methods.

3.1 Inputs and outputs

3.1.1 The inputs

To summarise, `saemix` requires to define the model and to fix some parameters used for the algorithms. First, it is necessary to define:

- the structural model, that is the regression function f defined in (1.1),
- the covariate model, that is the structure of the matrix μ defined in (1.2) and the covariates (c_i) .
- the variance-covariance model for the random effects, that is the structure of the variance-covariance matrix Ω defined in (1.2).
- the residual variance model, that is the regression function g .

The only mandatory elements for a `saemix` fit are:

- a data object, defined by at least:
 - the name of the data file
 - we advise to also specify the names of the columns containing the grouping variable, the predictor(s) and the response, although the program will attempt to recognise suitable columns
- a model object, defined by at least:
 - the name of a valid model function
 - the matrix of starting values `psi0`
 - * if no covariates are present in the model, a single line is sufficient, which will contain the starting values for the fixed effects μ in the model
 - * if covariates are present in the model: if `psi0` has more than 1 line, the next lines are assumed to represent the starting values for the covariate models (only parameters actually present in the model will be estimated, even if `psi0` contains non-null values; otherwise, values of 0 will be assumed).

Then, it is necessary to specify several parameters for running the algorithms:

- the SAEM algorithm requires to specify
 - the initial values of the fixed effects μ_0 , the initial variance covariance matrix Ω_0 of the random effects and the initial residual variance coefficients a_0 , b_0 and c_0 ,
 - the sequence of step sizes (γ_k) , that is the numbers of iterations (K_1, K_2) and the coefficients (a_1, a_2) defined in (2.5) and (2.6),
 - the number of burning iterations K_b used with the same value θ_0 before updating the sequence (θ_k) .
- the MCMC algorithm requires to set
 - the number of Markov Chains L ,
 - the numbers m_1 , m_2 , m_3 and m_4 of iterations of the Hasting-Metropolis algorithm,
 - the probability of acceptance ρ for kernel $q^{(3)}$ and $q^{(4)}$,
- the algorithm to estimate the conditional distribution of the (ϕ_i) requires to set
 - the width of the confidence interval ρ_{mcmc} (see (2.20)),
 - the number of iterations L_{mcmc} .
- the Simulated Annealing algorithm requires to set
 - the coefficient τ_1 and τ_2 defining the decrease of the temperature (see (2.14,2.15))

- the number of iterations K_{sa} .
- the Importance Sampling algorithm requires to set
 - the Monte Carlo number M used to estimate the observed likelihood (see (2.21)).
- the Gaussian Quadrature algorithm requires to set
 - the number of quadrature points N_{QG} used to compute each integral (see (2.4.3))
 - the width of each integral N_{QG}

In the R implementation of `saemix`, most of these parameters, as well as other variables used by the algorithm, are set through a list which is included in the object returned by an `saemix fit`. Table 3.1 shows the correspondance between the parameters and the elements in this list.

Parameter	Meaning	Option name	Default value
L	number of Markov Chains	<code>nb.chains</code>	1*
K_1, K_2	Number of iterations during the two periods	<code>nbiter.saemix</code>	<code>c(300,100)</code>
K_b	Number of burning iterations	<code>nbiter.burn</code>	5
m_1, m_2, m_3	Number of iterations of kernels $q^{(2)}$, $q^{(3)}$ and $q^{(4)}$ at each iteration of SAEM	<code>nbiter.mcmc</code>	<code>c(2,2,2)</code>
	Number of iterations during which simulated annealing is performed	<code>nbiter.sa</code>	
ρ	Probability of acceptance for kernels $q^{(2)}$ and $q^{(3)}$	<code>proba.mcmc</code>	0.4
	Stepsize for kernels $q^{(2)}$ and $q^{(3)}$	<code>stepsize.rw</code>	0.4
	Initial variance parameters for kernels $q^{(2)}$ and $q^{(3)}$	<code>rw.init</code>	0.5
τ	Parameter controlling cooling in the Simulated Annealing algorithm	<code>alpha.sa</code>	0.97
M	Number of Monte-Carlo samples used to estimate the likelihood by Importance Sampling	<code>nmc.is</code>	5000
ν	Number of degrees of freedom of the Student distribution used for the estimation of the log-likelihood by Importance Sampling	<code>nu.is</code>	4
K_{GQ}	Number of nodes used for Gaussian Quadrature	<code>nnodes.gq</code>	12
	Width of the distribution used for Gaussian Quadrature (in SD)	<code>nsd.gq</code>	4
L_{mcmc}	Number of iterations required to assume convergence for the conditional estimates	<code>ipar.lmcmc</code>	50
ρ_{mcmc}	Confidence interval for the conditional mean and variance	<code>ipar.rmcmc</code>	0.95

– *To be continued*

Table 3.1 – cont.

Parameter	Meaning	Option name	Default value
Other variables			
	Algorithms to be run in a call to <code>saemix()</code> : a vector of 3 values of 0/1, representing respectively individual parameter estimates (MAP), estimation of the Fisher information matrix and estimation of the LL by importance sampling	<code>algorithms</code>	<code>c(1,1,1)</code>
	Plot graphs during the estimation of the LL by IS	<code>print.is</code>	<code>FALSE</code>
	Maximum number of iterations for the estimation of fixed effects	<code>maxim.maxiter</code>	100
	Whether convergence plots should be drawn at regular intervals during the estimation	<code>displayProgress</code>	<code>TRUE</code>
	Interval (in number of iterations) between two convergence plots	<code>nbdisplay</code>	
	Seed to initialise the random number generator	<code>seed</code>	123456

Table 3.1: Parameters set as options in the `options` list. To set an option, one would define it as an element of this list (see examples), and any option not defined by the user is automatically set to its default value.

* *the default number of chains is 1, except when the number of subjects is smaller than 50, where it defaults to n_c where n_c is the smallest integer such that $n_c N \geq 50$*

Assuming the result of the `saemix` fit has been stored in an object `saemix.fit`, the list of options can be accessed using the following instruction (see section 3.2.2 for more details on how to access elements of objects in R):

```
saemix.fit["options"]
```

For example, to see the number of chains, one would type in R:

```
saemix.fit["options"]$nb.chains
```

The easiest way to set options is to pass them in a list when calling the main fitting function, as can be seen in the example section (section 4.1).

3.1.2 The outputs

In the R implementation of `saemix`, the object returned after a call to the main fitting function `saemix()` contains the following elements:

- `data`: the data object, created by a call to the `saemixData()` function, and containing the dataset to be used in the analysis
- `model`: the model object, created by a call to the `saemixModel()` function, and containing the model characteristics
- `options`: a list containing the options for the estimation algorithm (see above)
- `prefs`: a list containing the graphical preferences for plots, which will be described in the next section
- `results`: the results object
- `rep.data`: the replicated data (when available)
- `sim.data`: the simulated data (when available)

Assuming the result of a call to `saemix()` has been ascribed to the object `yfit`, these elements can be accessed, for example for the results element, with the following command:

```
yfit["results"]
```

The results object is an object of class `SaemixRes`. Most users will not need to access the elements since functions have been created to output the results. However, elements of the results object can also be accessed individually; for example, the likelihood estimated by importance sampling can be accessed as:

```
yfit["results"]["ll.is"]
```

More details on S4 structures (objects and methods), and on how to access the elements of S4 objects can be found in [3.2](#).

Table [3.2](#) shows the most important elements present in the results object (some of these are only present after a call to a specific function, or when the proper option has been set; for instance, estimates of individual parameters are only estimated when the first element of the `algorithm` element in `options` is 1).

Element	Meaning
npar.est	Number of parameter estimates
fixed.effects	Estimates of the fixed effects
se.fixed	Standard errors of estimation of the fixed effects
respar	Estimates of the parameters of the residual error model
se.repar	Standard errors of estimation of the residual parameters
omega	Estimates of the fixed effects
se.omega	Standard errors of the estimation of the fixed effects
ll.is	Log-likelihood estimated by importance sampling
aic.is	AIC using the log-likelihood estimated by importance sampling
bic.is	BIC using the log-likelihood estimated by importance sampling
ll.lin	Log-likelihood estimated by linearisation
aic.lin	AIC using the log-likelihood estimated by linearisation
bic.lin	BIC using the log-likelihood estimated by linearisation
ll.gq	Log-likelihood estimated by gaussian quadrature
aic.gq	AIC using the log-likelihood estimated by gaussian quadrature
bic.gq	BIC using the log-likelihood estimated by gaussian quadrature
map.psi	Individual estimates of the parameters (ψ), obtained as the mode of the conditional distribution (MAP)
map.phi	Estimate of the corresponding individual ϕ
map.eta	Estimate of the corresponding random effect
map.shrinkage	Shrinkage for the MAP estimates
cond.mean.psi	Individual estimates of the parameters, obtained as the mean of the conditional distribution
cond.mean.phi	Estimate of the corresponding individual ϕ
cond.mean.eta	Estimate of the corresponding random effect
cond.var.phi	Estimate of the variance of the individual ϕ
cond.shrinkage	Shrinkage for the conditional estimates
phi.samp	Samples from the individual conditional distribution of the ϕ
phi.samp.var	Variance of the samples from the individual conditional distribution of the ϕ
ypred	Population predictions, computed for the mean population parameters $ypred_{ij} = f(x_{ij}; h(\mathbb{E}_{\hat{\theta}}(\phi_i)))$
ppred	Population mean predictions, obtained as the expectation of the predictions $ppred_{ij} = \mathbb{E}_{\hat{\theta}}(f(x_{ij}; \psi_i))$
ipred	Individual predictions, computed using the MAP estimates of the individual parameters
icpred	Individual predictions, computed using the conditional estimates of the individual parameters
wres	Weighted population residuals, computed using ppred (see section 2.6)
pd	Prediction discrepancies

– To be continued

Table 3.2 – cont.

Element	Meaning
npde	Normalised prediction distribution errors
iwres	Individual weighted residuals, using the MAP estimates of the individual parameters (using the same computations as ipred)
icwres	Individual weighted residuals using the conditional estimates of the individual parameters (using the same computations as icpred)

Table 3.2: Elements contained in the results object.

A full list of all the elements in a results object can be obtained by the command:

```
getSlots("SaemixRes")
```

a) Estimation of the parameters:

The SAEM algorithm computes the maximum likelihood estimate $\hat{\theta}$ and estimates its covariance matrix $I(\hat{\theta})^{-1}/N$ using the Fisher Information Matrix, as defined in Section 2.2.

Recall that d is the number of individual parameters, then for $j = 1, 2 \dots d$, we can

1. estimate the vector of fixed effects μ (intercept and coefficients of the covariates) by $(\hat{\mu})$,
2. estimate the standard errors of μ ,
3. test if some components of μ are null by computing the significance level of the Wald test.

Let $\Omega = (\omega_{jl}, 1 \leq j, l \leq d)$. Then, for any $j, l = 1, 2 \dots d$, we can

1. estimate ω_{jl} by $\hat{\omega}_{jl}$, for all $1 \leq j, l \leq d$,
2. estimate the standard error of $\hat{\omega}_{jl}$, for all $1 \leq j, l \leq d$,

b) Estimation of the conditional distributions:

The MCMC algorithm provides an estimation of the conditional means, conditional modes and conditional standard deviations of the individual parameters and of the random effects.

The function can be called with an argument `nsamp` which runs several sampling chains in parallel, providing several independent samples from the individual conditional distribution for each subject. The number of iterations necessary to obtain convergence (that is, for the successive empirical conditional mean and sd to remain within the requested precision for all chains) is reported, and if the option `displayProgress` is `TRUE`, plots are produced during the estimation process showing the evolution of the different sampling chains.

- the conditional mode can be found in `saemix` in the `results` component of the object, as `map.psi` (there is also a `map.phi` component for the corresponding ϕ and a `map.eta` for the random effects)
- the conditional expectation can be found in `cond.mean.psi` and the variance in `cond.var.psi` (the corresponding ϕ and η are also available)

c) Estimation of the likelihood:

The `saemix` algorithm can provide three different approximations to the likelihoods, through importance sampling, linearisation or gaussian quadrature.

d) Hypothesis testing and model selection:

We can test the covariate model, the covariance model and the residual error model.

The AIC and BIC criteria are defined by

$$AIC = -2 \log \ell_M(y; \hat{\theta}) + 2P \quad (3.1)$$

$$BIC = -2 \log \ell_M(y; \hat{\theta}) + \log(N)P \quad (3.2)$$

where P is the total number of parameters to be estimated and N is the number of subjects. Note that the BIC defined using this formula is in fact the corrected BIC (BICc) proposed by Raftery to better account for the information in mixed-effect models [34]; it differs from the traditional BIC which uses a factor $\log(N_{tot})$ instead of $\log(N)$. The same formula is also used in MONOLIX.

A specific version of BIC can be used for the comparison of covariate models with fixed covariance structure of the random effects [9]:

$$BIC_h = -2 \log \ell_M(y; \hat{\theta}) + \log(N)P_R + \log(N_{tot})P_F \quad (3.3)$$

where P_R (resp. P_F) is the number of estimated parameters in μ that are related to the covariate effects on the random (resp. non random) components of the individual parameters. It is important to note that BIC_h is not appropriate if the compared models do not share the same covariance model.

Joint covariate and random effects selection often leads to the definition of many candidate models whose exhaustive comparison by an information criterion such as AIC or BIC is not possible in a reasonable time. An alternative approach is to use stepwise methods. The algorithm proposed in [10] follows such idea by iteratively combining the classical BIC (3.2) and the hybrid BIC (3.3) for covariance and covariate model selection respectively.

When comparing two nested models \mathcal{M}_0 and \mathcal{M}_1 with dimensions P_0 and P_1 (with $P_1 > P_0$), the Likelihood Ratio Test uses the test statistic

$$LRT = 2(\log \ell_{M,1}(y; \hat{\theta}_1) - \log \ell_{M,0}(y; \hat{\theta}_0))$$

According to the hypotheses to test, the limiting distribution of LRT under the null hypothesis is either a χ^2 distribution, or a mixture of a χ^2 distribution and a $\delta - Dirac$ distribution. For example:

- to test whether some fixed effects are null, assuming the same covariance structure of the random effects, one should use

$$LRT \xrightarrow[N \rightarrow \infty]{} \chi^2(P_1 - P_0)$$

- to test whether some correlations of the covariance matrix Ω are null, assuming the same covariate model, one should use

$$LRT \xrightarrow[N \rightarrow \infty]{} \chi^2(P_1 - P_0)$$

- to test whether the variance of one of the random effects is zero, assuming the same covariate model, one should use

$$LRT \xrightarrow[N \rightarrow \infty]{} \frac{1}{2}\chi^2(1) + \frac{1}{2}\delta_0$$

e) Estimation of the weighted residuals:

The Population Weighted Residuals ($PWRES_{ij}$), the Individual Weighted Residuals ($IWRES_{ij}$) and the Normalised Prediction Distribution Errors ($npde_{ij}$) are computed as described Section 2.6.

3.1.3 Plots

The generic function `plot.saemix` can be used to obtain a number of plots used to assess and diagnose the model. This function is called using the following arguments:

```
plot(saemix.fit,plot.type="plot.type")
```

where `saemix.fit` is the object returned after a successful call to `saemix`, and `"plot.type"` is the type of plot chosen. The following plot types are available:

- `"data"`: spaghetti plot of the data
- `"convergence"`: a plot of the convergence graphs; this is the default type when `type` is not given
- `"likelihood"`: estimate of the likelihood through importance sampling versus the number of MCMC samples
- `"individual.fit"`: plot of the individual fits overlayed on the data, for each subject in the dataset
- `"population.fit"`: plot of the fits obtained with the population parameters and the individual covariates and design, overlayed on the data, for each subject in the dataset
- `"both.fit"`: plot of the individual and population fits, overlayed on the data
- `"observations.vs.predictions"`: observations versus predictions(left: population predictions, right: individual predictions)
- `"random.effects"`: boxplot of the random effects. With the option `"m"`, a horizontal line is added representing the estimate of the population parameter
- `"parameters.versus.covariates"`: plot of a parameter versus all covariates in the model (uses the individual estimates); for continuous covariates, a scatterplot is produced, while for categorical covariates a boxplot is shown. With the option `"m"`, a horizontal line is added representing the estimate of the population parameter. With the options `"l"` or `"s"`, a curve representing a linear regression (`"l"`) or a spline regression (`"s"`) is added. Several options can be combined (see below)
- `"randeff.versus.covariates"`: plot of a random effect versus all covariates in the model (uses the individual estimates)
- `"correlations"`: matrix of scatterplot showing the correlations between pairs of random effects (uses the individual estimates)

- `"marginal.distribution"`: distribution of the random effects
- `"residuals.distribution"`: distribution of the standardised residuals, computed using the population predictions (weighted residuals), the individual predictions (individual weighted residuals) and optionally if available the npde. Both histograms and QQ-plots of the residuals are given
- `"residuals.scatter"`: scatterplot of standardised residuals versus the predictor (X) and versus the predictions. The residuals are computed using the population predictions (weighted residuals), the individual predictions (individual weighted residuals) and optionally if available the npde. The corresponding predictions are the individual predictions for individual residuals, and population predictions for npde and population residuals
- `"vpc"`: Visual Predictive Check; prediction intervals can be added to the plots. To produce prediction intervals, different methods are available for binning (grouping points), which can be selected through the `vpc.method` argument:

`equal`: the quantile of the data are used to define the breaks, yielding a similar number of points in each interval;

`width`: bins of equal width (if the option `xlog` is set to `TRUE`, the bins will be of equal width on the logarithmic scale);

`user`: user-defined breaks (set as the vector in `vpc.breaks` argument; it is possible to give only the inner breaks or to include the boundaries (min/max));

In the first three methods, there will be at most `vpc.bin` bins, and the boundaries of each interval, as well as the value used to plot the corresponding point, will be shown.

- `"npde"`: plots of the npde (distribution, histogram, and scatterplots versus the regression variable and versus predictions), as displayed in the npde library [4]. Tests comparing the empirical distribution of the npde to the theoretical $\mathcal{N}(0, 1)$ distribution by a combined test are also displayed.

Several plots can be produced by setting `plot.type` to be a vector. Partial matching will be used (so that `plot.type="individual"` will produce individual fits, but `plot.type="residuals"` will produce an error message because it could correspond to two different types of plots). After a successful fit, if the option `save.graphs` is `TRUE`, the following plots are produced by default and saved to a file named `diagnostic_graphs.ps` in the directory containing fit results: spaghetti plot of the data, convergence plots, likelihood by importance sampling, plots of predictions versus observations for population and individual estimates, boxplots of the random effects, correlation between the random effects. Individual fits are also saved, in a separate file called `individual_fits.ps`. Some of these plots may be missing if the corresponding estimates have not been requested (eg if the likelihood has not been computed by importance sampling, the plot won't be available).

Each plot can also be obtained individually using a specific function, which allows total flexibility over the layout, including options to change plotting symbols, colors, or which subjects are to be used. Table 3.3 gives the names of the individual functions corresponding to the plots listed above.

Plot function name	Brief description
<code>saemix.plot.data()</code>	Spaghetti plot of the data
<code>saemix.plot.convergence()</code>	Convergence plots for all estimated parameters
<code>saemix.plot.llis()</code>	Plot of the log-likelihood estimated by importance sampling
<code>saemix.plot.obsvspred()</code>	Plot of the predictions versus the observations
<code>saemix.plot.fits()</code>	Individual fit
<code>saemix.plot.distpsi()</code>	Estimated distribution of the random effects
<code>saemix.plot.randeff()</code>	Boxplot of a random effect
<code>saemix.plot.parcov()</code>	Plot of parameters versus covariates
<code>saemix.plot.randeffcov()</code>	Plot of random effects versus covariates
<code>saemix.plot.scatterresiduals()</code>	Scatterplots of residuals versus predictor and predictions
<code>saemix.plot.distribresiduals()</code>	Plot of the distribution of the residuals
<code>saemix.plot.vpc()</code>	Visual Predictive Check
<code>saemix.plot.npde()</code>	Plots of the npde

Table 3.3: Names of the individual functions used to obtain each type of plot. Please refer to the inline help for the arguments to provide to each function.

A help page describing these plots is available in the inline help:

```
?saemix.plot.data
```

A common argument to all the functions is a list of options. This list can be set using the function `saemix.plot.setoptions()`, and it is automatically set during the fit by `saemix()` and stored in the Slot `prefs` of the object. The options can then be modified through this list, for instance changing the new default color to red for all plots is done by setting the attribute `col` in the list:

```
saemix.fit["prefs"]$col<-"red"
```

Options can also be set on the fly for a given plot, by simply adding it to the call to `plot()` as an argument (see examples in section 4.1):

```
plot(saemix.fit,plot.type="data",col="red",main="Raw data")
```

The list of options that can be changed are given in table 3.4, along with their default value. Not all options apply to all graphs.

Parameter	Description	Default value
<i>General graphical options</i>		
ask	Whether users should be prompted before each new plot (if TRUE)	FALSE
new	Whether a new plot should be produced	TRUE
interactive	Whether users should be prompted before predictions or simulations are performed (if TRUE)	FALSE
mfrow	Page layout (NA: layout set by the plot function or before)	NA
main	Title	empty
xlab	Label for the X-axis	empty
ylab	Label for the Y-axis	empty
type	Type of the plot (as in the R plot function)	b (lines and symbols)
col	Main symbol color	black
xlog	Scale for the X-axis (TRUE: logarithmic scale)	FALSE
ylog	Scale for the Y-axis (TRUE: logarithmic scale)	FALSE
cex	A numerical value giving the amount by which plotting text and symbols should be magnified relative to the default	1
cex.axis	Magnification to be used for axis annotation relative to the current setting of 'cex'	1
cex.lab	Magnification to be used for x and y labels relative to the current setting of 'cex'	1
cex.main	Magnification to be used for main titles relative to the current setting of 'cex'	1
pch	Symbol type	20 (dot)
lty	Line type	1 (straight line)
lwd	Line width	1
xlim	Range for the X-axis (NA: ranges set by the plot function)	NA
ylim	Range for the Y-axis (NA: ranges set by the plot function)	NA
ablinecol	Color of the horizontal/vertical lines added to the plots	"DarkRed"
ablinelty	Type of the lines added to the plots	2 (dashed)
ablinelwd	Width of the lines added to the plots	2
<i>Options controlling the type of plots</i>		
ilist	List of subjects to include in the individual plots	all

– To be continued

Table 3.4 – cont.

Parameter	Description	Default value
level	Level of grouping to use (0=population, 1=individual)	0:1
smooth	Whether a smooth should be added to certain plots	FALSE
line.smooth	Type of smoothing (l=line, s=spline)	s
indiv.par	Type of individual estimates (map= conditional mode, eap=conditional mean)	map
which.par	Which parameters to use for the plot	all
which.cov	Which covariates to use for the plot	all
which.pres	Which type of residuals to plot at the population level (when level includes 0)	c("wres","npde")
which.resplot	Type of residual plot ("res.vs.x": scatterplot versus X, "res.vs.pred": scatterplot versus predictions, "dist.hist": histogram, "dist.qqplot": QQ-plot)	c("res.vs.x","res.vs.pred", "dist.qqplot","dist.hist")
Specific graphical options		
obs.col	Symbol color to use for observations	black
ipred.col	Symbol color to use for individual predictions	black
ppred.col	Symbol color to use for population predictions	black
obs.lty	Line type to use for observations	1
ipred.lty	Line type to use for individual predictions	2
ppred.lty	Line type to use for population predictions	3
obs.lwd	Line width to use for observations	1
ipred.lwd	Line width to use for individual predictions	1
ppred.lwd	Line width to use for population predictions	1
obs.pch	Symbol type to use for observations	20
ipred.pch	Symbol type to use for individual predictions	20
ppred.pch	Symbol type to use for population predictions	20
Options for marginal distribution		
indiv.histo	When TRUE, an histogram of the estimates of the individual parameters will be added to the plots of the distribution of the parameters	FALSE
cov.value	The value for each covariate to be used to condition on for the marginal distribution (NA: median will be used)	NA
range	Range (expressed in number of SD) over which to plot the marginal distribution	3

– To be continued

Table 3.4 – cont.

Parameter	Description	Default value
<i>Graphical options for VPC and residual plots</i>		
vpc.method	Method used to bin points (one of "equal", "width", "user" or "optimal"); at least the first two letters of the method need to be specified (the "optimal" method is not implemented yet)	"equal"
vpc.bin	number of binning intervals	10
vpc.interval	size of interval	0.95
vpc.breaks	vector of breaks used with user-defined breaks (vpc.method="user")	NULL
vpc.lambda	value of lambda used to select the optimal number of bins through a penalised criterion	0.3
vpc.pi	whether prediction intervals should be plotted for the median and the limits of the VPC interval	TRUE
vpc.obs	whether observations should be overlayed on the plot	TRUE
fillcol	Color used to fill histograms (individual parameter estimates) or to plot intervals in standard VPC-type plots (VPC, pd, npde)	"lightblue1"
col.fillmed	Color used to fill prediction intervals around the median (for VPC, pd, npde)	"pink"
col.fillpi	Color used to fill prediction intervals around the limits of intervals (for VPC, pd, npde)	"slategray1"
col.lmed	Color used to plot the median of simulated values (for VPC, pd, npde)	"indianred4"
col.lpi	Color used to plot the simulated limit of prediction intervals (for VPC, pd, npde)	"slategray4"
col.pobs	Color used to plot the symbols for observations (for VPC, pd, npde)	"steelblue4"
col.lobs	Color used to plot the line corresponding to given percentiles of observations (for VPC, pd, npde)	"steelblue4"
lty.lmed	Line type used to plot the median of simulated values (for VPC, pd, npde)	2
lty.lpi	Line type used to plot the simulated limit of prediction intervals (for VPC, pd, npde)	2
lty.lobs	Line type used to plot the line corresponding to given percentiles of observations (for VPC, pd, npde)	1

– To be continued

Table 3.4 – cont.

Parameter	Description	Default value
lwd.lmed	Line width used to plot the median of simulated values (for VPC, pd, npde)	2
lwd.lpi	Line width used to plot the simulated limit of prediction intervals (for VPC, pd, npde)	1
lwd.lobs	Line width used to plot the line corresponding to given percentiles of observations (for VPC, pd, npde)	2
Specific graphical options		
pcol	Main symbol color	black
lcol	Main line color	black

Table 3.4: Default graphical parameters. Any option not defined by the user is automatically set to its default value.

3.2 Classes in the saemix package

3.2.1 A very short introduction to S4 classes

This section is in progress. More information on S4 classes and Rpackages can be found in tutorials on the Web. I used extensively the following manual [14].

saemix has been programmed using the S4 classes in R. S4 classes implement Object oriented programming (OOP) in R, allowing to construct modular pieces of code which can be used as black boxes for large systems. Most packages in the base library and many contributed packages use the former class system, called S3. However, S4 classes are a more traditional and complete object oriented system including type checking and multiple dispatching. S4 is implemented in the methods package in base R.

Elements of an object are called "Slots". Slots can be accessed using the @ operator, instead of the \$ operator used for lists. However, the use of @ to access the class slots is heavily discouraged outside functions programmed directly by package developers. Instead, in saemix accessor functions ("get" functions) and replacement functions ("set" functions) have been defined to allow access to elements of an object in the same way as one would the elements of a list in R,

through the name of the slot. If `obj` is an object with a slot called "slot", we would display the value of the slot using the command:

```
obj["slot"]
```

Assuming that slot is a character string, we can replace its value by "my string" using the command:

```
obj["slot"]<-"my string"
```

Since an S4 class has built-in check for types, a command such as:

```
obj["slot"]<-3
```

would in this case produce an error.

3.2.2 S4 classes used in `saemix`

Visible S4 objects

The following classes have been defined in `saemix`:

- **SaemixData**: this object contains the structure and data of the longitudinal dataset
- **SaemixModel**: this object contains the structure representing a non-linear mixed effect model, used by the **SAEM** algorithm
- **SaemixRes**: this object contains the results obtained after a fit by `saemix()`; it is included in the **SaemixObject** as the Slot **results**
- **SaemixObject**: this is the object returned by a call to `saemix()`; this object has the following slots:
 - **data**: a **SaemixData** object, containing the structure and data of the longitudinal dataset
 - **model**: a **SaemixModel** object, containing the characteristics of the non-linear mixed effect model
 - **results**: a **SaemixRes** object, containing the results obtained after a fit by `saemix()`
 - **options**: a list of options

- `prefs`: a list of graphical preferences, that will replace the default graphical preferences if changed; the preferences set in this list can be superseded by setting an option in the call to the plot functions (see section 3.1.3)
- `rep.data`: an object of class `SaemixRepData` produced during the fit of the SAEM algorithm (for internal use only)
- `sim.data`: an object of class `SaemixSimData` containing data simulated according to the design of the original dataset and the fitted model, with the results obtained during the fit

The constructor functions for the first two objects are respectively `saemixData()` and `saemixModel()` (with a lowercase initial letter, to distinguish it from the object classes, which start with a capital letter, since in R lowercase and uppercase letters are different). These two functions are the functions intended to be used directly to produce the objects given as input to the `saemix()` function.

- `saemixData()`: the `saemixData()` function requires one mandatory argument, the name of a dataframe in R or of a file on disk containing the data. If the file has a header (or if the dataframe has column names), the program will attempt to recognise suitable names for the grouping, predictor and response variables. These may also be specified by the user, either as names or column numbers (see help page for `SaemixData`).
- `saemixModel()`: the `saemixModel()` function requires two mandatory arguments: the name of a Rfunction computing the model in the *saemix* format (see details and examples) and a matrix giving the initial estimates of the fixed parameters in the model. This matrix should contain at least one row, with the values of the initial estimates for the population mean parameters; if covariates are present in the data and enter the model, a second row should contain the values of the initial estimates for the covariate effects.

There is no constructor function for an `SaemixObject` object, since such an object should be returned by the `saemix()` function.

Hidden S4 objects

In addition to the visible objects, *saemix* also has 2 other classes which are not intended to be used directly by the user:

- `SaemixRepData`: this object is created during the fit by `saemix()`
- `SaemixSimData`: this object is created when simulating data

An `SaemixObject` contains instances of these two classes. The slot of class `SaemixRepData` is produced and filled during the fit, while the slot of class `SaemixSimData` is produced when simulations are performed (in particular, to compute weighted residuals and npde, and produce VPC plots).

3.2.3 Methods for S4 objects in `saemix`

Two types of functions have been developed for the `saemix` package:

- methods
- classical functions

Methods are a special type of functions, which apply to objects and benefit from multiple dispatch. R uses multiple dispatch extensively: one generic function call, such as for instance `print`, is capable of dispatching on the type of its argument and calls a printing function that is specific to the data supplied. For instance, using the `print()` function on a matrix will output the matrix, while using the same function on an object returned by the `lm()` function will produce a summary of the linear regression fit. We used this feature to produce notably `plot()` and `print()` functions (see next sections) which should apply to our `saemix` objects in a user-friendly way.

Generic methods

The following generic methods have been defined for `SaemixData`, `SaemixModel` and `SaemixObject` objects:

- `print`: the `print` function produces a summary of the object in a nice format
- `show`: this function is used invisibly by R when the name of the object is typed, and produces a short summary of the object
- `summary`: this function produces a summary of the object, and invisibly returns a list with a number of elements, which provides an alternative way to access elements of the class
 - for `SaemixData`, the list contains `ntot` (total number of observations), `nind` (vector containing the number of observations for each subject), `id` (vector of identifier), `xind` (matrix of predictors), `cov` (matrix of individual covariates), `y` (observations);
 - for `SaemixModel`, the list contains the model function, the error model, the list of parameters, the covariance structure, the covariate model;

- for **SaemixObject**, the list contains the estimated fixed effects, the estimated parameters of the residual error model, the estimated variability of the random effects, the correlation matrix, the log-likelihood by the different methods used, the Fisher information matrix, the population and individual estimates of the parameters for each subject, the fitted values, the residuals.
- **plot**: this produces plots of the different objects
 - for **SaemixData**, a plot of the data is produced. The default plot is a spaghetti plot of the response variable versus the predictor (if several predictors, this is the predictor given by `name.X`) with a different line for each individual
 - for **SaemixModel**, the model is used to predict the value of the response variable according to the value of the predictor(s) over a given range of values for the main predictor.
 - for **SaemixObject**, the plot function produces a number of different plots. By default, a series of plot are produced; when called with the `plot.type` argument, selected plots can be chosen.
- `[` function: the get function, used to access the value of the slots in an object
- `[<-` function: the set function, used to replace the value of the slots in an object

Examples of calls to these functions are given in the corresponding man pages and in the documentation (chapter 4). Additional generic methods for classes, such as `initialize()`, are not user-level in the *saemix* package.

Specific methods

Specific methods have been developed for the objects in the *saemix* package. Specific methods are methods which possibly apply to objects of several classes. For all purposes, they are used like generic methods.

The following methods apply to **SaemixObject** objects:

- **showall**: this method produces an extensive summary of the object. This method is also defined for **SaemixData** and **SaemixModel** objects.
- **predict**: this function uses the results from an SAEM fit to obtain model predictions for the data in the `data` element of the **SaemixObject** object
- **psi**, **phi**, **eta**: these three methods are used to access the estimates of the individual parameters and random effects. When the object passed to the function does not contain

these estimates, they are automatically computed. The object is then returned (invisibly) with these estimates added to the results

- `coef`: this method extracts the coefficients from an `SaemixObject` fit, returning a list with three components (some components may be empty (eg MAP estimates) if they have not been computed during or after the fit)
 - `fixed`: estimated fixed effects in the model
 - `population`: population parameter estimates for each subject; the estimation of population parameters includes individual covariates if some enter the model; this is a list with two components, `map` and `cond`, which are respectively the MAP estimates and the conditional mean estimates
 - `individual`: individual parameter estimates: a list with two components `map` and `cond`; this is a list with two components, `map` and `cond`, which are respectively the MAP estimates and the conditional mean estimates of the individual parameters

Additional specific methods have been defined but are not user-level (`read.saemixData()` is used by the constructor function).

3.2.4 Accessing S4 objects in `saemix`

Help for S4 objects and methods

Aliases for the `SaemixData`, `SaemixModel` and `SaemixObject` objects have been created, so that the usual online help can be called:

```
help(SaemixData)
?SaemixData
```

Classic methods are accessed by the usual help function, for example:

```
?saemix
```

will produce the help file for the main `saemix()` function, fitting the non-linear mixed effect model.

The help files for generic and methods on the other hand can be accessed by the following (non-intuitive) commands:

```
help("plot,SaemixData")
```

Typing:

```
help(plot)
```

will only give the help page for the generic `Rplot` function. In the same way, we access the help page for the `plot` function applied to the object resulting from a call to `saemix()` (which contains links to the page describing the specific plots):

```
help("plot,SaemixObject")
```

Elements for S4 objects defined in *saemix*

The elements, or slots, of the objects with class `SaemixData`, `SaemixModel` and `SaemixObject` are described in the respective help pages. When an object is first created, some of its slots may be empty or filled in with default values.

In the following, we create the object `saemix.data` by a call to the constructor function:

```
data(theo.saemix)
saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA,
name.group=c("Id"),name.predictors=c("Dose","Time"),name.response=c("Concentration"),
name.covariates=c("Weight","Sex"),units=list(x="hr",y="mg/L",covariates=c("kg","-")),
name.X="Time")
```

We can then access the number of subjects in the dataset by the `get` function:

```
saemix.data["N"]
```

Warning: modifying the elements in the objects outside of dedicated functions or methods can have unwanted side-effects. For instance, if one was to change the number of subjects in the `data` slot of an object created by a call to `saemix()`, the consistency of the object would not be guaranteed, and this could cause strange behaviour when trying to print or plot the object, or use it in subsequent functions. For this reason it is strongly recommended to only use the functions and methods defined in *saemix* to access and modify *saemix* objects. For instance, to apply the SAEM algorithm only to a subset of the subjects, it is preferable to apply the function `saemixData()` to a subset of the data instead of trying to change directly the `SaemixData` object.

Chapter 4

Examples

4.1 Theophylline pharmacokinetics

4.1.1 One-compartment model

Boeckmann, Sheiner and Beal (1994) report data from a study by Dr. Robert Upton of the kinetics of the anti-asthmatic drug theophylline [42]. Twelve subjects were given oral doses of the anti-asthmatic drug theophylline, then serum concentrations (in mg/L) were measured at 11 time points over the next 25 hours. In the present package, we removed the data at time 0 to avoid some unexplained non-zero values in a supposedly single-dose study. In the original dataset shipped with the **NONMEM**, doses are given as doses per kilo body weight. Here, we therefore also transformed the doses to doses in mg instead of mg/kg.

These data are analyzed in Davidian and Giltinan [6] and Pinheiro and Bates [30] using a two-compartment open pharmacokinetic model. These data are also available in R in the library **datasets** under the name **Theoph** in a slightly modified format and including the data at time 0.

Subject i receives an initial dose D_i at time 0 and serum concentrations y_{ij} are measured at time t_{ij} . Serum concentration is modeled by a first-order one compartment model, according to the following equation:

$$y_{ij} = \frac{D_i k_{ai} k_{ei}}{CL_i (k_{ai} - k_{ei})} \left(e^{-k_{ai} t_{ij}} - e^{-k_{ei} t_{ij}} \right) + \epsilon_{ij} \quad (4.1)$$

where CL_i is the clearance of subject i , k_{ai} is the absorption rate constant, k_{ei} is the elimination rate constant and is expressed as a function of CL_i and the volume of distribution V_i as $k_{ei} = \frac{CL_i}{V_i}$. For subject i :

- the vector of regression (or design) variables is $x_{ij} = (D_i, t_{ij})$
- the vector of individual parameters is $\theta_i = (\ln(k_{ai}), \ln(CL_i), \ln(V_i))$
 - k_{ai} , CL_i and V_i are assumed to be independant log-normal random variables
 - we assumed a relationship between the clearance and the subject's body weight BW_i

$$\begin{aligned}\ln(k_{ai}) &= \mu_1 + \eta_1 \\ \ln(V_i) &= \mu_2 + \eta_2 \\ \ln(CL_i) &= \mu_3 + \beta BW_i + \eta_3\end{aligned}\tag{4.2}$$

- we can use a simple homoscedastic error model where $\text{Var}(\epsilon_{ij}) = a^2$

The data is shown in figure 4.1.

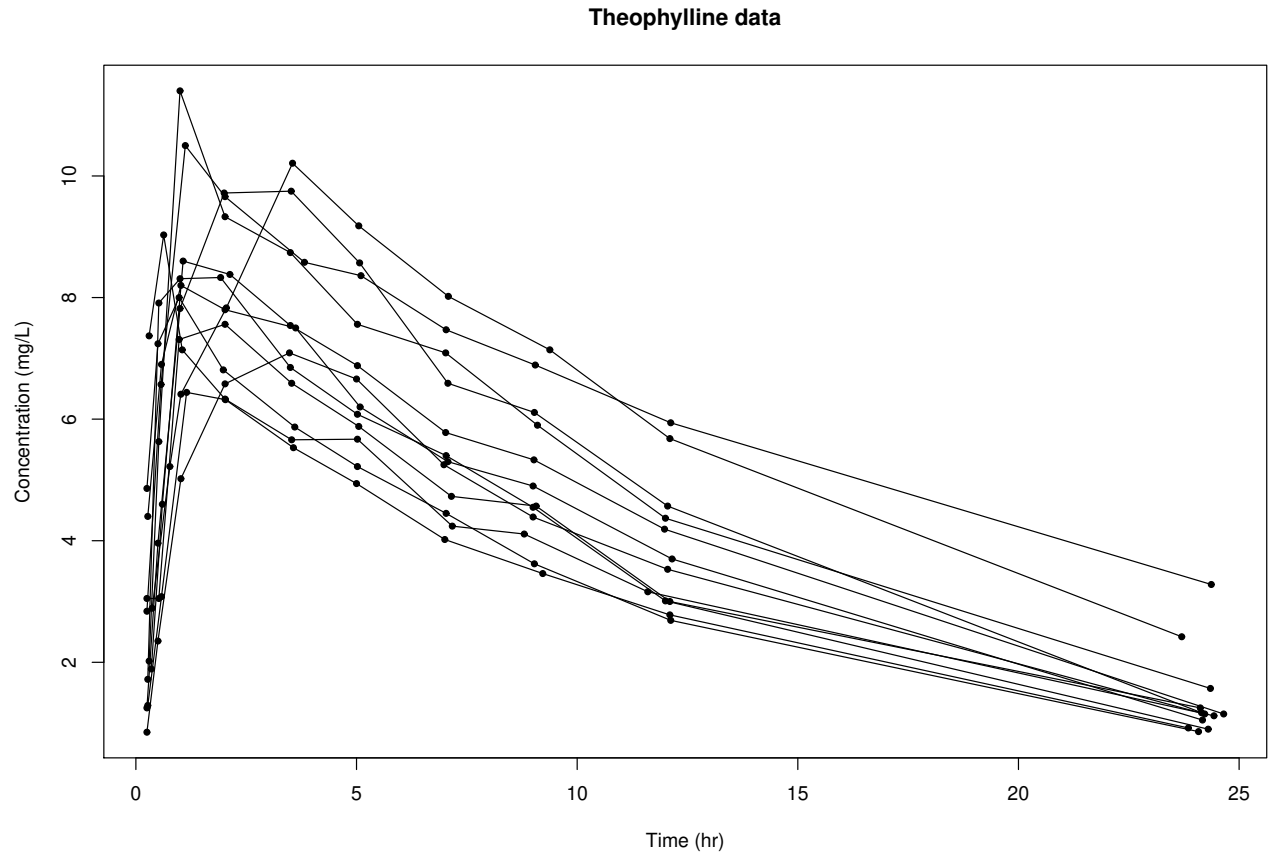


Figure 4.1: Theophylline concentrations versus time for the 12 subjects included in the study.

The following code was used in R to read the data, using the `saemixData` to create the structured object we will use in the analysis, and the `plot` function developed for `SaemixData` objects to obtain the plot in figure 4.1:

```
library(saemix)

data(theo.saemix)
saemix.data<-saemixData(name.data=theo.saemix,header=TRUE,sep=" ",na=NA,
  name.group=c("Id"),name.predictors=c("Dose", "Time"),name.response=c("Concentration"),
  name.covariates=c("Weight","Sex"),units=list(x="hr",y="mg/L",covariates=c("kg","-")),
  name.X="Time")

plot(saemix.data)
```

The `name.X` is used to specify which of the predictors should be used on the X-axis, it is optional when the first predictor in the list.

The following code is used to specify the model:

```
model1cpt<-function(psi,id,xidep) {
  tim<-xidep[,2]
  dose<-xidep[,1]
  ka<-psi[id,1]
  V<-psi[id,2]
  CL<-psi[id,3]
  k<-CL/V
  ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
  return(ypred)
}
saemix.model<-saemixModel(model=model1cpt,
  description="One-compartment model with first-order absorption",
  psi0=matrix(c(1.,20,0.5,0.1,0,-0.01),ncol=3, byrow=TRUE,dimnames=list(NULL,
    c("ka", "V", "CL"))),transform.par=c(1,1,1),
  covariate.model=matrix(c(0,0,1,0,0,0),ncol=3,byrow=TRUE),fixed.estim=c(1,1,1),
  covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
  omega.init=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE), error.model="constant")
```

In this example, we specify the vector of starting values through `psi0`, which is defined as the following matrix:

	ka	V	CL
[1,]	1.0	20	0.50
[2,]	0.1	0	-0.01

The first line is renamed as `Pop.CondInit` when the model object is created (see output from the commands given in the snippet of code above), and contains the initial estimates of the

population parameters k_a , V and CL . The second line, renamed `Cov.CondInit` in this example, contains the initial values for the parameter-covariate relationships in the model. In this example, we have assumed an effect of the covariate `Weight` on the clearance CL , and the initial value of the corresponding fixed effect is -0.01. In this model there is no relationship between either of the two covariates in the model and k_a , so that the 0.1 value given in `psi0` will not be used. If we also had relationships between the covariate `Sex` and the model parameters, the same starting values would be used (using the vector recycling principle `R`), however we could add other lines to `psi0` to specify different starting values. For example, assuming we want to estimate an effect of weight on V and CL , as well as a gender effect on CL , we could replace the `covariate.model` argument with:

```
covariate.model=matrix(c(0,1,1,0,0,1),ncol=3,byrow=TRUE)
```

and give different starting values for each parameter-relationship in `psi0`, for example 0.1 for both weight effects and -0.1 for the gender effect:

```
psi0=matrix(c(1.,20,0.5,0,0.1,0.1,0,0,-0.1),ncol=3,byrow=TRUE,
  dimnames=list(NULL, c("ka","V","CL")))
```

Note that the model requires two predictors, dose and time. The user is responsible for writing the model function and checking the consistency between the model function and the data. Here, the first predictor (first column) is dose and the second predictor is time so that we need both items in the dataset, and we need to give the names of the two predictors in the proper order (the order corresponding to the way the model function is written here) when creating the data object. This is a single-dose administration and therefore the dose column contains the same dose repeated for each time-point. However, for graphs we want the observations to be plotted versus time and not versus dose; by default, the program will use the first predictor as the X axis, but we override this behaviour here by setting the option `name.X="Time"` in the creation of the data object, so that the graphs will use time on the X-axis.

Then we fit the model using the `saemix()` function:

```
saemix.fit<-saemix(saemix.model,saemix.data,list(seed=632545,nb.chains=5,
  nbiter.saemix = c(300, 150)))
```

We use 5 chains here to stabilise the estimation because there are only 12 subjects in the dataset (by default, the algorithm will increase the number of chains if there are less than 50 subjects in the dataset, and set it to a higher value as describe in section 3.1), and we increase the number of steps in the second stage to 150 (default: 100) to show how to set this option. Increasing the number of iterations in the second stage helps to obtain a more stable conditional distribution for the individual parameters.

This produces the following output:

```

.....
Nonlinear mixed-effects model fit by the SAEM algorithm
-----
----          Data          ----
-----

Object of class SaemixData
  longitudinal data for use with the SAEM algorithm
Dataset theo.saemix
  Structured data: Concentration ~ Dose + Time | Id
  X variable for graphs: Time (hr)
  covariates: Weight (kg), Sex (-)
First 10 lines of data:
   Id   Dose  Time Concentration Weight Sex
1   1 319.992 0.25           2.84   79.6   1
2   1 319.992 0.57           6.57   79.6   1
3   1 319.992 1.12          10.50   79.6   1
4   1 319.992 2.02           9.66   79.6   1
5   1 319.992 3.82           8.58   79.6   1
6   1 319.992 5.10           8.36   79.6   1
7   1 319.992 7.03           7.47   79.6   1
8   1 319.992 9.05           6.89   79.6   1
9   1 319.992 12.12          5.94   79.6   1
10  1 319.992 24.37           3.28   79.6   1
-----
----          Model          ----
-----

Nonlinear mixed-effects model
  Model function: One-compartment model with first-order absorption
function(psi,id,xidep) {
  dose<-xidep[,1]
  tim<-xidep[,2]
  ka<-psi[id,1]
  V<-psi[id,2]
  CL<-psi[id,3]
  k<-CL/V
  ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)-exp(-ka*tim))
  return(ypred)
}
  Nb of parameters: 3
  parameter names:  ka V CL

```

```

distribution:
Parameter Distribution
[1,] ka          log-normal
[2,] V           log-normal
[3,] CL          log-normal
Variance-covariance matrix:
ka V CL
ka  1 0  0
V   0 1  0
CL  0 0  1
Error model: constant , initial values: a=1
Covariate model:
ka V CL
Weight 0 0 1
Initial values
ka V CL
PopCI 1.0 20 0.50
CovCI 0.1 0 -0.01
-----
---- Key algorithm options ----
-----
Algorithms: MAP, FIM, LL by IS
Number of iterations: K1=300, K2=150
Number of chains: 5
Seed: 632545
Number of MCMC iterations for IS: 5000
Simulations:
nb of simulated datasets used for npde: 1000
nb of simulated datasets used for VPC: 100
Input/output
save the results to file pop_parameters.txt in directory: newdir
save graphs
-----
---- Results ----
-----
----- Fixed effects -----
-----
Parameter      Estimate SE      CV(%) p-value
[1,] ka         1.567  0.2998  19.1 -
[2,] V          31.475  1.3838   4.4 -
[3,] CL          1.581  1.0155  64.2 -
[4,] beta_Weight(CL) 0.008  0.0092 113.5 0.19

```



```

[5,] a          0.743   0.0569   7.7 -
-----
----- Variance of random effects -----
-----
      Parameter Estimate SE      CV(%)
ka omega.ka  0.388    0.175 45
V  omega.V   0.015    0.009 59
CL omega.CL  0.070    0.034 49
-----
----- Correlation matrix of random effects -----
-----
      omega.ka omega.V omega.CL
omega.ka 1      0      0
omega.V  0      1      0
omega.CL 0      0      1
-----
----- Statistical criteria -----
-----
Likelihood computed by linearisation
      -2LL= 343.4919
      AIC = 359.4919
      BIC = 363.3712

Likelihood computed by importance sampling
      -2LL= 344.8896
      AIC = 360.8896
      BIC = 364.7689
-----

```

By default, the results are saved in a file called `pop_parameters.txt` in the `newdir` directory, and graphs are produced.

Table 4.1 reports the parameters obtained on a Linux Ubuntu distribution running R version 2.11.1 for this example. In this example, the fixed effect representing the influence of weight on CL is not significant ($p=0.19$, NS according to a Wald test).

A series of diagnostic plots can be produced simply by applying the function `plot()` to the object returned by `saemix()`:

```
plot(saemix.fit)
```

By using the `plot.type=""` argument, specific graphs can be produced (see section 3.1.3). For example, the convergence plot shown in figure 4.2 can be produced by:

Parameter	Population estimate (SE%)	IIV Variance (SE%)
k_a (hr^{-1})	1.57 (19%)	0.39 (45%)
CL (L.hr^{-1})	1.58 (64%)	0.07 (49%)
$\beta_{BW,CL}$ (-)	0.008 (110%)	-
V (L)	31.5 (4%)	0.02 (59%)
a (mg.L^{-1})	0.74 (6%)	-

Table 4.1: Pharmacokinetic parameters estimated by `saemix` for the theophylline data.

```
plot(saemix.fit,plot.type="convergence")
```

In this figure we can see all the parameters converging quickly to their estimated value.

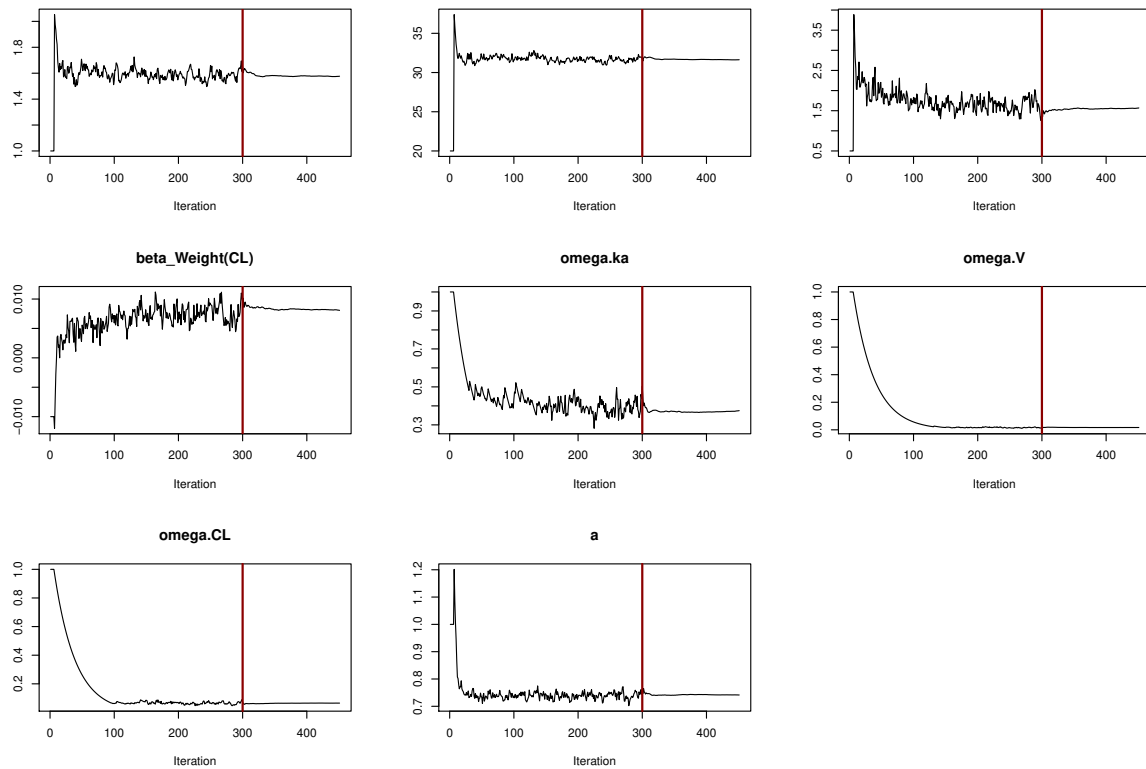


Figure 4.2: Convergence plots for the estimated pharmacokinetic parameters and the variabilities.

Figure 4.3 shows the evolution of the log-likelihood during the importance sampling step. Figure 4.4 shows the predicted values compared to the observed concentrations, for the population predictions (left) and the individual predictions (right). Figure 4.5 shows the individual data for the 12 subjects, with the individual predictions overlayed (smoothed predictions were obtained). Both

plots indicate good model adequacy.

-2xLL by Importance Sampling

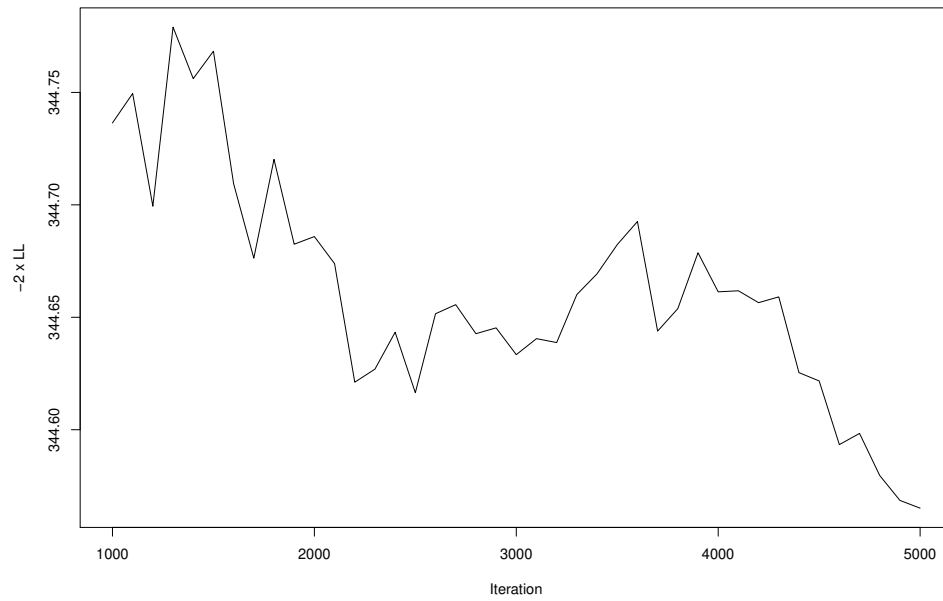


Figure 4.3: Estimating the log-likelihood by Importance Sampling.

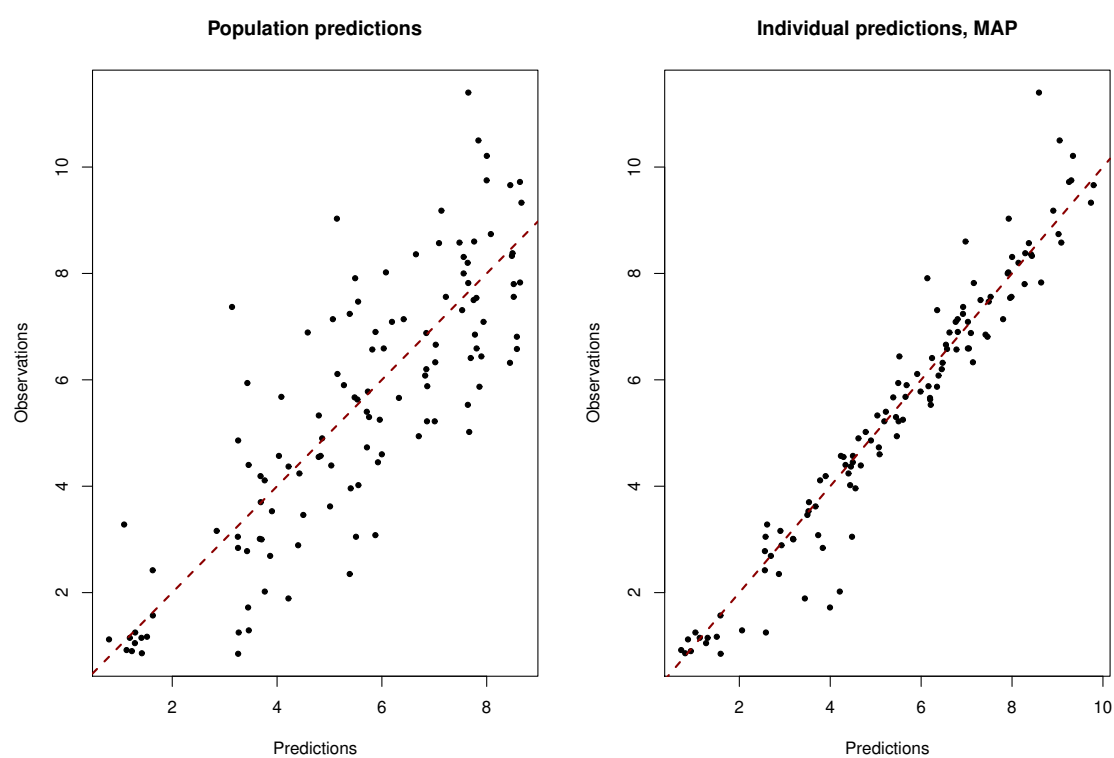


Figure 4.4: Observations versus predictions (left: population predictions, right: individual predictions).

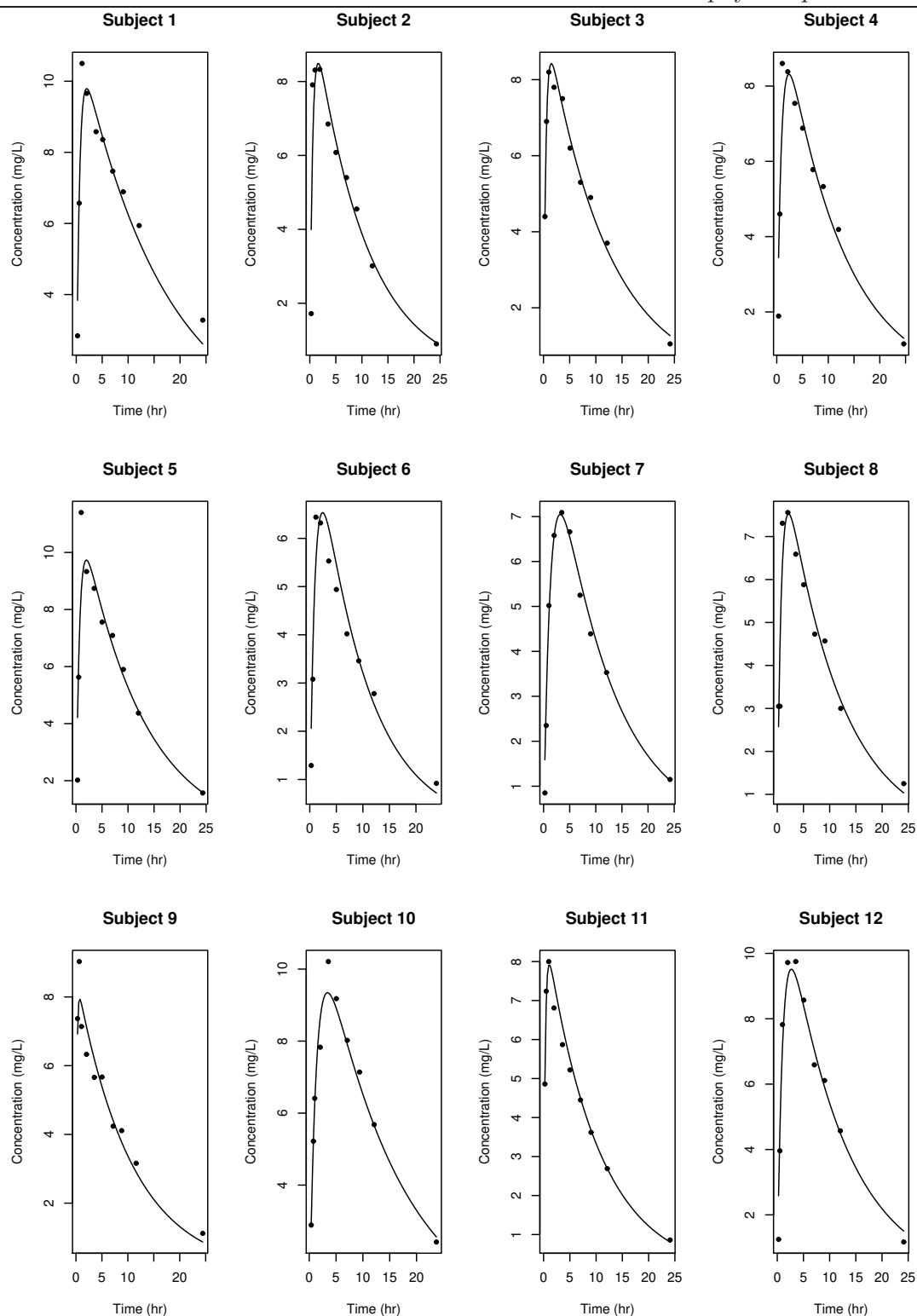


Figure 4.5: Individual plots for the 12 subjects in the study. Dots represent observations and the line shows the smoothed profile predicted using the individual estimated parameters.

The following example shows how to use the functions defined in section 3.1.3 to plot the individual fits for the first 4 subjects in the theophylline example, including a smoothed prediction line, and changing the color of the line and the plotting symbol. A logarithmic scale is used for the Y-axis. The resulting plot is shown in figure 4.6

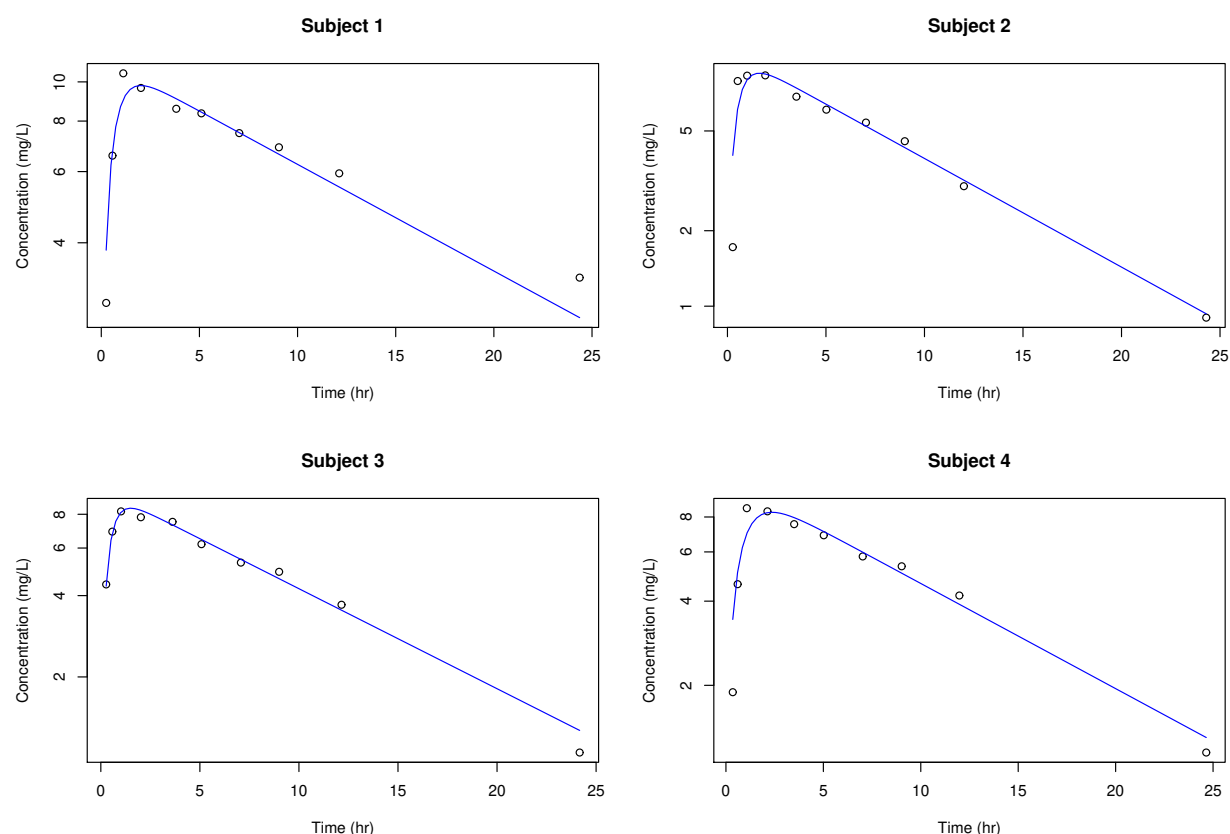


Figure 4.6: Individual plots for the first 4 subjects in the study, with different options.

To obtain these plots, we can use the generic function `plot()`, by setting the `plot.type` argument to "individual.fit", to produce these plots:

```
# Plotting individual fits with selected options
par(mfrow=c(2,2))
plot(saemix.fit,plot.type="individual.fit",new=FALSE,ilist=1:4,smooth=TRUE,ylog=T,
pch=1, col="Blue",xlab="Time in hr",ylab="Theophylline concentrations (mg/L)")
```

We can also use directly the `saemix.plot.fits()` function with the same graphical options, which gives the exact same graph:

```
# Plotting individual fits with selected options
par(mfrow=c(2,2))
saemix.plot.fits(saemix.fit,new=FALSE,ilist=1:4,smooth=TRUE,ylog=T,pch=1,
col="Blue",xlab="Time in hr",ylab="Theophylline concentrations (mg/L)")
```

Other diagnostic plots include Visual Predictive Checks, shown in figure 4.7, and residual plots.

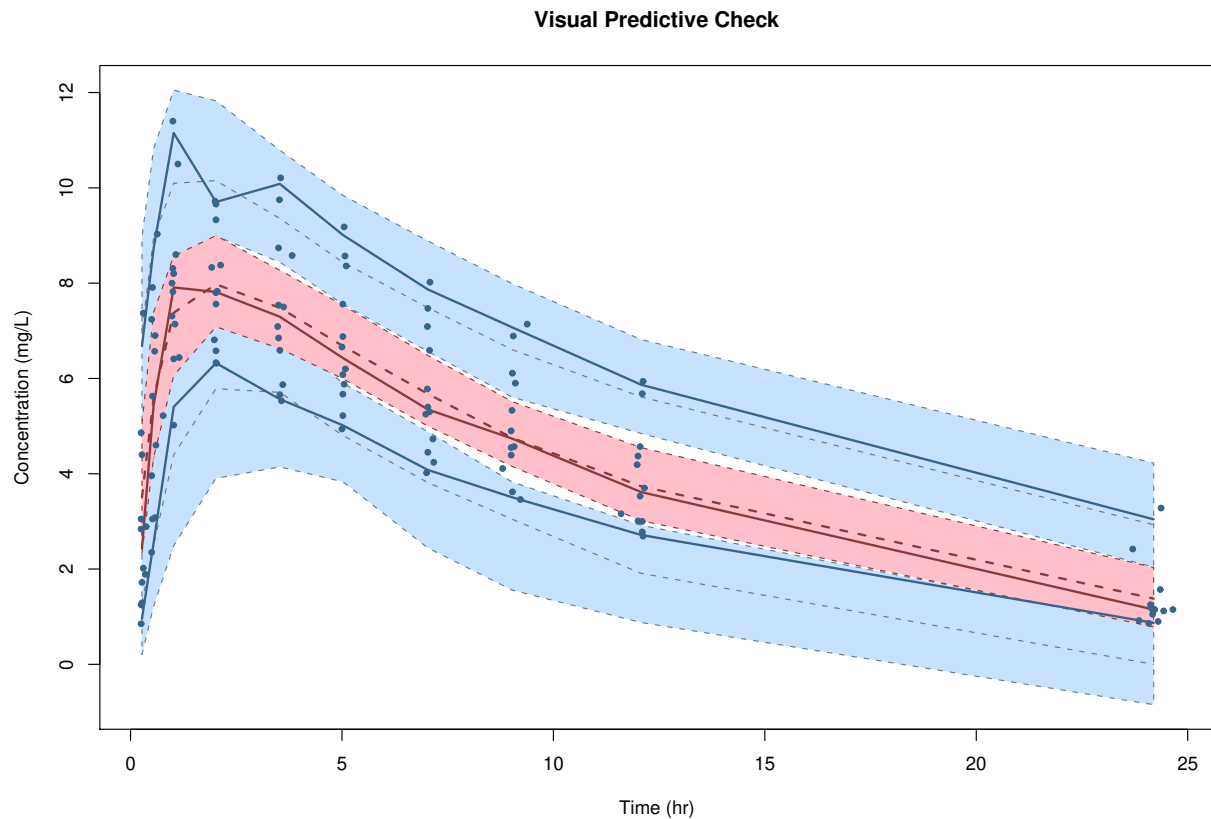


Figure 4.7: VPC for the theophylline data.

The following code can be used to first simulate from the model in order to compute simulation-based metrics (residuals and VPC), and then produce VPC and scatterplots of residuals versus time and predictions (figure 4.8).

```
# Scatterplots of residuals
plot(saemix.fit, plot.type="residuals.scatter")

# VPC
```

```
plot(saemix.fit, plot.type="vpc")
```

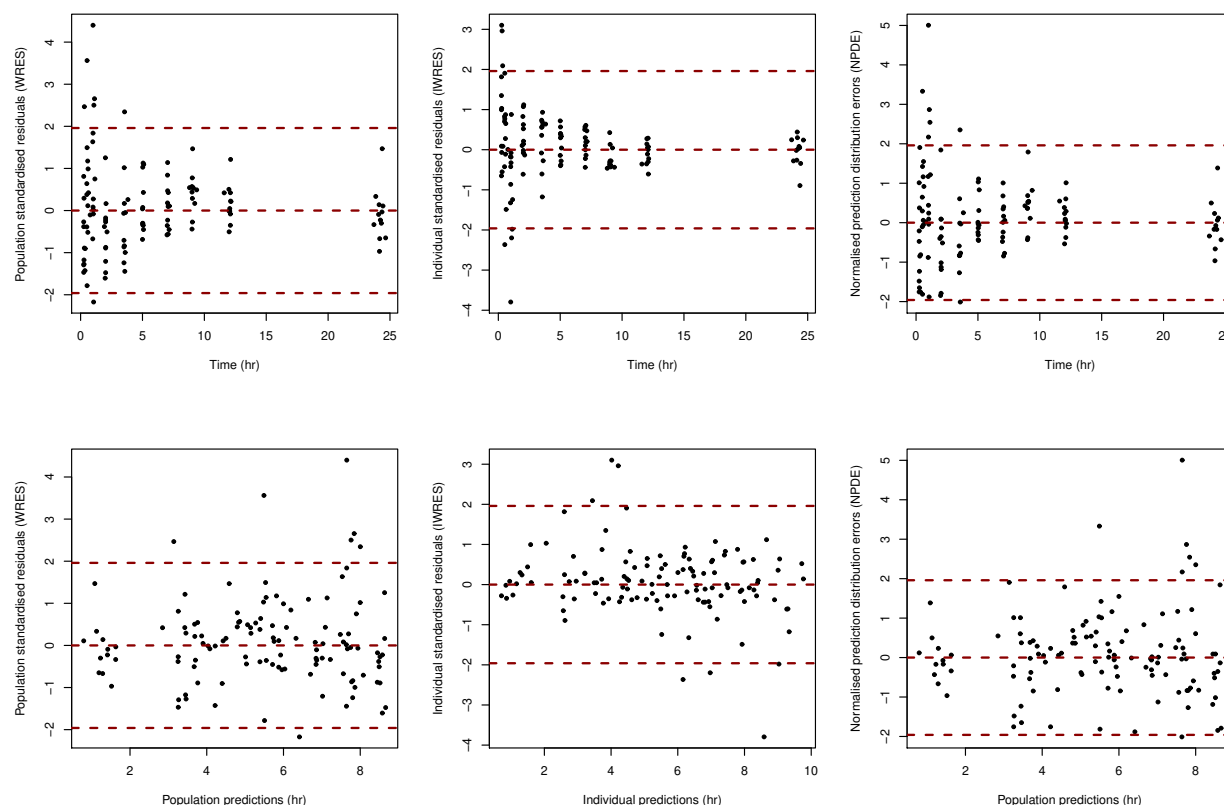


Figure 4.8: Scatterplots of the residuals (left: weighted population residuals; middle: individual weighted residuals; right: npde) versus time (top) and predictions (bottom).

Finally, note that the SAEM algorithm is relatively robust to the initial choice of parameter estimates, but different initial choices may lead to different population estimates. Here, if we had set all the initial parameters to 1 as in the following code, the model converges to very different values and a flip-flop occurs (k_a becomes smaller than the elimination rate constant $k = CL/V$). The resulting fit however has a lower likelihood and the VPC graphs indicate poor estimates of the variability (not shown), which can give an indication of problems with the model.

```
saemix.model<-saemixModel(model=model1cpt,
description="One-compartment model with first-order absorption",
psi0=matrix(c(1.,1.,1.,0.1,0,-0.01),ncol=3, byrow=TRUE,dimnames=list(NULL,
c("ka","V","CL"))),transform.par=c(1,1,1),
covariate.model=matrix(c(0,0,1,0,0,0),ncol=3,byrow=TRUE), fixed.estim=c(1,1,1),
covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
```



```
omega.init=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE), error.model="constant")
```

Thus, it is always good policy during data analysis to check the stability of the final model estimates by changing the initial estimates and running the algorithm again, and to compare the magnitude of the parameter estimates with a reference, such as prior information or literature values.

4.1.2 One-compartment model at steady-state

In the theophylline example, we described the pharmacokinetics using the single-dose, first-order absorption and elimination model. The following code shows how to fit the same data with the same model at steady-state, assuming a 24 hours dosing interval:

```
data(theo.saemix)
# Include a column for the inter-dose interval (tau)
theo.saemix2<-cbind(theo.saemix,tau=24)
saemix.data2<-saemixData(name.data=theo.saemix2,header=TRUE,sep=" ",na=NA,
  name.group=c("Id"),name.predictors=c("Dose","Time","tau"),
  name.response=c("Concentration"),name.covariates=c("Weight","Sex"),
  units=list(x="hr",y="mg/L",covariates=c("kg","-")), name.X="Time")
# Define the model for steady-state
modelSS<-function(psi,id,xidep) {
  dose<-xidep[,1]
  tim<-xidep[,2]
  tau<-xidep[,3]
  ka<-psi[id,1]
  V<-psi[id,2]
  CL<-psi[id,3]
  k<-CL/V
  ypred<-dose*ka/(V*(ka-k))*(exp(-k*tim)/(1-exp(-k*tau))-
exp(-ka*tim)/(1-exp(-ka*tau)))
  return(ypred)
}
saemix.model2<-saemixModel(model=modelSS,
  description="One-compartment model with first-order absorption, Steady-state",
  psi0=matrix(c(1.,20,0.5,0.1,0,-0.01),ncol=3,byrow=TRUE,
  dimnames=list(NULL, c("ka","V","CL"))),transform.par=c(1,1,1))

# Run SAEMIX again
saemix.options<-list(seed=632545)
saemix.fit2<-saemix(saemix.model2,saemix.data2,saemix.options)
```

4.2 Simulated pharmacodynamic model

A symposium dedicated to Comparison of Algorithms Using Simulated Data Sets and Blind Analysis, took place in Lyon, France, September 2004, organised by P. Girard and F. Mentré. During this symposium, a blind comparison of several PK/PD modelling software was performed, using simulated datasets. This example uses two datasets simulated for this comparison.

The two datasets contain 100 individuals, each receiving 3 different doses: (0, 10, 90), (5, 25, 65) or (0, 20, 30). It is assumed that doses were given in a cross-over study with sufficient wash-out period to avoid a carry-over effect. Responses y_{ij} have been simulated with an E_{\max} model, a standard pharmacodynamic model:

$$y_{ij} = E_{0,i} \frac{D_{ij} E_{\max,i}}{D_{ij} + ED_{50,i}} + \epsilon_{ij} \quad (4.3)$$

For subject i :

- the regression variable is the dose received $x_{ij} = (D_{ij})$
- the vector of individual parameters is $\theta_i = (\ln(E_{0,i}), \ln(E_{\max,i}), \ln(ED_{50,i}))$
- the only available covariate is the gender w_i of the individual (0 for male and 1 for female)

The individual parameters were simulated assuming a log-normal distribution for all parameters, and a gender effect on $ED_{50,i}$:

$$\begin{aligned} \ln(E_{0,i}) &= \ln(E_0) + \eta_{i1} \\ \ln(E_{\max,i}) &= \ln(E_{\max}) + \eta_{i2} \\ \ln(ED_{50,i}) &= \ln(ED_{50}) + \beta w_i + \eta_{i3} \end{aligned} \quad (4.4)$$

In the simulations, the fixed effects were set to $(\ln(E_0), \ln(E_{\max}), \ln(ED_{50})) = (24, 100, 12)$. The covariance matrix of the random effects was a diagonal matrix. The variances of the random effects were (0.12, 0.26, 0.05). The residual variance was a constant variance, with $\sigma^2 = 20$. The two data sets were simulated with different values of β :

- the first dataset was simulated with a gender effect, $\beta = 0.3$, and is available in the package under the name PD1.saem
- the second dataset was simulated under the null hypothesis, $\beta = 0$, and is available in the package under the name PD2.saem

The data is shown in figure 4.9.

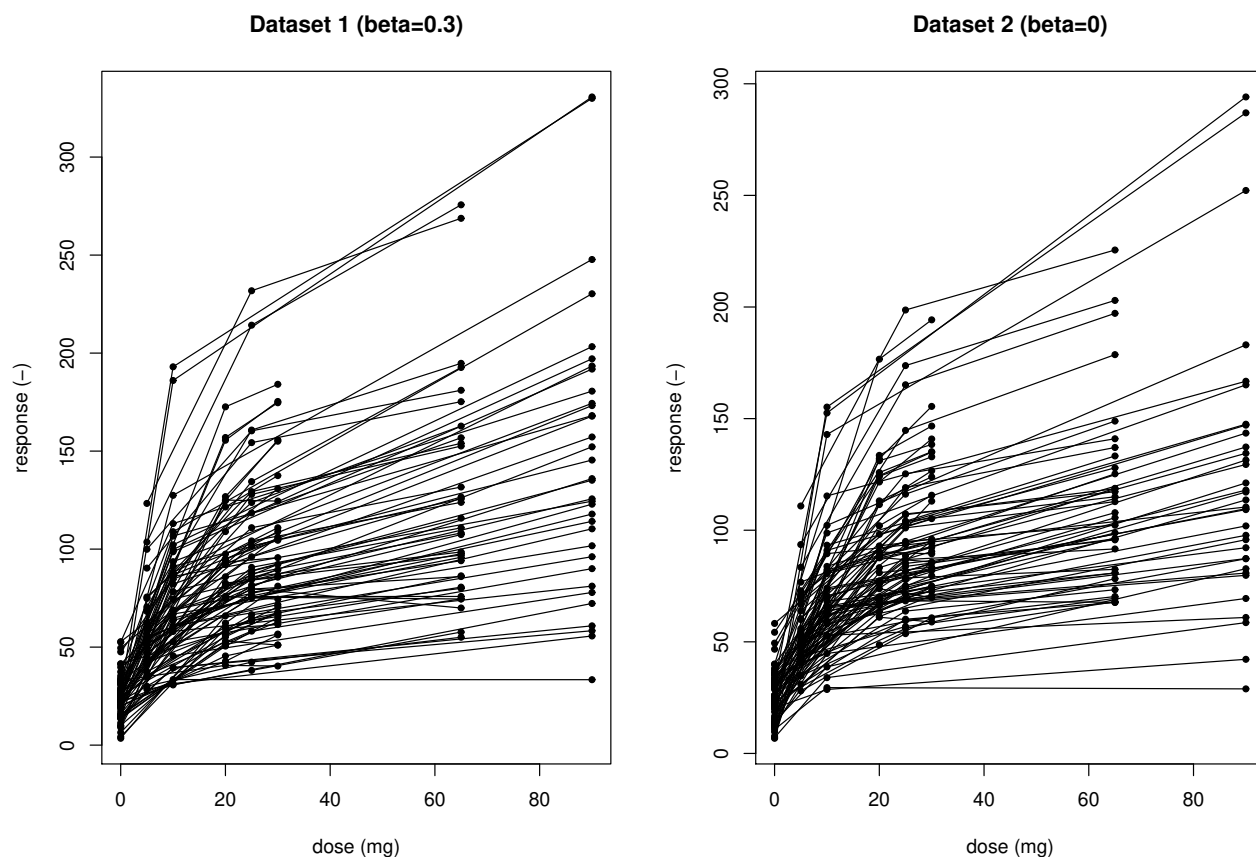


Figure 4.9: Effect versus dose for the data simulated with an E_{\max} model, with a gender effect on ED_{50} (left) and without a gender effect (right).

```
library(saemix)
data(PD1.saemix)
data(PD2.saemix)
saemix.data1<-saemixData(name.data=PD1.saemix,header=TRUE,name.group=c("subject"),
name.predictors=c("dose"),name.response=c("response"),name.covariates=c("gender"),
units=list(x="mg",y="-",covariates="-"))

saemix.data2<-saemixData(name.data=PD2.saemix,header=TRUE,name.group=c("subject"),
name.predictors=c("dose"),name.response=c("response"),name.covariates=c("gender"),
units=list(x="mg",y="-",covariates="-"))

modelemax<-function(psi,id,xidep) {
# input:
#  psi : matrix of parameters (3 columns, E0, Emax, EC50)
#  id  : vector of indices
```

```
# xidep : dependent variables (same nb of rows as length of id)
# returns:
# a vector of predictions of length equal to length of id
dose<-xidep[,1]
e0<-psi[id,1]
emax<-psi[id,2]
e50<-psi[id,3]
f<-e0+emax*dose/(e50+dose)
return(f)
}
saemix.model<-saemixModel(model=modelemax,description="Emax model",
  psi0=matrix(c(20,300,20,0,0,0),ncol=3,byrow=TRUE,
  dimnames=list(NULL,c("E0","Emax","EC50"))),transform.par=c(1,1,1),
  covariate.model=matrix(c(0,0,1),ncol=3,byrow=TRUE),
  fixed.estim=c(1,1,1), error.model="constant",
  covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE))
```

We can check the predictions for the model using the plot function with the model and data objects as arguments (predictions will be obtained for the parameters in the psi0 argument of the model), as well as change the parameters on the fly using the psi=c(XXX) argument. If additional predictors are needed by the model, we would add them using predictors=c(XXX). The ilist argument is used to specify which subjects to plot (by default, the first 20 subjects are used):

```
# Predictions for psi0
plot(saemix.model, saemix.data)

# Predictions for psi=c(0, 200, 50)
plot(saemix.model, saemix.data, psi=c(0, 200, 50), ilist=1:12)
```

Note: when applied directly, the plot function requires the first predictor (here, the only one, dose) in the model to be the one on the X-axis, if there are several predictors in the model (see the online help example for how to permute the two predictors in the theophylline example to use this function). The plot function can also be applied after a model fit to the two elements (model and data) in an SaemixObject object, in which case the mapping of the predictors will be automatic and predictions can be obtained for the population or individual parameters.

The following code was used in R to run this example on the two datasets:

```
saemix.options<-list(directory=directory,algorithms=c(1,1,1),nb.chains=1,
  save=FALSE,save.graphs=FALSE)
```

```
# Fitting the model on the two PD datasets
saemix.fit1<-saemix(saemix.model,saemix.data1,saemix.options)
saemix.fit2<-saemix(saemix.model,saemix.data2,saemix.options)
```

Table 4.2 shows the parameter estimates for the two datasets. The estimates for the three fixed effects are similar for both datasets, while the estimate of β is close to the values simulated for both. For PD2.saemix, the SE on β is very large, as is the SE on the estimate of the variability of EC_{50} .

Parameter	PD1.saemix		PD2.saemix	
	Estimate (SE%)	IIV (SE%)	Estimate (SE%)	IIV (SE%)
E_0 (-)	22.71 (5%)	0.13 (22%)	23.18 (5%)	0.16 (20%)
E_{\max} (-)	106.46 (6%)	0.31 (15%)	96.14 (5%)	0.22 (15%)
EC_{50} (mg)	11.25 (8%)	0.03 (55%)	12.38 (6%)	0.01 (151%)
$\beta_{gender,EC_{50}}$ (-)	0.35 (26%)	-	-0.06 (116%)	-
a (mg.L ⁻¹)	4.94 (8%)	-	4.67 (8%)	-

Table 4.2: Pharmacokinetic parameters estimated by `saemix` for the simulated PD data.

The convergence plots are shown in figures 4.10 and 4.11, where we can see β converging to a non-zero value for the first dataset, while the estimate fluctuates around 0 for the second dataset. The Wald test performed for the fixed effect representing the effect of gender on EC_{50} shows that this parameter is significantly different from 0 in the first dataset ($p=6.10^{-5}$).

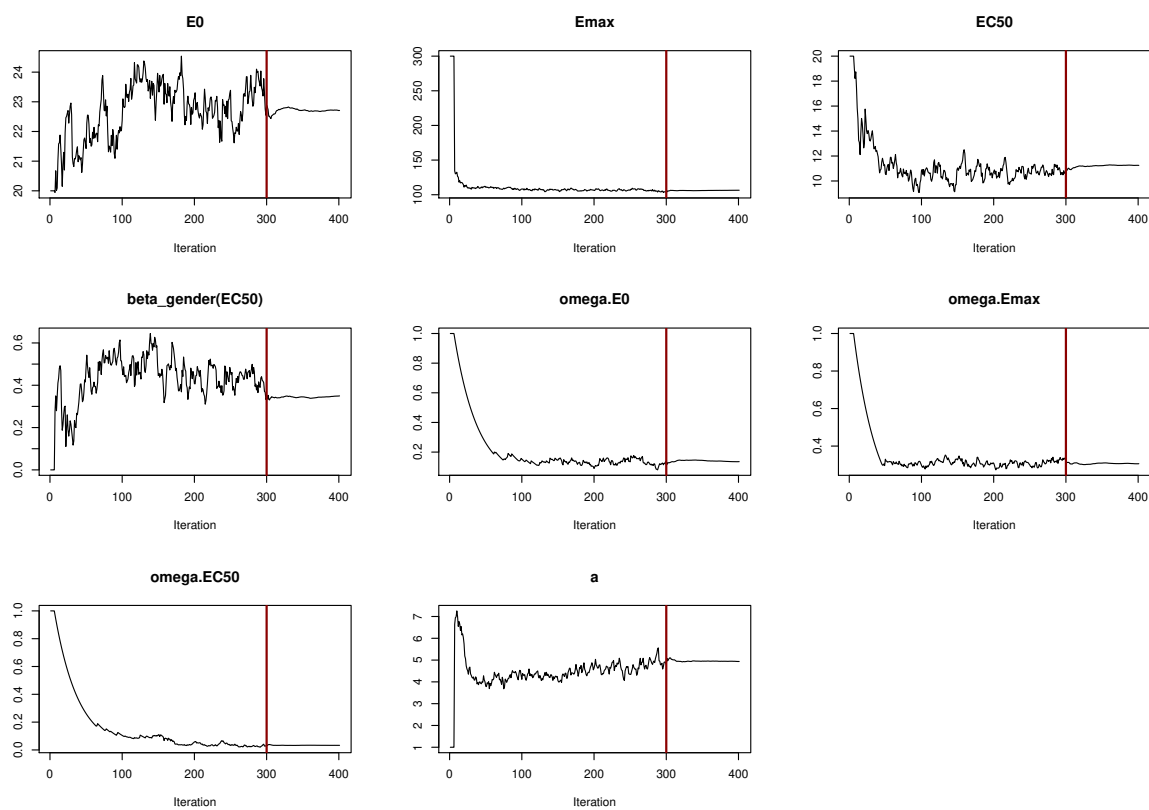


Figure 4.10: Convergence plots for the estimated pharmacokinetic parameters and the variabilities, for the first dataset.

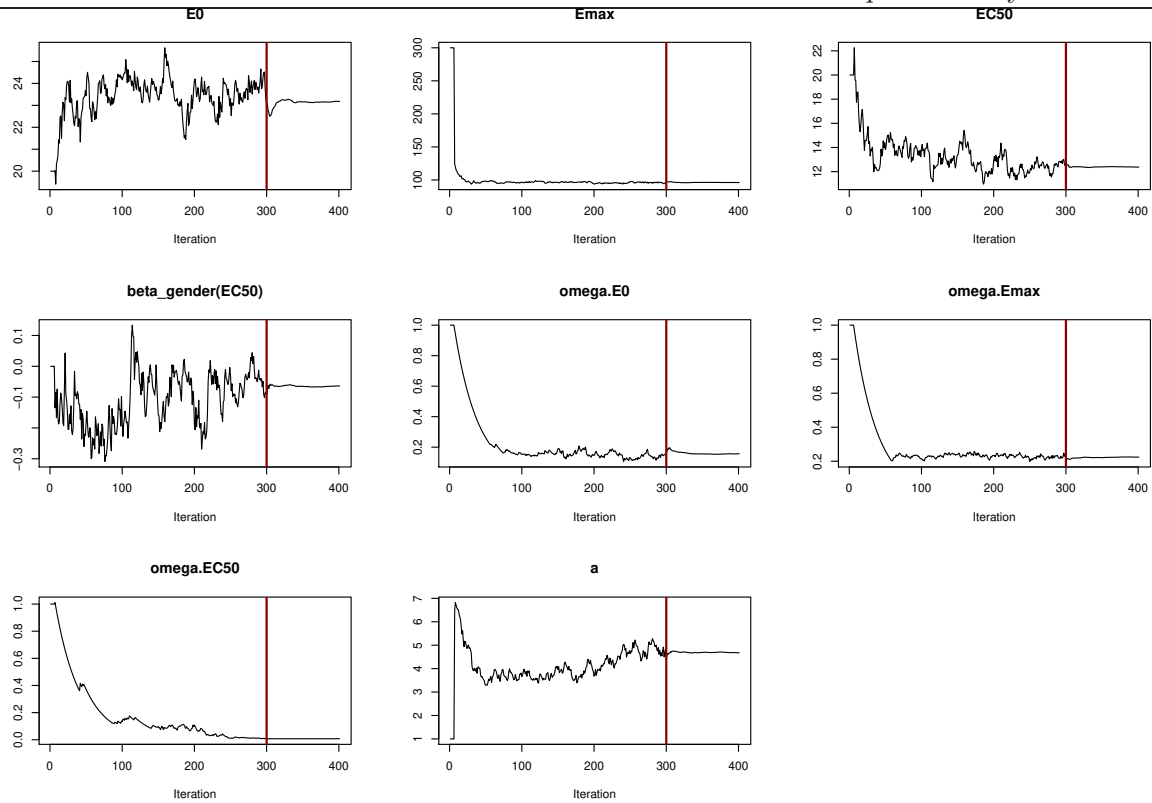


Figure 4.11: Convergence plots for the estimated pharmacokinetic parameters and the variabilities, for the second dataset.

Finally, figure 4.12 shows the individual data for the first 12 subjects in the first dataset, with the individual predictions overlayed. A smoothed prediction was obtained. The model fits the data extremely well, which is unsurprising given that this is simulated data, with a rather small residual variability.

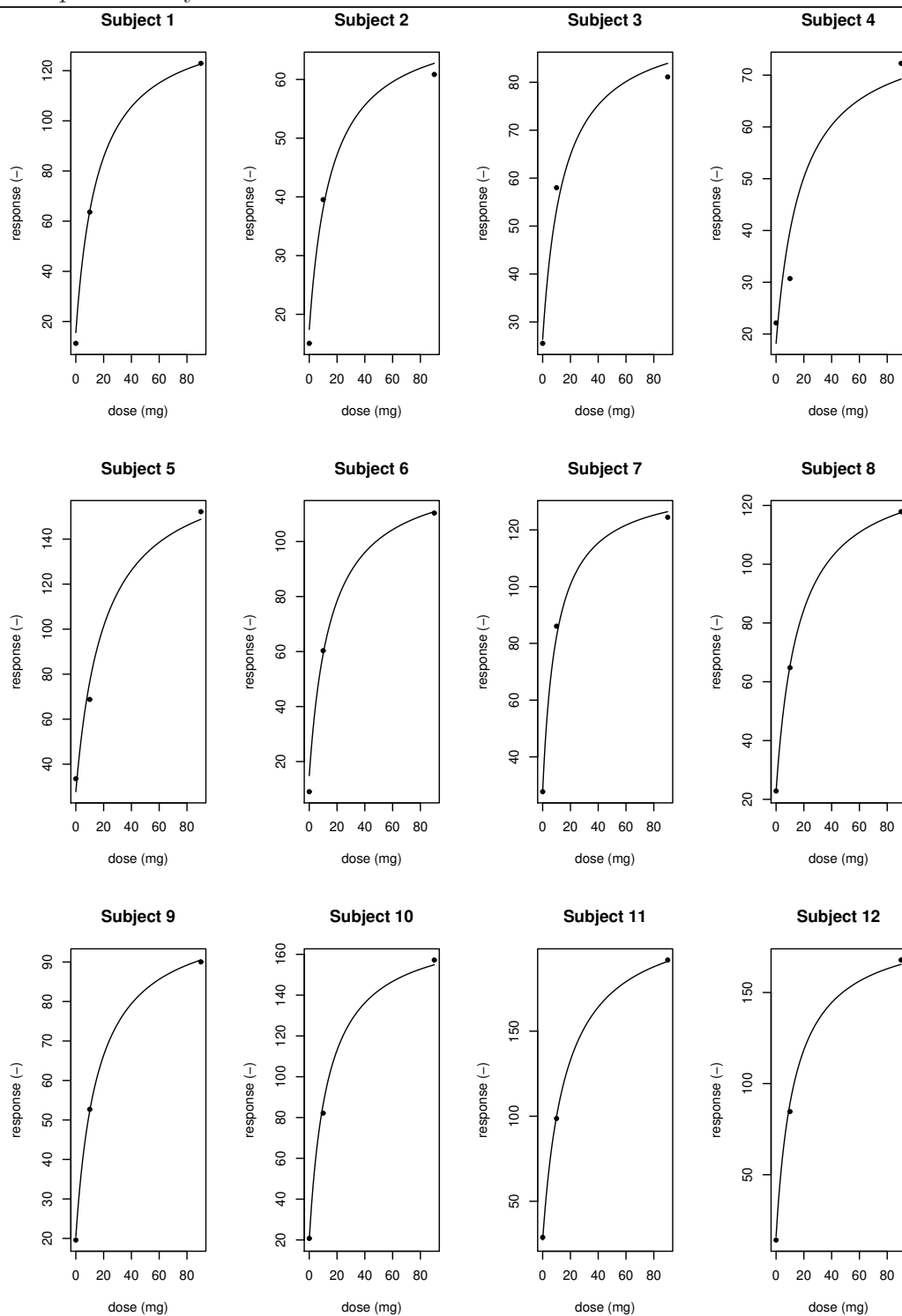


Figure 4.12: Individual plots for the 12 subjects in the first dataset (PD1.saemix). Dots represent observations and the line shows the profile predicted using the individual estimated parameters.

4.3 Weight gain of cows

The data used in this example is the evolution of the weight (in kg) of 560 cows. The weight of each cow was recorded on 9 or 10 occasions. An exponential model was assumed to describe the weight gain with time:

$$y_{ij} = A_i (1 - B_i e^{-K_i t_{ij}}) + \epsilon_{ij} \quad (4.5)$$

For subject i :

- the regression variable is the time (in days) $x_{ij} = (t_{ij})$
- the vector of individual parameters is $\theta_i = (A_i, B_i, K_i)$
- there were 3 covariates in the file:
 1. the year of birth (between 1988 and 1998)
 2. existence of a twin (no=1, yes=2)
 3. the rank of birth (between 3 and 7)

The data is shown in figure 4.13.

The following code was used in R to run this example:

```
library(saemix)
data(cow.saemix)
saemix.data<-saemixData(name.data=cow.saemix,header=TRUE,name.group=c("cow"),
name.predictors=c("time"),name.response=c("weight"),
name.covariates=c("birthyear","twin","birthrank"),
units=list(x="days",y="kg",covariates=c("yr","-","-")))

growthcow<-function(psi,id,xidep) {
# input:
#   psi : matrix of parameters (3 columns, ka, V, CL)
#   id : vector of indices
#   xidep : dependent variables (same nb of rows as length of id)
# returns:
#   a vector of predictions of length equal to length of id
x<-xidep[,1]
a<-psi[id,1]
b<-psi[id,2]
k<-psi[id,3]
```

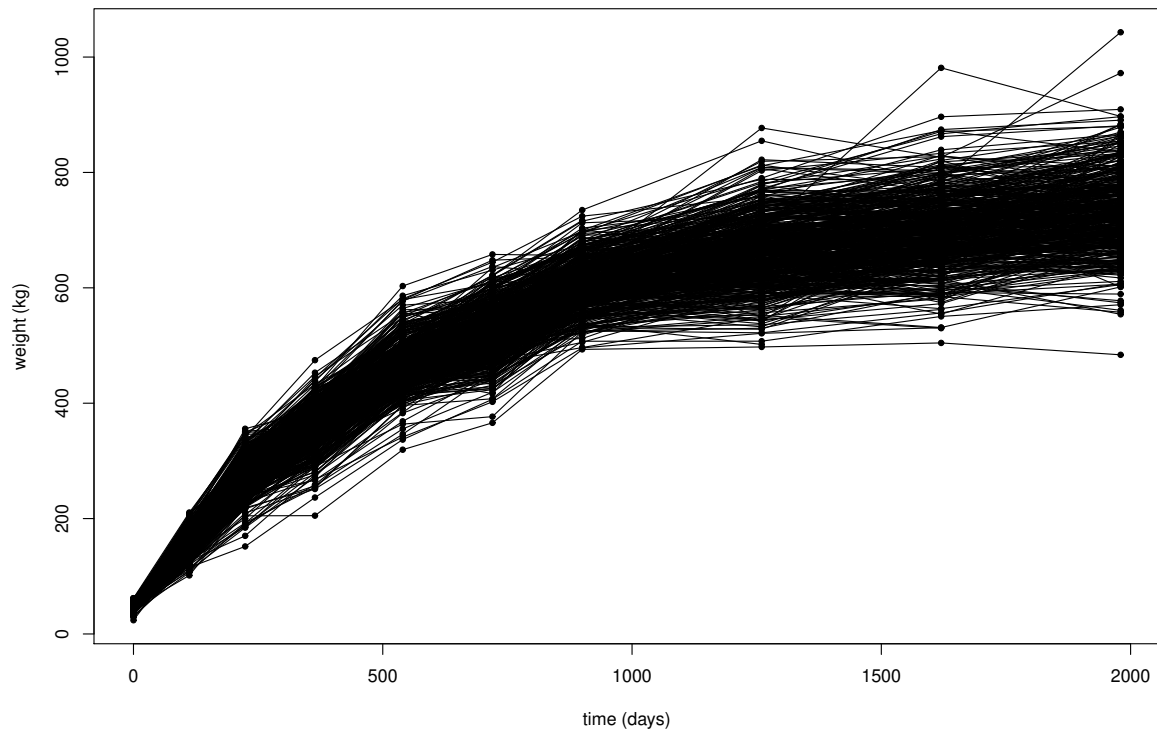


Figure 4.13: Weight gain of 560 cows recorded repeatedly over time.

```
f<-a*(1-b*exp(-k*x))
return(f)
}
saemix.model<-saemixModel(model=growthcow,description="Exponential model",
psi0=matrix(c(700,0.9,0.02,0,0,0),ncol=3,byrow=TRUE,
dimnames=list(NULL,c("A","B","k"))),transform.par=c(1,1,1),fixed.estim=c(1,1,1),
covariate.model=matrix(c(0,0,0,0,0,0,0,0,0),ncol=3,byrow=TRUE),
covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),
omega.init=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=TRUE),error.model="constant")

saemix.options<-list(algorithms=c(1,1,1),nbiter.saemix=c(200,100),nb.chains=1,
save=FALSE,save.graphs=FALSE)

# Fitting the models
saemix.fit<-saemix(saemix.model,saemix.data,saemix.options)
```

As an alternative, we can compute the estimate of the likelihood by Gaussian Quadrature:

```
saemix.fit<-llgq.saemix(saemix.fit)
```

The three estimates of the likelihood were found to be in good agreement in this example:

```
-----
----- Statistical criteria -----
-----
Likelihood computed by linearisation
      -2LL= 53723.42
      AIC = 53737.42
      BIC = 53767.71

Likelihood computed by importance sampling
      -2LL= 53723.88
      AIC = 53737.88
      BIC = 53768.18

Likelihood computed by Gaussian quadrature
      -2LL= 53723.04
      AIC = 53737.04
      BIC = 53767.34
-----
```

The fits to the data from the first 4 animals can be plotted using the function `saemix.plot.fits`. First, default plot options are set in a list called `saemix.plot.options` using the function `saemix.plot.setoptions`. Second, the option controlling the list of subjects to be plotted is set (here, we choose to plot the graphs for the first four animals), and the option `smooth` indicates that we want an smoothed version of the plots (using interpolated weights):

```
plot(saemix.fit,plot.type="individual.fit",ilist=1:4,smooth=TRUE)
```

The result is shown in figure [4.14](#).

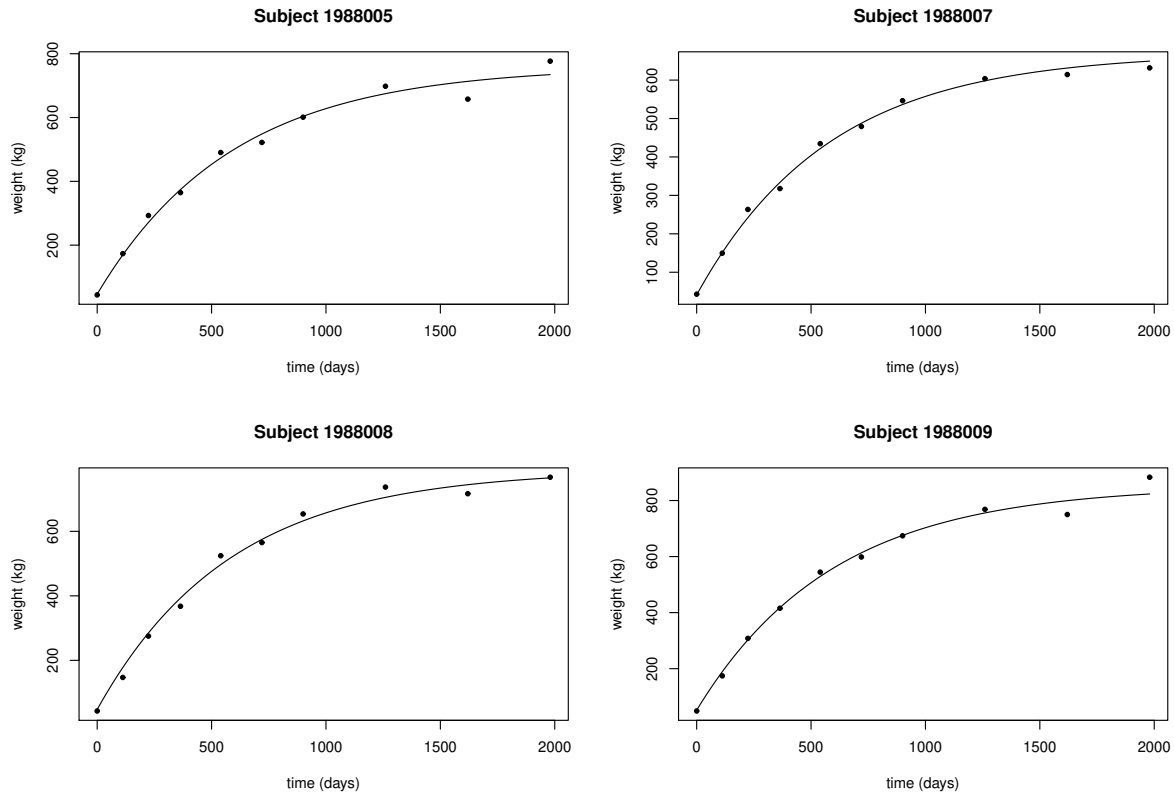


Figure 4.14: Fit for the first four subjects in the cow dataset.

4.4 Height of Oxford boys

saemix can be used even for linear models. The dataset `oxboys.saemix` was taken from the library `nlme` [31]. It describes the evolution with age of the height of boys from Oxford, England. There is no covariate in the model, and we use a simple linear model to account for the increase in height over this age range:

$$y_{ij} = \text{Base}_i + \text{Slope}_i \text{age}_{ij} + \epsilon_{ij} \quad (4.6)$$

where Base_i is the baseline height at the entrance of subject i in the study and Slope_i the slope for the increase of height with age age_{ij} . For subject i :

- the vector of regression (or design) variables is $x_{ij} = (\text{age}_{ij})$
- the vector of individual parameters is $\theta_i = (\text{Base}_i, \text{Slope}_i)$
 - the individual parameters are assumed to have a normal distribution

- we can use a simple homoscedastic error model where $\text{Var}(\epsilon_{ij}) = a^2$

The data is shown in figure 4.15.

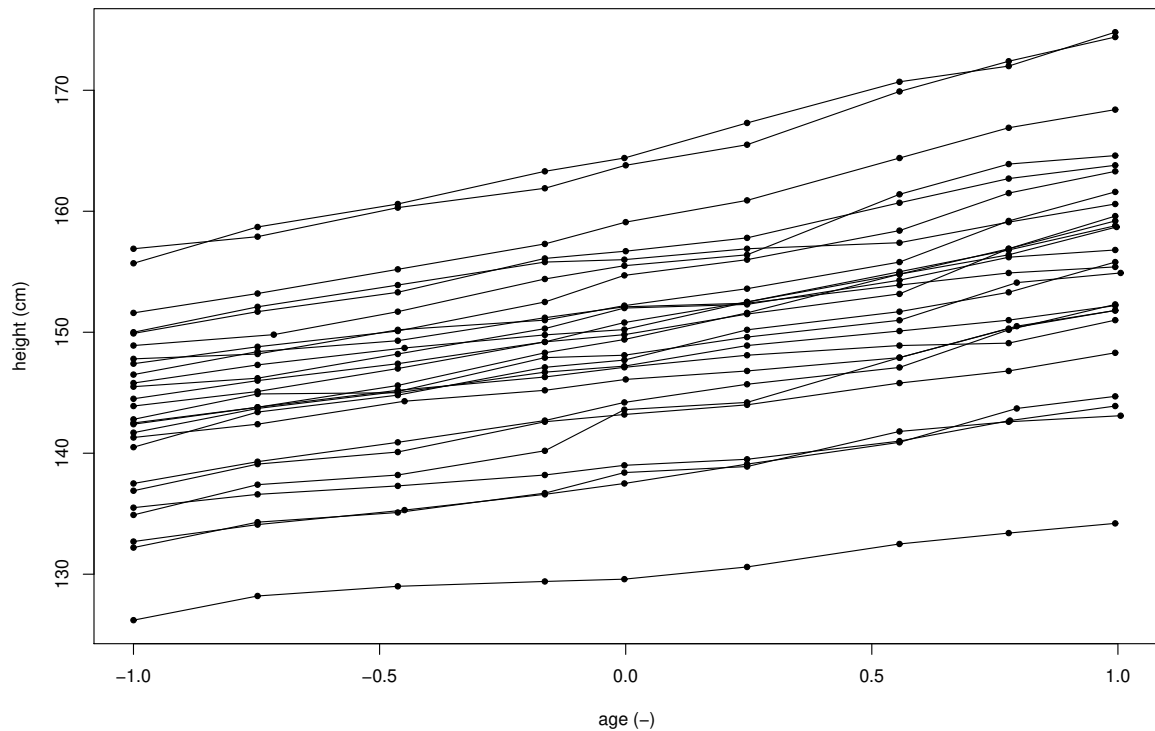


Figure 4.15: Evolution with age of the height of boys from Oxford.

The following code was used in R to run this example:

```
library(saemix)

data(oxboys.saemix)
saemix.data<-saemixData(name.data=oxboys.saemix,header=T,name.group=c("Subject"),
name.predictors=c("age"),name.response=c("height"), units=list(x="-",y="yr"))

growth.linear<-function(psi,id,xidep) {
# input:
#   psi : matrix of parameters (2 columns, base and slope)
#   id  : vector of indices
#   xidep : dependent variables (same nb of rows as length of id)
# returns:
#   a vector of predictions of length equal to length of id
```

```

x<-xidep[,1]
base<-psi[id,1]
slope<-psi[id,2]
f<-base+slope*x
return(f)
}
saemix.model<-saemixModel(model=growth.linear,description="Linear model",
psi0=matrix(c(140,1),ncol=2,byrow=T,dimnames=list(NULL,c("base","slope"))),
transform.par=c(1,0), covariance.model=matrix(c(1,1,1,1),ncol=2,byrow=T),
error.model="constant")

saemix.options<-list(algorithms=c(1,1,1),nb.chains=1)

saemix.fit<-saemix(saemix.model,saemix.data,saemix.options)

```

4.5 A yield model

The data used in this study were from 37 winter wheat experiments carried out between 1990 and 1996 on commercial farms in the Paris Basin, France. Each experiment was from a different site. Two soil types were represented, a loam soil and a chalky soil. Common winter wheat varieties were used. Each experiment consisted of five to eight different nitrogen fertilizer rates, for a total of 224 nitrogen treatments. Nitrogen fertilizer was applied in two applications during the growing season. For each nitrogen treatment, grain yield (adjusted to 150 g.kg⁻¹ grain moisture content) was measured. In addition, end-of-winter mineral soil nitrogen (NO₃- plus NH₄⁺) in the 0- to 90-cm layer was measured on each site-year during February when the crops were tillering. See [9] for a more complete description of the plant sampling and nitrogen analysis. Yield and end-of-winter mineral soil nitrogen measurements were in the ranges 3.44- 11.54 t.ha⁻¹ , and 40-180 kg.ha⁻¹ respectively.

The data is shown in figure 4.16.

Let y_{ij} denote the j^{th} measurement of the yield response in the i^{th} site-year when the nitrogen fertilizer dose d_{ij} is applied. The only available covariate is the amount of soil mineral nitrogen at the end of winter (w_i).

A first model is a linear-plus-plateau function (LP) defined by:

$$y_{ij} = \begin{cases} Y_{max,i} + B_i(d_{ij} - X_{max,i}) & \text{if } d_{ij} \leq X_{max,i} \\ Y_{max,i} & \text{if } d_{ij} \geq X_{max,i} \end{cases} \quad (4.7)$$

This model includes three individual random parameters, $\phi_i = (Y_{max,i}, X_{max,i}, B_i)$. $Y_{max,i}$ is the maximal yield value in the i^{th} site-year and $X_{max,i}$ is the fertilizer dose that maximizes yield. The three parameters were assumed to follow a normal distribution.

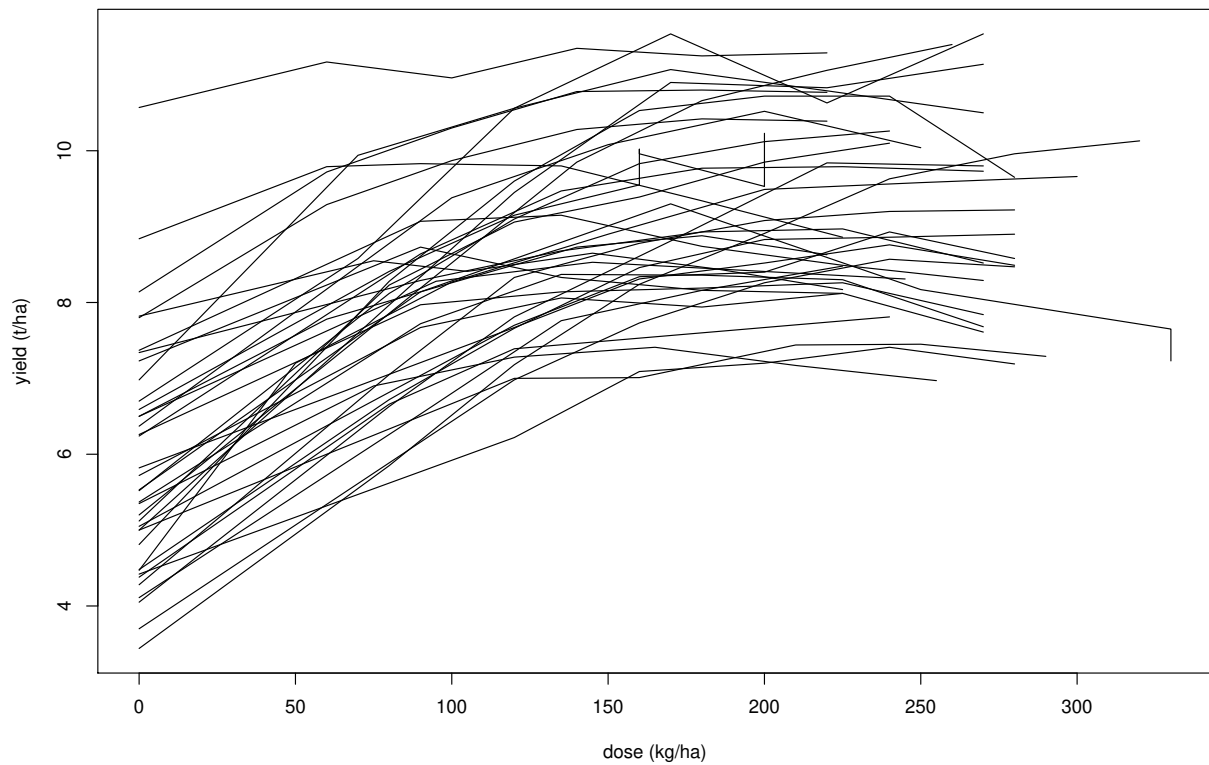


Figure 4.16: Yield from 37 winter wheat experiments.

A second model is a square-root-plus-plateau function (QP) defined by:

$$y_{ij} = \begin{cases} Y_{max,i} + B_i(\sqrt{d_{ij}} - \sqrt{X_{max,i}}) & \text{if } d_{ij} \leq X_{max,i} \\ Y_{max,i} & \text{if } d_{ij} \geq X_{max,i} \end{cases} \quad (4.8)$$

We use the following code to run these two models:

```
library(saemix)

data(yield.saemix)
saemix.data<-saemixData(name.data=yield.saemix,header=TRUE,name.group=c("site"),
name.predictors=c("dose"),name.response=c("yield"), name.covariates=c("soil.nitrogen"),
units=list(x="kg/ha",y="t/ha", covariates=c("kg/ha")))

yield.LP<-function(psi,id,xidep) {
  x<-xidep[,1]
  ymax<-psi[id,1]
```

```

    xmax<-psi[id,2]
    slope<-psi[id,3]
    f<-ymax+slope*(x-xmax)
#   cat(length(f),"  ",length(ymax),"\n")
    f[x>xmax]<-ymax[x>xmax]
    return(f)
}

yield.QP<-function(psi,id,xidep) {
  x<-xidep[,1]
  ymax<-psi[id,1]
  xmax<-psi[id,2]
  slope<-psi[id,3]
  f<-ymax+slope*(x**0.5-xmax**0.5)
#   f<-ymax+slope*sqrt(abs(x-xmax))
  f[x>xmax]<-ymax[x>xmax]
  return(f)
}

saemix.model1<-saemixModel(model=yield.LP,description="Linear + plateau model",
psi0=matrix(c(8,100,0.2,0,0,0),ncol=3,byrow=T, dimnames=list(NULL,c("Ymax","Xmax",
"slope"))), covariate.model=matrix(c(0,0,0),ncol=3,byrow=T),
transform.par=c(0,0,0),covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=T),
error.model="constant")

saemix.model2<-saemixModel(model=yield.QP,description="Quadratic + plateau model",
psi0=matrix(c(10,120,0.005,0,0,0),ncol=3,byrow=T, dimnames=list(NULL,c("Ymax","Xmax",
"slope"))), covariate.model=matrix(c(0,0,0),ncol=3,byrow=T), transform.par=c(0,0,0),
covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=T),error.model="constant")

saemix.options<-list(algorithms=c(1,1,1),nb.chains=1, nbiter.saemix=c(400,100),
nmc.is=25000, save=FALSE,save.graphs=FALSE)

# Fitting the models
saemix.fit1<-saemix(saemix.model1,saemix.data,saemix.options)
saemix.fit2<-saemix(saemix.model2,saemix.data,saemix.options)

```

The two models perform very similarly in terms of log-likelihood, with a slight advantage to the LP model: the statistical criterion (-2 times the log-likelihood) was equal to 406.86 for the LP model and to 416.28 for the QP model. Figure 4.17 shows the plots of predictions versus observations for the two models, again very similar.

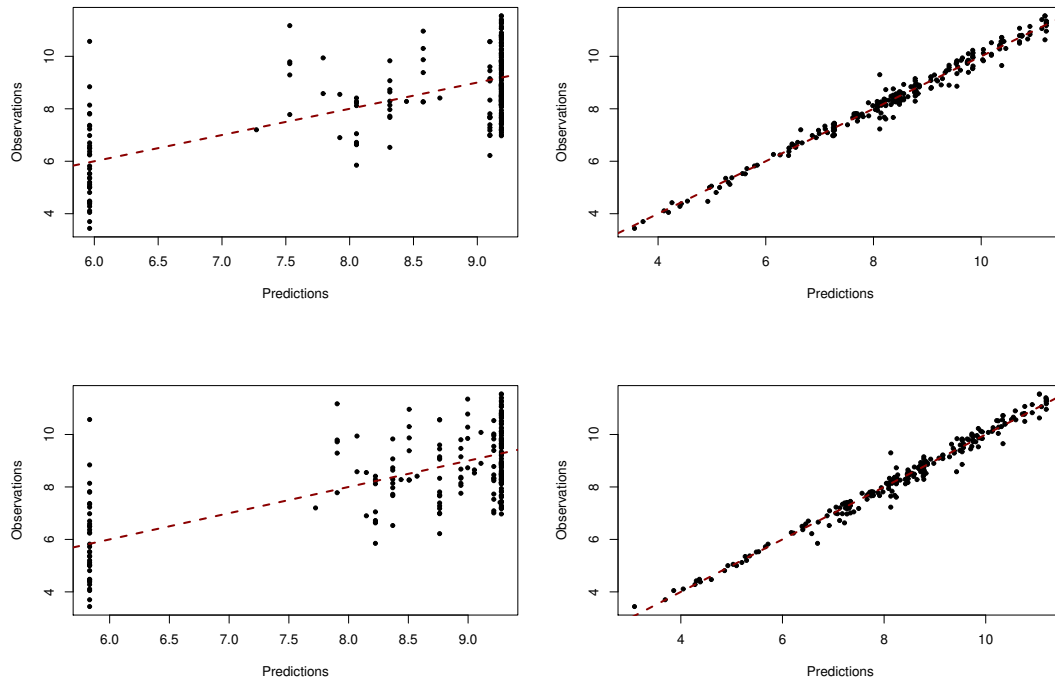


Figure 4.17: Observations versus predictions for the LP model (upper panel) and QP model (lower panel), with population predictions on the left and individual predictions on the right.

Figure 4.18 shows the fit of the two models for the first four subjects. The figure was obtained using the following code:

```
par(mfrow=c(4,2))
for(i in 1:4) {
  plot(saemix.fit1,plot.type="individual.fit",ilist=i,smooth=TRUE,new=F)
  plot(saemix.fit2,plot.type="individual.fit",ilist=i,smooth=TRUE,new=F)
}
```

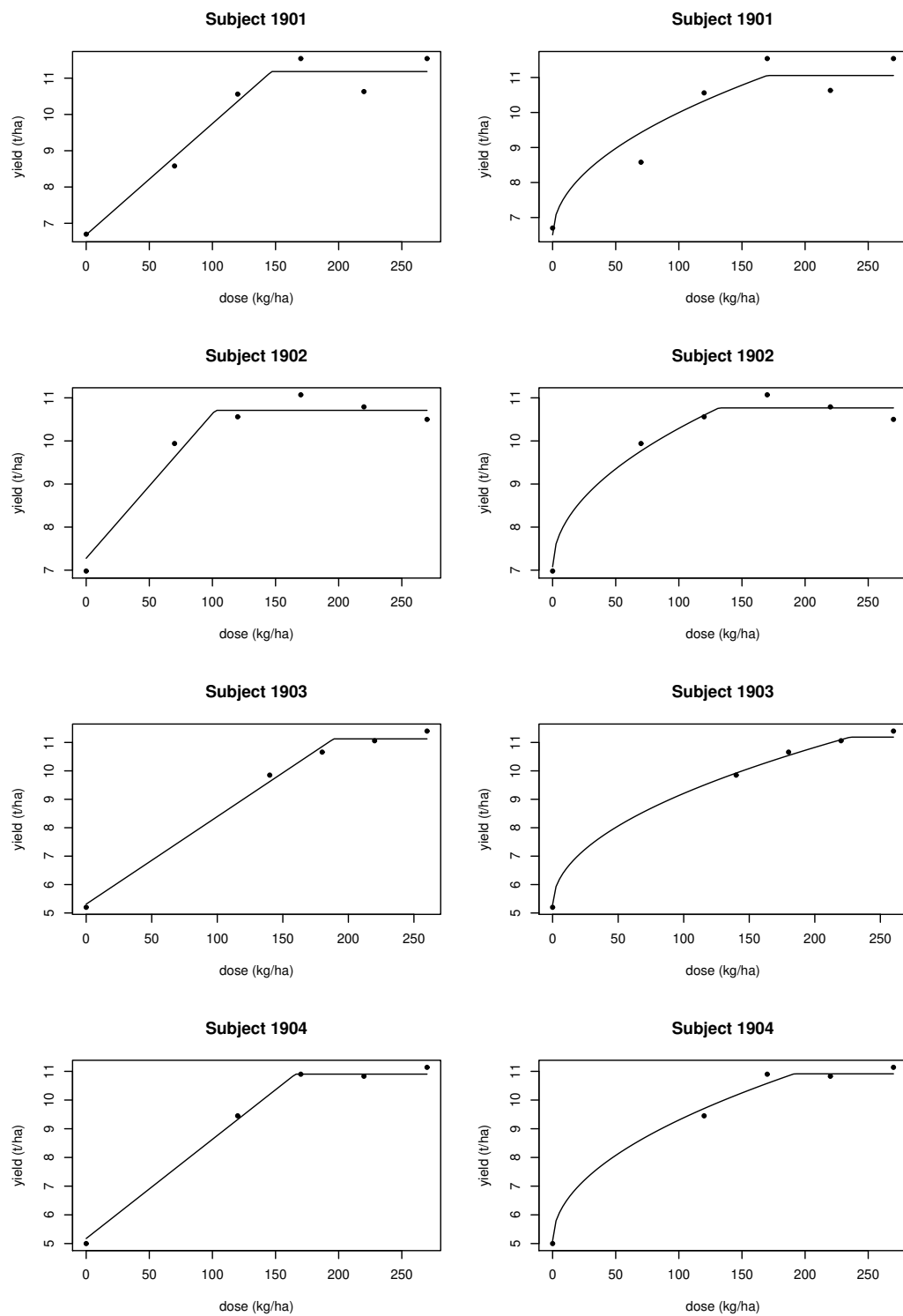


Figure 4.18: Fits for the LP model (left) and QP model (right) for the first 4 subjects.

We can explore the covariates using diagnostic plots. For instance, the following code plots the estimated individual parameters versus the covariates in the model (here, soil nitrogen), assuming the fit is in the object `saemix.fit`:

```
plot(saemix.fit1, plot.type="parameters.vs.covariates")
```

Figure 4.19 shows the result, and indicates a decreasing trend in X_{max} with increasing amounts of soil nitrogen.

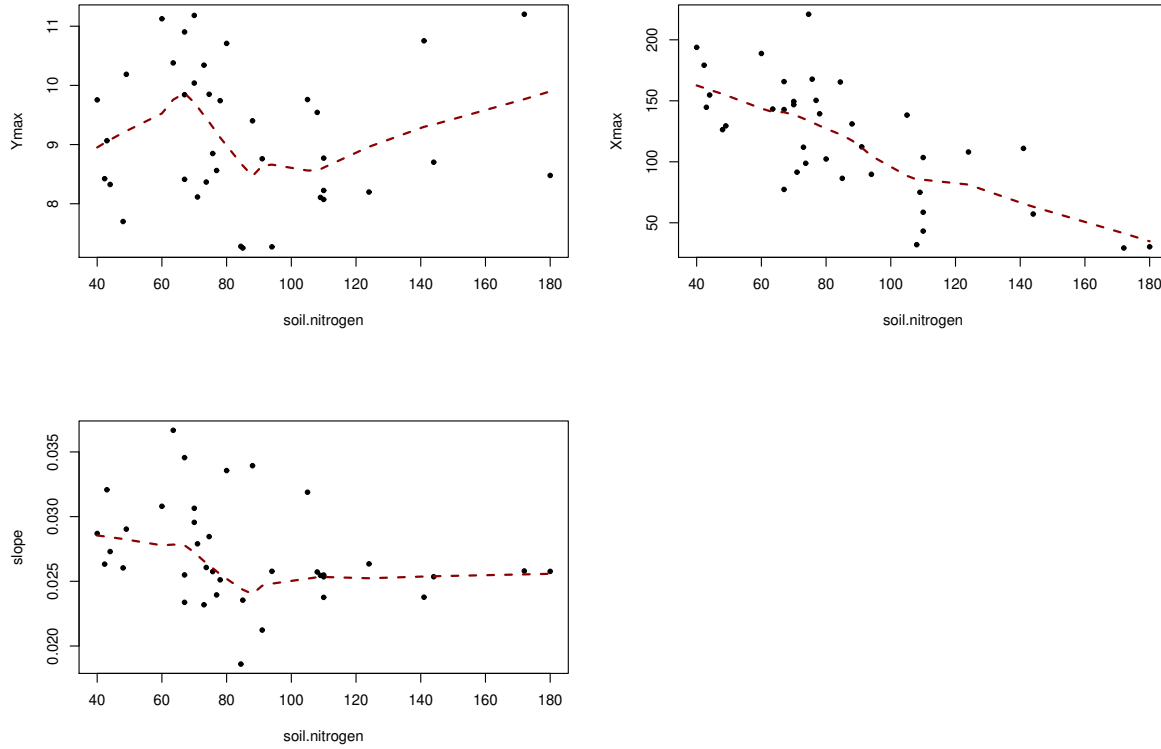


Figure 4.19: Graphs of the estimated (individual) parameters versus covariate.

In this example, we can then show with `saemix` that the effect of the amount of soil mineral nitrogen at the end of winter is statistically significant for explaining the fluctuations of the parameter X_{max} in both LP and QP models, with a drop of over 20 points in the statistical criterion. For example, the following code shows how to fit the LP-model with and without covariate effect on X_{max} , and outputs the resulting log-likelihoods:

```
saemix.model3<-saemixModel(model=yield.LP,description="Linear + plateau model",
psi0=matrix(c(8,100,0.2,0,0,0),ncol=3,byrow=T, dimnames=list(NULL,c("Ymax","Xmax",
```

```

"slope"))), covariate.model=matrix(c(0,1,0),ncol=3,byrow=T),
transform.par=c(0,0,0),covariance.model=matrix(c(1,0,0,0,1,0,0,0,1),ncol=3,byrow=T),
error.model="constant")

saemix.fit3<-saemix(saemix.model3,saemix.data,saemix.options)

{
  cat("LP model:\n")
  cat("    without covariate, -2xLL=", (saemix.fit1["results"] ["ll.is"])*(-2), "\n")
  cat("    with covariate, -2xLL=", (saemix.fit3["results"] ["ll.is"])*(-2), "\n")
}

# The same can be done for the QP model

```

4.6 Discrete data

4.6.1 Binary data

saemix 3.0 can now be used to estimate the parameters of a model where the outcome is a discrete response. The most simple of this, but also the least informative type, is a binary response that can take the values 0 or 1. In this section, we illustrate the use of saemix to model binary data from a randomised clinical trial comparing two treatments for fungal toenail infection. We use the toenail dataset available in R in the packages prLogistic or HSAUR3.

Data are from [8], a multi-center randomised comparison of two oral treatments (A and B) for toenail infection. 294 patients are measured at seven visits, i.e. at baseline (week 0), and at weeks 4, 8, 12, 24, 36, and 48 thereafter, comprising a total of 1908 measurements. The primary end point was the absence of toenail infection and the outcome of interest is the binary variable "onycholysis" which indicates the degree of separation of the nail plate from the nail-bed (categorised as 0=none or mild versus 1=moderate or severe). Figure 4.20 shows the evolution of the number of events (left) and the proportion of events (right) in the two treatment groups over the 7 visits of the study.

Several analyses have been made in the literature [24, 25], and here we fit the logistic random effect model developed by [16]. This model includes a random intercept (θ_1 , normally distributed with a standard deviation ω_1), a time effect (β_2 , normally distributed with a standard deviation ω_2). Treatment (A or B) (β) is included as a covariate on time. The treatments were randomised at baseline so we don't include a treatment effect on the intercept. The probability $p_{ij} = P(Y_{ij} = 1 | \theta_{1,i}, \theta_{2,i})$ associated with an event Y_{ij} at time t_{ij} is given by the following equation for the logit:

$$\text{logit}(p_{ij}) = \ln \left(\frac{p_{ij}}{1 - p_{ij}} \right) = \theta_{1,i} + \theta_{1,i} t_{ij} \quad (4.9)$$

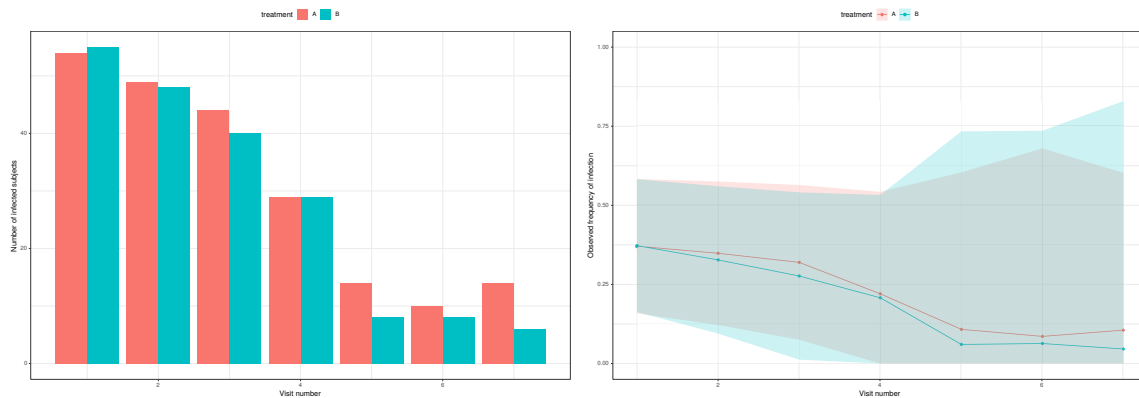


Figure 4.20: Toenail data. Left: number of events at each visit; right: proportion of infected subjects at each visit.

For non-gaussian models, the model function must be written to return the log-pdf, that is, the logarithm of the probability of the observed response given a set of parameters. To do this we need to pass the response as one of the predictors.

```
data(toenail.saemix)
saemix.data<-saemixData(name.data=toe,name.group=c("id"),name.predictors=c("time","y"),
  name.response="y", name.covariates=c("treatment"),name.X=c("time"))
```

To tell saemix that we are now dealing with non-continuous responses, we add the argument `modelType='likelihood'` to the definition of the model using the function `saemixModel`. We assume only the intercept has interindividual variability, and follows a normal distribution. We set the covariate model for a treatment effect on θ_2 .

```
binary.model<-function(psi,id,xidep) {
  tim<-xidep[,1]
  y<-xidep[,2]
  inter<-psi[id,1]
  slope<-psi[id,2]
  logit<-inter+slope*tim
  pevent<-exp(logit)/(1+exp(logit))
  pobs = (y==0)*(1-pevent)+(y==1)*pevent
  logpdf <- log(pobs)
  return(logpdf)
}
```

```
saemix.model<-saemixModel(model=binary.model,description="Binary model",
```

```

modeltype="likelihood",
psi0=matrix(c(-5,-.1,0,0),ncol=2,byrow=TRUE,dimnames=list(NULL,
c("theta1","theta2"))),
transform.par=c(0,0), covariate.model=c(0,1),
covariance.model=matrix(c(1,0,0,1),ncol=2))

```

We then fit the model, setting the option `fim=FALSE` as the approximation used in the computation of the FIM by linearisation is not appropriate in discrete models. Since binary data contains very limited information, it is advised to increase the number of chains to stabilise the estimation. Here we set the number of chains to 10.

```

saemix.options<-list(seed=1234567,save=FALSE,save.graphs=FALSE,
displayProgress=FALSE,nb.chains=10,fim=FALSE)
binary.fit<-saemix(saemix.model,saemix.data,saemix.options)

```

Important note: The linear approximation of the FIM does not apply well to discrete response models. Exact computation methods to estimate the FIM without linearisation have been proposed by [35] using Hamiltonian Monte-Carlo and [45] using adaptive Gaussian quadrature. These methods can be applied to estimate SE for the parameters but are not automatically available yet in `saemix`.

Diagnostics: automated visualisation or diagnostic plots have not yet been implemented for discrete response models, but we can of course create our own in R by simulating from the model. To do this we need to define a simulation function associated with the structural model, with the same arguments as the model function, and returning simulated responses. For the binary model, this function would be the following, where we replace the line defining the log-probability `logp` in `binary.model` with a line simulating from a Binomial distribution with parameter *pevent* for each value of time, and returning those simulated events.

```

simulBinary<-function(psi,id,xidep) {
  tim<-xidep[,1]
  y<-xidep[,2]
  inter<-psi[id,1]
  slope<-psi[id,2]
  logit<-inter+slope*tim
  pevent<-1/(1+exp(-logit))
  ysim<-rbinom(length(tim),size=1,prob=pevent)
  return(ysim)
}
nsim<-100
binary.fit <- simulateDiscreteSaemix(binary.fit, simulBinary, nsim=nsim)

```

In figure 4.21 we use the simulated data in the `datasim` dataframe contained in the `sim.data` element of the object after the call to the function to compute a 90% prediction interval on the proportion of events at each visit. This VPC plot shows a reasonable model fit in the two treatment groups.

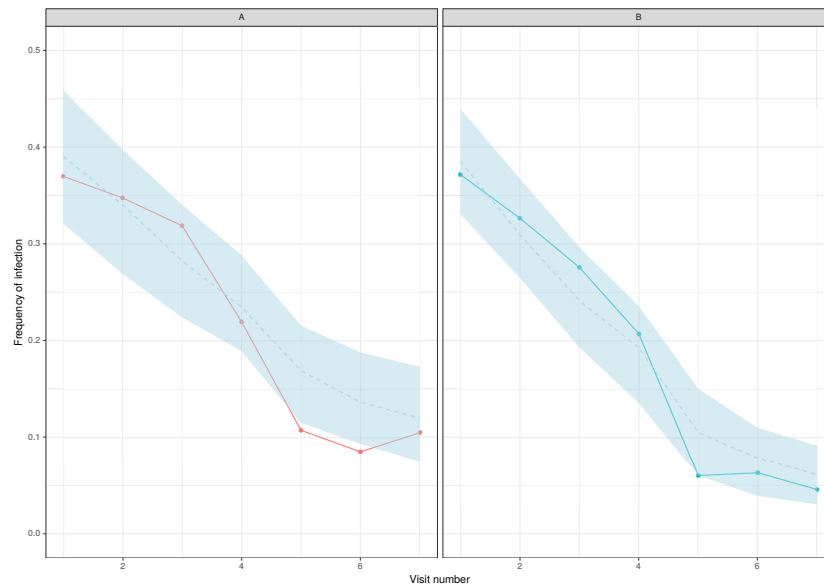


Figure 4.21: Proportion of expected events compared to the observed proportion across time, for the model fit to the toenail data.

The following code was used to produce this plot:

```
simdat <- binary.fit@sim.data@datasim
simdat$visit <- rep(toenail.saemix$visit, nsim)
simdat$treatment <- rep(toenail.saemix$treatment, nsim)

# requires
# library(tidyverse)
# library(ggplot2)
# library(gridExtra)

ytab <- NULL
for(irep in 1:nsim) {
  xtab <- simdat[simdat$irep == irep,]
  xtab1 <- xtab %>%
    group_by(visit, treatment) %>%
    summarise(nev = sum(ysim), n = n()) %>%
    mutate(freq = nev/n)
```

```
  ytab<-rbind(ytab,xtab1[,c("visit","freq","treatment")])
}
gtab <- ytab %>%
  group_by(visit, treatment) %>%
  summarise(lower=quantile(freq, c(0.05)), median=quantile(freq, c(0.5)),
    upper=quantile(freq, c(0.95))) %>%
  mutate(treatment=ifelse(treatment==1,"B","A"))
gtab$freq<-1

# summarising data
toe1 <- toenail.saemix %>%
  group_by(visit, treatment) %>%
  summarise(nev = sum(y), n=n()) %>%
  mutate(freq = nev/n, sd=sqrt((1-nev/n)/nev)) %>%
  mutate(treatment=ifelse(treatment==1,"B","A"))

plot2 <- ggplot(toe1, aes(x=visit, y=freq, group=treatment)) +
  geom_line(aes(colour=treatment)) +
  geom_point(aes(colour=treatment)) +
  geom_line(data=gtab, aes(x=visit, y=median), linetype=2, colour='lightblue') +
  geom_ribbon(data=gtab,aes(ymin=lower, ymax=upper), alpha=0.5, fill='lightblue') +
  ylim(c(0,0.5)) + theme_bw() + theme(legend.position = "none") +
  facet_wrap(~treatment) +
  xlab("Visit number") + ylab("Frequency of infection")
print(plot2)
```


4.6.2 Categorical data

The `knee.saemix` data represents pain scores recorded in a clinical study in 127 patients with sport related injuries treated with two different therapies. The pain occurring during knee movement was observed after 3, 7 and 10 days of treatment. It was taken from the `catdata` package in R [41] (dataset `knee`) and reformatted as follows. A time column was added representing the day of the measurement (with 0 being the baseline value) and each observation corresponds to a different line in the dataset. Treatment was recoded as 0/1 (placebo/treatment), gender as 0/1 (male/female) and `Age2` represents the squared of centered Age. Figure 4.22 shows barplots of the different pain scores as a function of time in study, illustrating a recovery as the proportion of lower pain scores increases.

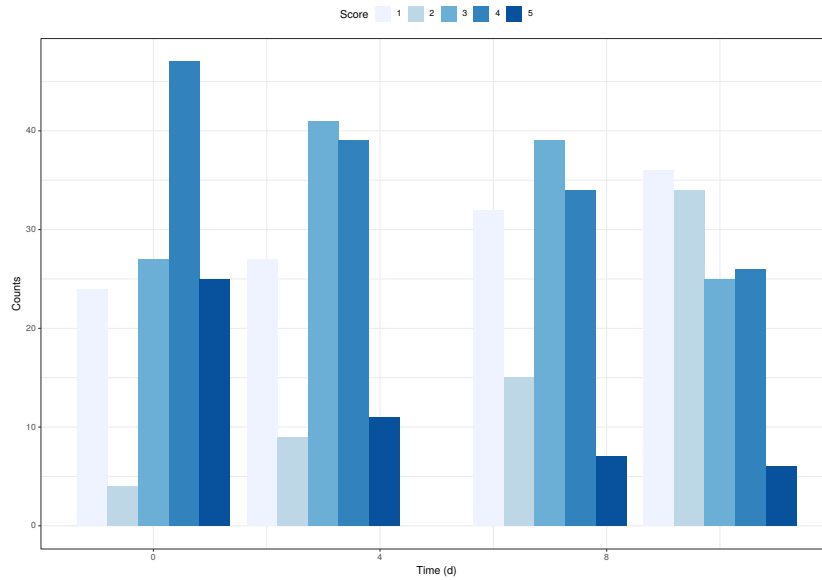


Figure 4.22: Barplot of the evolution of pain scores with time in the `knee.saemix` dataset.

```
data(knee.saemix)
```

The dataset is part of the datasets analysed in [44] with various methods (please refer to the different vignettes in the documentation of `knee`), but mainly as logistic regression on the response after 10 days, or as mixed binary regression after dichotomising the response. Here, we will fit the following proportional odds model to the full data: The probability $p_{ij} = P(Y_{ij} = 1 | \theta_{1,i}, \theta_{2,i})$

associated with an event Y_{ij} at time t_{ij} is given by the following equation for the logit:

$$\begin{aligned}
 \text{logit}(P(Y_{ij} = 1|\psi_i)) &= \alpha_{1,i} + \beta_i t_{ij} \\
 \text{logit}(P(Y_{ij} = 2|\psi_i)) &= \alpha_{1,i} + \alpha_2 \\
 \text{logit}(P(Y_{ij} = 3|\psi_i)) &= \alpha_{1,i} + \alpha_2 + \alpha_3 \\
 \text{logit}(P(Y_{ij} = 4|\psi_i)) &= \alpha_{1,i} + \alpha_2 + \alpha_3 + \alpha_4 \\
 P(Y_{ij} = 4|\psi_i) &= 1 - \sum_k = 1^4 P(Y_{ij} = k|\psi_i)
 \end{aligned} \tag{4.10}$$

where α_1 and β have interindividual variability, β is the effect of time, α_1 is the probability of a pain score of 1 and the other parameters represent an incremental risk to move into the higher pain category.

When the response has more than one category, we define the probability for (K-1) categories and combine them to obtain the likelihood for each observation.

```

ordinal.model<-function(psi,id,xidep) {
  y<-xidep[,1]
  time<-xidep[,2]
  alp1<-psi[id,1]
  alp2<-psi[id,2]
  alp3<-psi[id,3]
  alp4<-psi[id,4]
  beta<-psi[id,5]

  logit1<-alp1 + beta*time
  logit2<-logit1+alp2
  logit3<-logit2+alp3
  logit4<-logit3+alp4
  pge1<-exp(logit1)/(1+exp(logit1))
  pge2<-exp(logit2)/(1+exp(logit2))
  pge3<-exp(logit3)/(1+exp(logit3))
  pge4<-exp(logit4)/(1+exp(logit4))
  pobs<-(y==1)*pge1+(y==2)*(pge2-pge1)+(y==3)*(pge3-pge2)+(y==4)*(pge4-pge3)+(y==5)*(1-pge4)
  logpdf <- log(pobs)
  return(logpdf)
}

saemix.model<-saemixModel(model=ordinal.model,
  description="Ordinal categorical model",modeltype="likelihood",
  psi0=matrix(c(0,0.2, 0.6, 3, 0.2),ncol=5,byrow=TRUE,
    dimnames=list(NULL,c("alp1","alp2","alp3","alp4","beta"))),
  transform.par=c(0,1,1,1,1),

```

```

covariance.model = diag(c(1,0,0,0,1))

saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, nb.chains=10,
  fim=FALSE)

ord.fit<-saemix(saemix.model,saemix.data,saemix.options)

```

Figure 4.23, representing a VPC of the mean value of pain score at each time point, stratified over treatment, shows some model misfit especially for treatment 0. This figure was produced using simulations from the fitted model as in the binary data examples using the simulation function below.

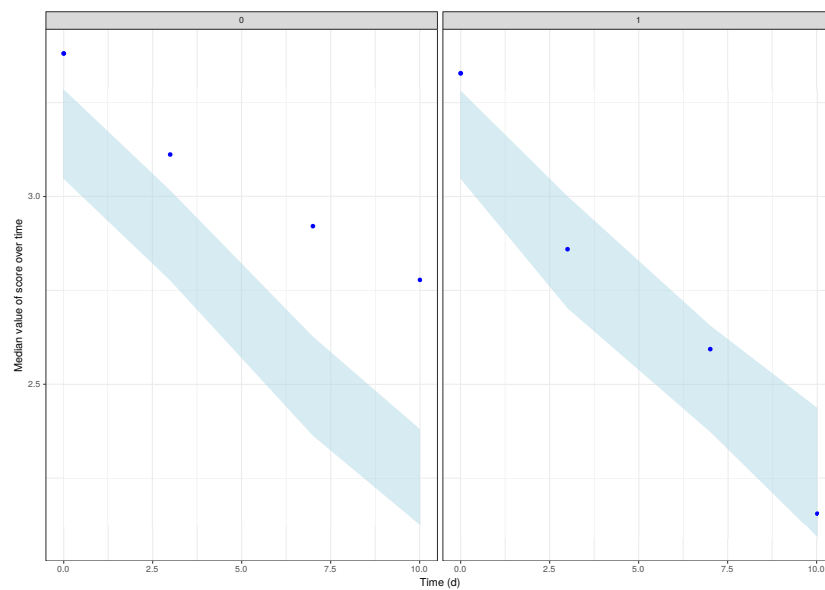


Figure 4.23: VPC of the mean value of pain score in the model adjusted to the knee.saemix dataset.

```

simulateOrdinal<-function(psi,id,xidep) {
  y<-xidep[,1]
  time<-xidep[,2]
  alp1<-psi[id,1]
  alp2<-psi[id,2]
  alp3<-psi[id,3]
  alp4<-psi[id,4]
  beta<-psi[id,5]

```

```

logit1<-alp1 + beta*time
logit2<-logit1+alp2
logit3<-logit2+alp3
logit4<-logit3+alp4
pge1<-exp(logit1)/(1+exp(logit1))
pge2<-exp(logit2)/(1+exp(logit2))
pge3<-exp(logit3)/(1+exp(logit3))
pge4<-exp(logit4)/(1+exp(logit4))
x<-runif(length(time))
ysim<-1+as.integer(x>pge1)+as.integer(x>pge2)+as.integer(x>pge3)+as.integer(x>pge4)
return(ysim)
}

```

4.6.3 Count data

Epilepsy data

We first show a simple example using the epilepsy data from the MASS package. We can fit the Poisson model to the data, which assumes that the probability to observe a count value equal to n is given by:

$$P(Y = n) = \frac{\lambda^n e^{-\lambda}}{n!} \quad (4.11)$$

where $\lambda > 0$ is the parameter of the model. We assume a log-normal distribution for λ .

```

epilepsy<-MASS::epil
saemix.data<-saemixData(name.data=epilepsy, name.group=c("subject"),
  name.predictors=c("period","y"),name.response=c("y"),
  name.covariates=c("trt","base", "age"),
  units=list(x="2-week",y="", covariates=c("", "", "yr")))
## Poisson model with one parameter
countPoi<-function(psi,id,xidep) {
  y<-xidep[,2]
  lambda<-psi[id,1]
  logp <- -lambda + y*log(lambda) - log(factorial(y))
  return(logp)
}
saemix.model<-saemixModel(model=countPoi,description="Count model Poisson",
  modeltype="likelihood",
  psi0=matrix(c(0.5),ncol=1,byrow=TRUE,dimnames=list(NULL, c("lambda"))),

```

```
transform.par=c(1))
saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE,
  displayProgress=FALSE)
poisson.fit<-saemix(saemix.model,saemix.data,saemix.options)
```

RAPI data

This dataset was kindly made available by David Atkins (University of Washington) in his tutorial on modelling count data [1]. The RAPI data studies gender differences across two years in alcohol-related problems, as measured by the Rutgers Alcohol Problem Index (RAPI) [46]. Students were asked to report every six months the number of alcohol-related problems, and the dataset includes 3,616 repeated measures of these counts in 818 subjects, 561 of whom had the full 5 measurements over a period of 2 years. Interesting features of this dataset are first, the longitudinal aspect which allow to evaluate changes over time, and second, the shape of the distribution of counts. Counts are often positively skewed, bounded by zero, with a large stack of data points at zero, indicating individuals and/or occasions without drinking, use, or related problems. This dataset was used in [1] to illustrate mixed effects count regression using the `glmer()` function from the `lme4`.

The dataset is available in `saemix` under the name `rapi.saemix` so we read it and create our `saemixData` object in the usual way. Because we need the value of the outcome to compute the corresponding likelihood, the `rapi` column is used both as a predictor and as the response:

```
data(rapi.saemix)
saemix.data<-saemixData(name.data=rapi.saemix, name.group=c("id"),
  name.predictors=c("time","rapi"),name.response=c("rapi"),
  name.covariates=c("gender"),
  units=list(x="months",y="",covariates=c("")))
hist(rapi.saemix$rapi, main="", xlab="RAPI score", breaks=30)
```

`saemix` currently does not have automated plots for discrete outcome data, but we can produce our own histogram (here, across all measurements without taking time into account) to notice that indeed, there seems to be many subjects reporting no alcohol related problems over some periods.

Poisson model: the first model we can fit to this data is, as previously, the Poisson model, but this time we add a time effect. Here we will write the same model as in `glmer()` to compare our results. In `glmer()` a logarithmic link function is used to transform the mean of the Poisson model (λ) into a linear predictor of time and covariates. Random effects are then added to the parameters of the linear model. To take into account the change with time in `saemix`, we need to rewrite the previous model to use normal distributions for the parameters and explicitly write the linear model in the function, as follows:

```
count.poisson<-function(psi,id,xidep) {
  time<-xidep[,1]
  y<-xidep[,2]
  intercept<-psi[id,1]
  slope<-psi[id,2]
  lambda<- exp(intercept + slope*time)
  logp <- -lambda + y*log(lambda) - log(factorial(y))
  return(logp)
}
```

The expression of $\log p$ in the model function is unchanged, but now we define a log-normal distribution for λ within the model so that we can use two parameters and time as a predictor. The statistical model also changes to reflect this, as our parameters intercept and slope are now on the scale of the random effects, so they are given a normal distribution. Defining and fitting this model in `saemix`, we have:

```
saemix.model.poi<-saemixModel(model=count.poisson,description="Count model Poisson",
  modeltype="likelihood",
  psi0=matrix(c(log(5),0.01),ncol=2,byrow=TRUE,dimnames=list(NULL, c("intercept","slope")),
  transform.par=c(0,0), omega.init=diag(c(0.5, 0.5)))
saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, displayProgress=FALSE)
poisson.fit<-saemix(saemix.model.poi,saemix.data,saemix.options)
```

Note that when parameters enter the model through a normal distribution, we may need to adjust the initial values of the Ω matrix (argument `omega.init`) to avoid failure to find valid initial parameter estimates.

We can also add the covariate gender to both parameters as well as a correlation between the two random effects:

```
modsmx.poi.cov2<-saemixModel(model=count.poisson,
  description="Count model Poisson",modeltype="likelihood",
  psi0=matrix(c(log(5),0.01),ncol=2,byrow=TRUE,dimnames=list(NULL,
  c("intercept","slope"))), transform.par=c(0,0),
  omega.init=diag(c(0.5, 0.5)),
  covariance.model=matrix(data=1, ncol=2, nrow=2),
  covariate.model=matrix(c(1,1), ncol=2, byrow=TRUE))
poisson.fit.cov2<-saemix(modsmx.poi.cov2,saemix.data,saemix.options)
```

Comparing the parameter estimates from this fit to the estimates obtained by `glmer()` using a Laplace approximation in Table 2 of [1], we find very good agreement with the SAEM algorithm.

Note: saemix does not provide adequate standard errors of estimation for the parameters in version 3.0. The FO-approximation of the FIM implemented in the current version of the algorithm is known to be very poor for discrete outcome models.

Some diagnostics for this model can be obtained by simulating from the model. To do this we need to define a simulation function associated with the structural model, with the same arguments as the model function, and returning simulated responses. For the Poisson model, this function would be the following, where we replace the line defining the log-probability `logp` in `count.poisson` with a line simulating from a Poisson distribution with parameter λ for each value of time, and returning those simulated counts.

```
saemix.simulatePoisson<-function(psi, id, xidep) {
  time<-xidep[,1]
  y<-xidep[,2]
  intercept<-psi[id,1]
  slope<-psi[id,2]
  lambda<- exp(intercept + slope*time)
  y<-rpois(length(time), lambda=lambda)
  return(y)
}
```

We then use the `simulateDiscreteSaemix` function to obtain simulations from the model, using the estimated parameters. Here we set the number of simulations to 100 as the dataset is large and we are interested in global diagnostics.

```
yfit1<-simulateDiscreteSaemix(poisson.fit.cov2, simulate.function=saemix.simulatePoisson,
  nsim=100)
cat("Observed proportion of 0's",
  length(yfit1@data@data$rap1[yfit1@data@data$rap1==0])/yfit1@data@ntot.obs,"\n")
cat("      Poisson model, p=",
  length(yfit1@sim.data@datasim$ysim[yfit1@sim.data@datasim$ysim==0])/
  length(yfit1@sim.data@datasim$ysim),"\n")
```

Handling overdispersion: the model predicts a lower proportion of subjects without alcohol-related problems than we observe in data, a sign of overdispersion (with a Poisson model, the mean of the Poisson distribution, λ , is equal to the variance, an assumption which is violated here). Several models can be used to take this feature into account. First, we can use the Zero-Inflated Poisson model, where the number of counts equal to 0 is increased. This model can be built as a mixture between a distribution of 0's with probability p_0 and a standard Poisson model. We modify our model function above to:

```
count.poissonzip<-function(psi,id,xidep) {
  time<-xidep[,1]
  y<-xidep[,2]
  intercept<-psi[id,1]
  slope<-psi[id,2]
  p0<-psi[id,3] # Probability of zero's
  lambda<- exp(intercept + slope*time)
  logp <- log(1-p0) -lambda + y*log(lambda) - log(factorial(y)) # Poisson
  logp0 <- log(p0+(1-p0)*exp(-lambda)) # Zeroes
  logp[y==0]<-logp0[y==0]
  return(logp)
}
```

and fit the model using:

```
modsmx.zip<-saemixModel(model=count.poissonzip,description="count model ZIP",
  modeltype="likelihood",
  psi0=matrix(c(1.5, 0.01, 0.2),ncol=3,byrow=TRUE,dimnames=list(NULL,
    c("intercept", "slope","p0"))),
  transform.par=c(0,0,3), covariance.model=diag(c(1,1,0)),
  omega.init=diag(c(0.5,0.3,0)),
  covariate.model = matrix(c(1,1,0),ncol=3, byrow=TRUE))
```

```
zippoisson.fit <- saemix(modsmx.zip,saemix.data,saemix.options)
```

where we assume a logit-normal distribution for the added parameter p_0 through the `transform.par` argument. We could also model $\text{logit}(p_0)$ on a normal scale if we needed to add a time effect. The same approach as above can then be used to simulate from the model, this time using the simulation function:

```
saemix.simulatePoissonZIP<-function(psi, id, xidep) {
  time<-xidep[,1]
  y<-xidep[,2]
  intercept<-psi[id,1]
  slope<-psi[id,2]
  p0<-psi[id,3] # Probability of zero's
  lambda<- exp(intercept + slope*time)
  prob0<-rbinom(length(time), size=1, prob=p0)
  y<-rpois(length(time), lambda=lambda)
  y[prob0==1]<-0
  return(y)
}
```


and we can check that the proportion of simulated 0's is now closer to the observed value.

A second type of model used in [1] is the hurdle model, which combines a binary logistic model for the probability of having counts greater than 0, with a truncated Poisson model for counts greater than 0. To implement this, we fit separately the two models: the binary logistic model is fit to the data where we use as a response the binary outcome with values 0 for counts of 0 and 1 for counts strictly positive, then the Poisson model is fit to the data excluding zero counts.

Other possible models include the negative binomial model and generalised Poisson models with additional parameters handling the overdispersion.

Diagnostics: the simulation functions can be used to produce diagnostic plots. As an example, we can compare the expected proportion of 0's, representing subjects without alcohol problems, versus time and stratified by gender, to compare the different models (the code available as a notebook on the github for saemix). The results, shown in Figure 4.24. We could also look at the counts for different categories or the evolution of the median counts.

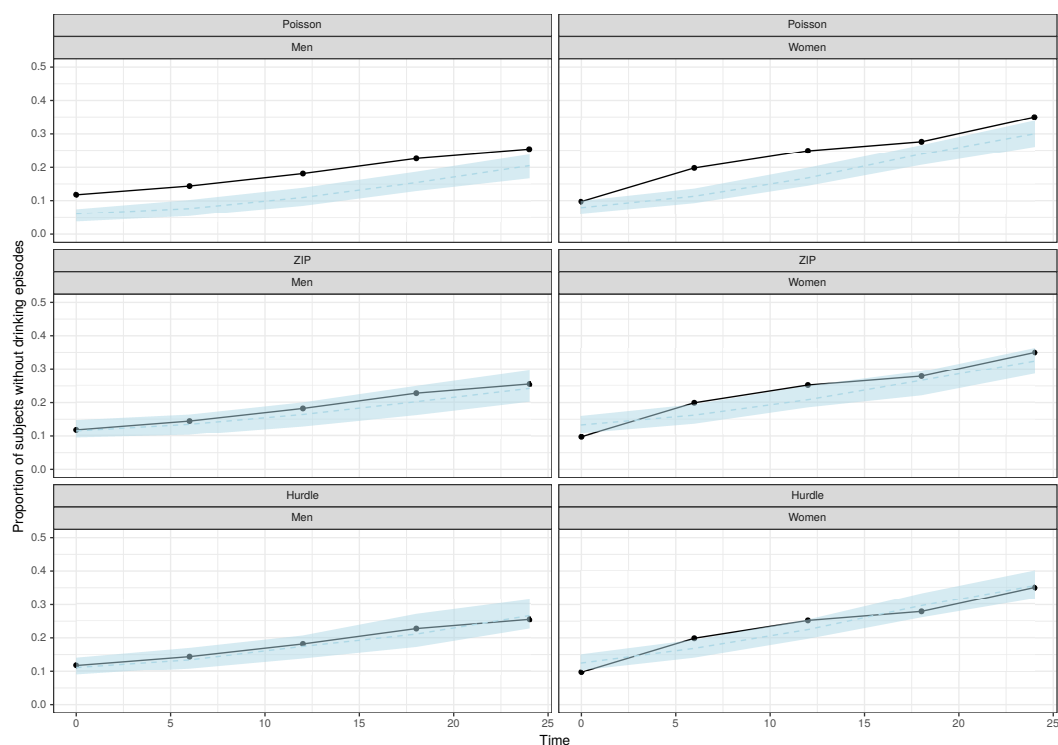


Figure 4.24: Proportion of subjects without drinking problems versus time, for men and women, observed and expected for the Poisson, ZIP and hurdle models, for the RAPI data.

4.7 Time-to-event data

4.7.1 Single event

The example chosen to illustrate the analysis of time-to-event data is the NCCTG Lung Cancer Data, describing the survival in patients with advanced lung cancer from the North Central Cancer Treatment Group [26]. Covariates measured in the study include performance scores rating how well the patient can perform usual daily activities. We reformatted the cancer dataset provided in the survival package in R [43] in SAEM format: patients with missing age, sex, institution or physician assessments were removed from the dataset. Status was recoded as 1 for death and 0 for a censored event, and a censoring column was added to denote whether the patient was dead or alive at the time of the last observation. A line at time=0 was added for all subjects. Finally, subjects were numbered consecutively from 0 to 1.

We can use a Weibull model for the hazard, parameterised as λ and β . For individual i , the hazard function of this model is:

$$h(t, \psi_i) = \frac{\beta_i}{\lambda_i} \left(\frac{t}{\lambda_i} \right)^{\beta_i - 1}. \quad (4.12)$$

Here, the vector of individual parameters is $\psi_i = (\lambda_i, \beta_i)$. These two parameters are assumed to be independent and lognormally distributed:

$$\log(\lambda_i) \sim \mathcal{N}(\log(\lambda_{\text{pop}}), \omega_\lambda^2), \quad (4.13)$$

$$\log(\beta_i) \sim \mathcal{N}(\log(\beta_{\text{pop}}), \omega_\beta^2). \quad (4.14)$$

Then, the vector of population parameters is $\theta = (\lambda_{\text{pop}}, \beta_{\text{pop}}, \omega_\lambda, \omega_\beta)$.

The survival function for this model is:

$$S(t) = e^{-\left(\frac{t}{\lambda}\right)^\beta}$$

The model function for saemix needs to define the log-pdf for each observation. At time 0, it is 0 (no event has occurred yet). For a censored event, the log-likelihood is equal to the logarithm of the survival function since the beginning of the observation period, while for an observed event we add the logarithm of the hazard at the time of the event. In the model below, we pass individual censoring times as the third predictor, so that each individual may have his or her own follow-up duration.

```
weibulltte.model<-function(psi,id,xidep) {
  T<-xidep[,1]
  y<-xidep[,2] # events (1=event, 0=no event)
```

```

cens<-which(xidep[,3]==1) # censoring times (subject specific)
init <- which(T==0)
lambda <- psi[id,1] # Parameters of the Weibull model
beta <- psi[id,2]
Nj <- length(T)

ind <- setdiff(1:Nj, append(init,cens)) # indices of events
hazard <- (beta/lambda)*(T/lambda)^(beta-1) # H'
H <- (T/lambda)^beta # H
logpdf <- rep(0,Nj) # ln(l(T=0))=0
logpdf[cens] <- -H[cens] + H[cens-1] # ln(l(T=censoring time))
logpdf[ind] <- -H[ind] + H[ind-1] + log(hazard[ind]) # ln(l(T=event time))
return(logpdf)
}

```

Important note: In TTE models with a single event, there is not enough information to estimate interindividual variability, but `saemix` needs at least one parameter to run. In this case, we include a random effect in the model but it cannot be estimated properly.

Diagnostics: Automated visualisation or diagnostic plots have not yet been implemented for discrete response models, but we can of course create our own in R. In Figure 4.25 we used the linear approximation of the FIM and the delta-method to compute a very rough estimate of the confidence interval on the predicted survival curve, overlaying it to the non-parametric Kaplan-Meier estimate provided by the `survival` package. Here despite its shortcomings the FIM approximation seems to be adequate.

We can also use simulations to compute the normalised prediction discrepancies (npd) developed by [3].

4.7.2 Repeated time-to-event

For repeated time-to-event data, we use the same model function as above, as the likelihood of an event will be defined relative to the previous event until censoring occurs.

Repeated events were generated using `simulx` (`mlxR` package in R), for $N = 100$ individuals, using the Weibull model (4.12) with $\lambda_{\text{pop}} = 10$, $\omega_{\lambda} = 0.3$, $\beta_{\text{pop}} = 3$ and $\omega_{\beta} = 0.3$ and assuming a right censoring time $\tau_c = 20$.

The following code was used in R to run this example:

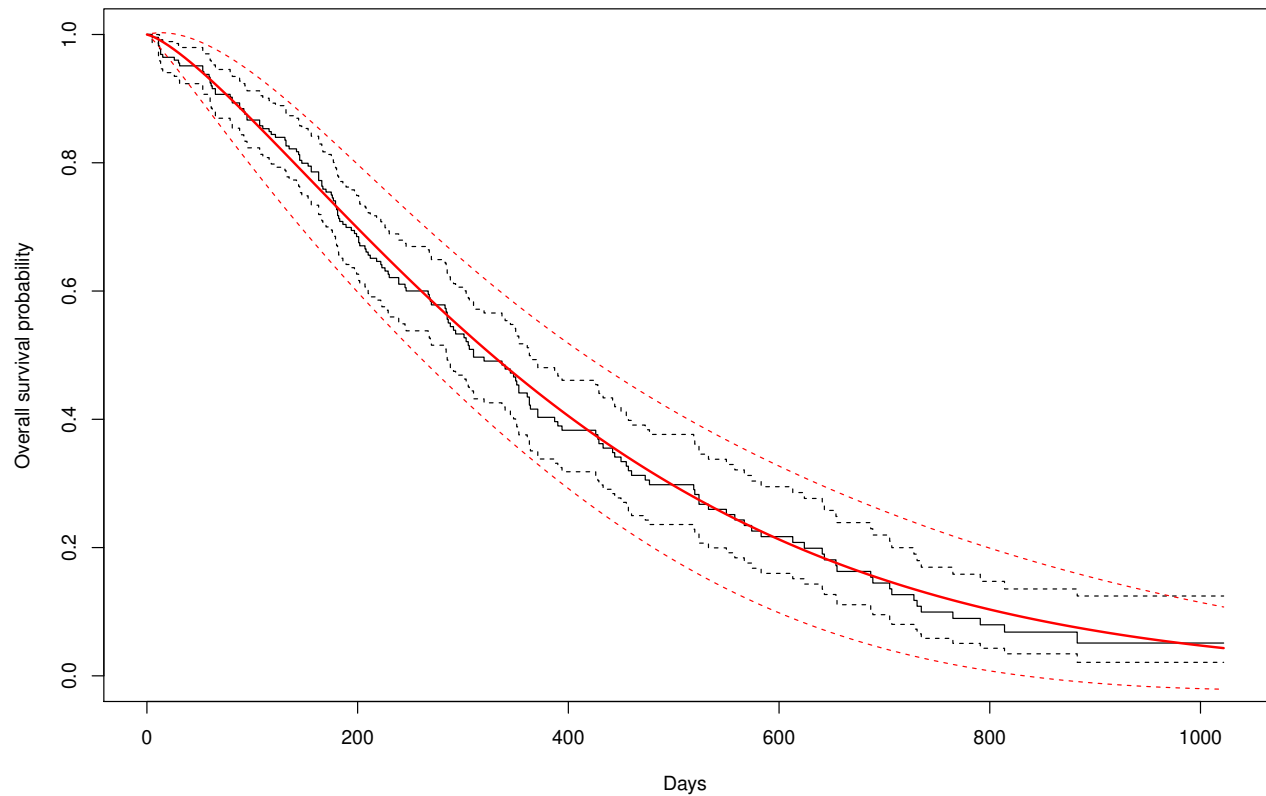


Figure 4.25: Survival function, as a Kaplan-Meier estimate (black) and fitted by a Weibull model with `saemix` (red).

```
data(tte.saemix)
saemix.data<-saemixData(name.data=tte.saemix,header=TRUE,sep=" ",na=NA,
  name.group=c("id"),name.response=c("y"),name.predictors=c("time","y"), name.X=c("time"))

timetoevent.model<-function(psi,id,xidep) {
  T<-xidep[,1]
  N <- nrow(psi)
  Nj <- length(T)
  censoringtime = 20
  lambda <- psi[id,1]
  beta <- psi[id,2]
  init <- which(T==0)
  cens <- which(T==censoringtime)
```

```

ind <- setdiff(1:Nj, append(init,cens))
hazard <- (beta/lambda)*(T/lambda)^(beta-1)
H <- (T/lambda)^beta
logpdf <- rep(0,Nj)
logpdf[cens] <- -H[cens] + H[cens-1]
logpdf[ind] <- -H[ind] + H[ind-1] + log(hazard[ind])
return(logpdf)
}

saemix.model<-saemixModel(model=timetoevent.model,description="time model",
  type="likelihood",
  psi0=matrix(c(2,1),ncol=2,byrow=TRUE,dimnames=list(NULL,c("lambda","beta"))),
  transform.par=c(1,1),covariance.model=matrix(c(1,0,0,1),ncol=2,byrow=TRUE))

saemix.options<-list(map=F,fim=F,ll.is=F, nb.chains = 1, nbiter.saemix =c(200,100), display=
saemix.fit<-saemix(model,saemix.data,saemix.options)

```

Figure 4.26 shows the convergence of the population parameters for this example. The results are summarised in the following table:

```

-----
----                      Results                      ----
-----
----- Fixed effects -----
-----
      Parameter Estimate
[1,] lambda      5.0
[2,] beta        2.8
-----
----- Variance of random effects -----
-----
      Parameter      Estimate
lambda omega2.lambda 0.039
beta   omega2.beta   0.921
-----
----- Correlation matrix of random effects -----
-----
              omega2.lambda omega2.beta
omega2.lambda 1              0
omega2.beta   0              1

```

Figure 4.26: Convergence plot obtained for the RTTE data

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