Saemix 3 - showcasing binary and categorical models

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Version

Use saemix version ≥ 3.2

Objective

Run binary and categorical models in saemix

This notebook uses additional result files from the **saemix** development github (https://github.com/saemixdevelopment/saemixextension), not integrated in the package to avoid bloating. The *workDir* folder in the next chunk of code points to the folder where the user stored this code, and is needed to run the notebook (*workDir* defaults to the current working directory). Specifically, the notebook loads the results for the bootstrap runs performed using different approaches (see Comets et al. Pharm Res 2021). Bootstraps can be run instead by switching the *runBootstrap* variable to TRUE in the first chunk of code:

- in the code, the number of bootstraps is set to 10 for speed but we recommend to use at least 200 for a 90% CI
- this can be changed in the following change of code by uncommenting the line *nboot*<-200 and setting the number of bootstrap samples (this may cause memory issues in **Rstudio** with older machines, if this is the case we recommend executing the code in a separate script)

The current notebook can be executed to create an HMTL or PDF output with comments and explanations. A script version containing only the R code is also given as $saemix3_categoricalModel.R$ in the same folder.

Binary response model

Data description The *toenail.saemix* dataset in the **saemix** package contains binary data from a randomised clinical trial comparing two treatments for fungal toenail infection. The original data is available in **R** as the *toenail* dataset in the package **prLogistic** and has been reformatted for **saemix**.

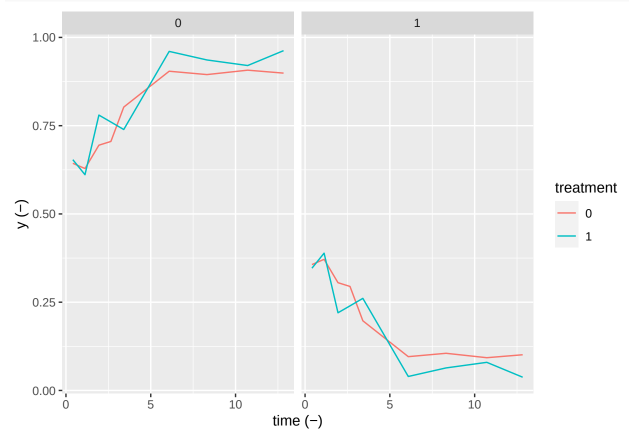
The data was collected in a multi-center randomised comparison of two oral treatments (A and B) for toenail infection. 294 patients are measured at seven visits, i.e. at baseline (week 0), and at weeks 4, 8, 12, 24, 36, and 48 thereafter, comprising a total of 1908 measurements. The primary end point was the presence 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 (0=none or mild versus 1=moderate or severe).

To create the data object using saemixData, we need to specify the response column both as a response (name.response="y") and as a predictor (here, time is the first predictor and we add the response in the argument name.predictors).

Exploring data The usual plot of the data object is not very informative as it alternates between 0 and 1's. Instead we plot the evolution of the frequency of infection over time in the population, stratifying by treatment. We use the *plotDiscreteData()* function from the package, setting the *outcome* argument to 'binary'.

```
# Distribution of times
if(FALSE) hist(toenail.saemix$time, breaks=c(-1,0.25,1.25,2.25, 3.25, 7,10,15,20), freq=T)
table(cut(toenail.saemix$time, breaks=c(-1,0.25,1.25,2.25, 3.25, 7,10,15,20)))
##
##
     (-1,0.25] (0.25,1.25] (1.25,2.25] (2.25,3.25]
                                                         (3.25,7]
                                                                        (7,10]
                                                 240
                                                                           245
##
           294
                        278
                                    272
                                                              291
                    (15, 20]
##
       (10, 15]
##
           282
```

```
# Proportion of 0's and 1's across time
plotDiscreteData(saemix.data, outcome='binary', which.cov="treatment")
```



We can also present the results as a barplot of the proportion of events, using **ggplot2** and **tidyverse**.

```
# Barplots across time
toe1 <- toenail.saemix %>%
  group_by(visit, treatment) %>%
  summarise(nev = sum(y), n=n()) %>%
  mutate(freq = nev/n, sd=sqrt((1-nev/n)/nev)) %>%
  mutate(lower=freq-1.96*sd, upper=freq+1.96*sd)
```

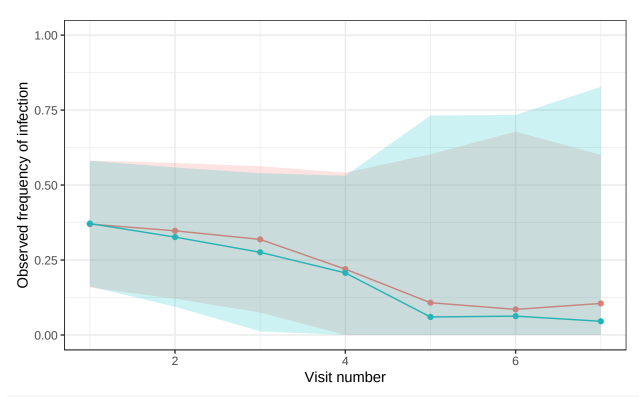
`summarise()` has grouped output by 'visit'. You can override using the
`.groups` argument.

```
toe1$lower[toe1$lower<0] <-0 # we should use a better approximation for CI
toe1$treatment <- factor(toe1$treatment, labels=c("A","B"))

plot1<-ggplot(toe1, aes(x=visit, y=freq, group=treatment)) + geom_line(aes(colour=treatment)) +
    geom_point(aes(colour=treatment)) +
    geom_ribbon(aes(ymin=lower, ymax=upper, fill=treatment), alpha=0.2) +
    ylim(c(0,1)) + theme_bw() + theme(legend.position = "top") +
    xlab("Visit number") + ylab("Observed frequency of infection")

print(plot1)</pre>
```

treatment - A - B



```
if(saveFigs) {
  namfig<-"toenail_infectionFreq.eps"
  cairo_ps(file = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, wideline plot(plot1)
  dev.off()
}</pre>
```

Statistical model The model fit here is a logistic random effect model developed by (Hedeker et al. 1994). This model includes a random intercept (θ_1 and ω_1), a time effect (θ_2) and treatment (A or B) (β) as a covariate affecting the slope θ_2 . We considered the interaction term between time and treatment but no treatment effect alone as it would impact the intercept which shouldn't be different between arms due to the randomisation process. The time course was assumed to be similar across subjects as no significant interindividual variability was found when testing different models.

- Model for repeated binary data
 - the probability that y_{ij} outcome observed in subject i at visit j (time t_{ij} in months) is 1 is modelled

as a logistic model

- linear model on the logit scale (logit(p) = ln $\left(\frac{p}{1-p}\right)$)

$$logit(P(y_{ij} = 1)) = \theta_{1,i} + \theta_{2,i}t_{ij}$$

- Statistical model
 - $-\theta_{1,i}$ assumed to follow a normal distribution $N(\mu_{\theta_1},\omega_{\theta_1})$
 - $-\theta_{2,i}$ assumed to depend on treatment as in $\theta_{2,i} = \mu_{\beta} + 1_{trt=B}$

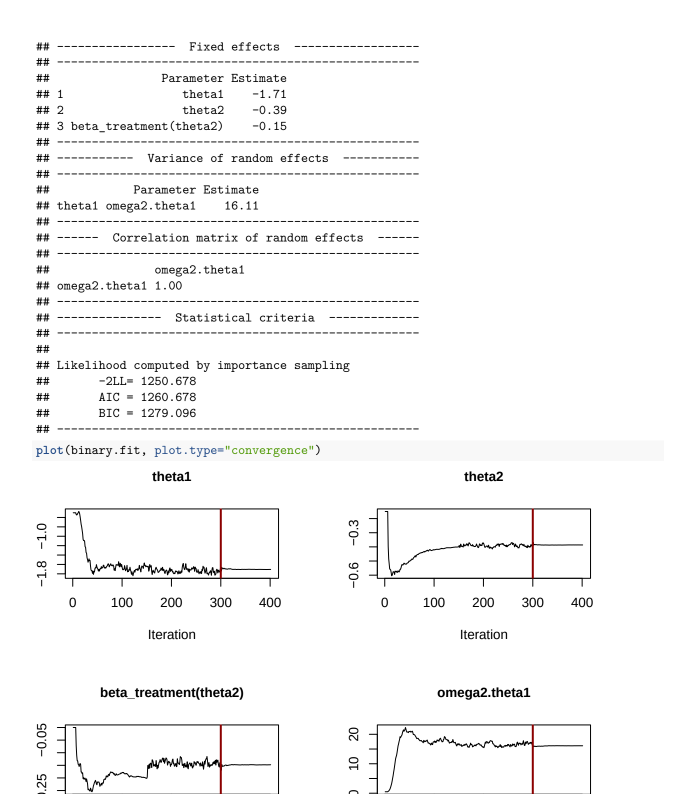
In the following chunk of code, we define the model and the simulate function, then create the saemixModel object.

```
# saemix model
binary.model<-function(psi,id,xidep) {</pre>
       tim<-xidep[,1]</pre>
       y < -xidep[,2]
       inter<-psi[id,1]</pre>
       slope<-psi[id,2]</pre>
       logit<-inter+slope*tim</pre>
       pevent<-exp(logit)/(1+exp(logit))</pre>
       logpdf<-rep(0,length(tim))</pre>
       P.obs = (y==0)*(1-pevent)+(y==1)*pevent
       logpdf <- log(P.obs)</pre>
       return(logpdf)
}
# simulation function (used for diagnostics)
simulBinary<-function(psi,id,xidep) {</pre>
       tim<-xidep[,1]
       y < -xidep[,2]
       inter<-psi[id,1]</pre>
       slope<-psi[id,2]</pre>
       logit<-inter+slope*tim</pre>
       pevent<-1/(1+exp(-logit))</pre>
       ysim<-rbinom(length(tim),size=1, prob=pevent)</pre>
       return(ysim)
saemix.model<-saemixModel(model=binary.model,description="Binary model",simulate.function=simulBinary, named to the same same named to the same named t
                                                                                               psi0=matrix(c(-0.5,-.15,0,0),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1",
                                                                                               transform.par=c(0,0), covariate.model=c(0,1),covariance.model=matrix(c(1,0,0,
```

Fitting the model

- Fit with saemix and store the results to the object binary.fit
 - setting options
 - 10 chains
 - don't display intermediate graphs or save graphs
 - don't compute the FIM (approximation not suited to discrete data models)

```
# saemix fit
saemix.options<-list(seed=1234567,save=FALSE,save.graphs=FALSE, displayProgress=FALSE, nb.chains=10, fit
binary.fit<-saemix(saemix.model,saemix.data,saemix.options)
summary(binary.fit)</pre>
```



- Results numerical output - note that the estimated value of μ_{θ_1} is -1.71, corresponding to an estimated probability of event of 0.154 - this is lower than the observed probability of infection at time 0 (around 0.37) because the

Iteration

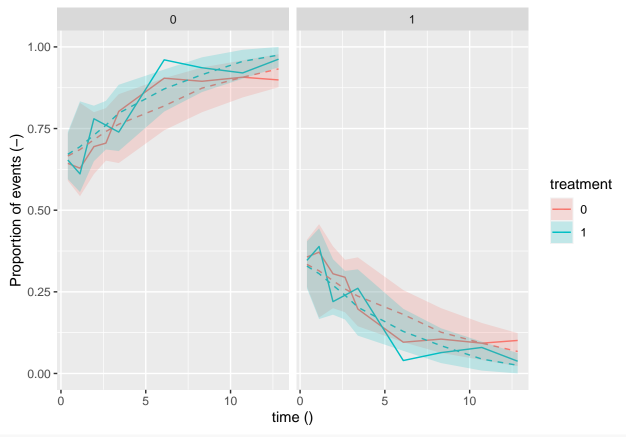
Iteration

logistic model is highly non-linear and $E(f(\theta))$ is very different from $f(E(\theta))$ - convergence plots show good convergence for all parameters

Diagnostics

- Simulation function to simulate from a binary model
 - the model function defines directly the log-pdf, so the user needs to define a function to simulate from the appropriate binomial function
 - note the similarities between the model function (binary.model()) and the simulation function (simulBinary())
 - * same setting of dependent variables (tim and y) from xidep and parameters (inter and slope) from psi
 - · note that we don't use y in simulBinary()
 - * same definition of pevent (= $P(Y_{ij} = 1)$, the probability of observing an event)
 - * in binary.model() we then compute the probability of the observed outcome using the observed value of Y_{ij} contained in y for each observation
 - * in simulBinary(), we use the individual $P(Y_{ij} = 1)$ predictions to simulate from a Bernouilli distribution using the rbinom() function
- once the simulation function has been defined and associated with the model component of the object, we use the *simulateDiscreteSaemix()* function from the **saemix** package to simulate *nsim* values (here 100) with the population parameters estimated in *binary.fit*
 - this adds a *simdata* element to the *binary.fit*
 - the simulations are used to produce VPC via the discreteVPC() function, again specifying the outcome to be binary. Here, we stratify the plot on the treatment covariate

```
# $1_{Y_{ij}=0} \times (1-P(Y_{ij}=1)) + 1_{Y_{ij}=1} \times P(Y_{ij}=1) $
# simulate from model (nsim=100)
nsim<-1000
binary.fit <- simulateDiscreteSaemix(binary.fit, nsim=nsim)
discreteVPC(binary.fit, outcome="binary", which.cov="treatment")</pre>
```



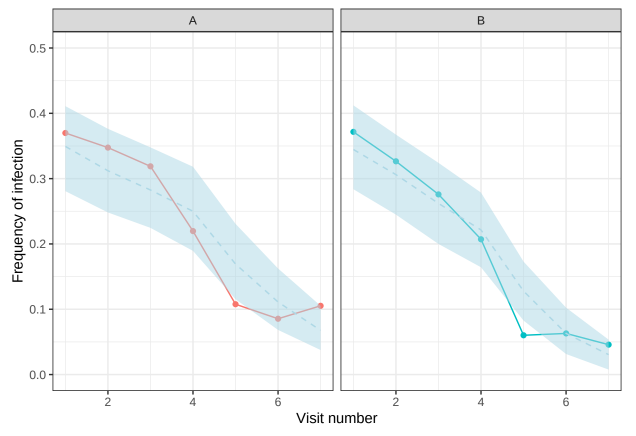
```
if(saveFigs) {
  namfig<-"toenail_vpcByTreatment.eps"
  cairo_ps(file = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widdiscreteVPC(binary.fit, outcome="binary", which.cov="treatment")
  dev.off()
}</pre>
```

We can also extract dataframe with the simulated data (binary.fit@sim.data@datasim) and produce diagnostics in R. Below, using **tidyverse**, we add columns visit and treatment to plot the frequency of infection over time for each treatment.

```
simdat <-binary.fit@sim.data@datasim</pre>
simdat$visit<-rep(toenail.saemix$visit,nsim)</pre>
simdat$treatment<-rep(toenail.saemix$treatment,nsim)</pre>
# VPC-type diagnostic
ytab<-NULL
for(irep in 1:nsim) {
  xtab<-simdat[simdat$irep==irep,]</pre>
  suppressMessages(
  xtab1 <- xtab %>%
    group_by(visit, treatment) %>%
    summarise(nev = sum(ysim), n=n()) %>%
    mutate(freq = nev/n)
  )
  ytab<-rbind(ytab,xtab1[,c("visit","freq","treatment")])</pre>
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

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")</pre>
```



```
if(saveFigs) {
  namfig<-"toenail_ggplot2VPCTreatment.eps"
  cairo_ps(file = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widely plot(plot2)
  dev.off()
}</pre>
```

Work in progress

- npde for categorical data (submitted)
 - TODO using code from Marc Cerou

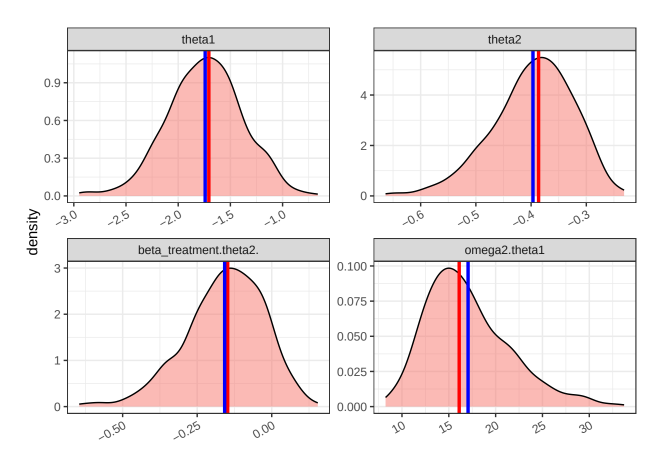
npd

Standard errors of estimation The computation of the FIM in saemix uses the so-called FOCE method, an approximation where the model function f is linearised around the conditional expectation of the individual parameters. This approximation is particularly poor for discrete data models, which is why currently saemix doesn't provide estimation errors for categorical/binary data models. In this document we show how to obtain SE through the computation of the exact FIM using numerical integration, as well as a bootstrap approach. Because all subjects have different times, for the binary data we will use bootstrap approaches as computing the exact FIM is time-consuming and we would need to compute it for each subject separately before summing the individual FIMs.

Different bootstrap approaches can be used in non-linear mixed effect models and have been implemented for saemix in Comets et al. 2021, with code available on the github.

Case bootstrap The first bootstrap approach we can use is case bootstrap, where we resample at the level of the individual. We plot the bootstrap distribution for the 4 parameters (intercept, slope, treatment effect on slope, and variability of intercept). The red vertical line represents the estimate obtained on the original data while the blue line shows the mean of the bootstrap distribution.

```
if(!runBootstrap) {
  case.bin <- read.table(file.path(saemixDir, "bootstrap", "results", "toenail caseBootstrap.res"), header</pre>
 nboot<-dim(case.bin)[1]</pre>
  else case.bin <- saemix.bootstrap(binary.fit, method="case", nboot=nboot)</pre>
head(case.bin)
##
     Replicate
                  theta1
                              theta2 beta_treatment.theta2. omega2.theta1
## 1
             1 -1.851662 -0.2905759
                                                 -0.21093899
                                                                   13.21270
## 2
             2 -1.582608 -0.3956728
                                                                   21.50126
                                                 -0.14496853
## 3
             3 -1.290633 -0.4133164
                                                 -0.43242325
                                                                   23.04858
## 4
             4 -1.851579 -0.4228992
                                                 -0.17280887
                                                                   19.78489
## 5
             5 -1.453970 -0.4218880
                                                 -0.01963166
                                                                   11.77697
## 6
             6 -1.728966 -0.3965398
                                                 -0.18952473
                                                                   13.28420
# Bootstrap distributions
if(nboot<200) cat("The number of bootstrap samples is too low to provide good estimates of the confiden
  resboot1<-case.bin
  ypd2<-NULL
  for(icol in 1:4) {
    ypd2<-rbind(ypd2,data.frame(rep=resboot1[,1],Param=colnames(resboot1)[(icol+1)],value=resboot1[,(icol+1)]</pre>
  ypd2$Param<-factor(ypd2$Param, levels = unique(ypd2$Param))</pre>
  ypd2.fix<-ypd2[ypd2$Param %in% unique(ypd2$Param)[1:3],]</pre>
  ypd2.iiv<-ypd2[ypd2$Param %in% unique(ypd2$Param)[4],]</pre>
  ypd <- ypd2
  par.estim<-c(binary.fit@results@fixed.effects,diag(binary.fit@results@omega)[binary.fit@results@indx.
  mean.bootDist<-apply(resboot1, 2, mean)[-c(1)]</pre>
  df<-data.frame(Param=unique(ypd2$Param), mean.boot=mean.bootDist, est.saemix=par.estim, Bootstrap="Ca
  plot.density2<-ggplot(data=ypd2) + geom_density(aes(value,fill="red4"), alpha=0.5) +
    geom_vline(data=df,aes(xintercept=est.saemix),colour="red",size=1.2) +
    geom_vline(data=df,aes(xintercept=mean.boot),colour="blue",size=1.2) +
    theme_bw() + theme(axis.title.x = element_blank(),axis.text.x = element_text(size=9, angle=30, hjus
    facet_wrap(~Param, ncol=2, scales = 'free')
  print(plot.density2)
```



Conditional bootstrap We can also use conditional bootstrap, a non-parametric residual bootstrap which bootstraps samples from the conditional distributions and preserves the exact structure of the original dataset.

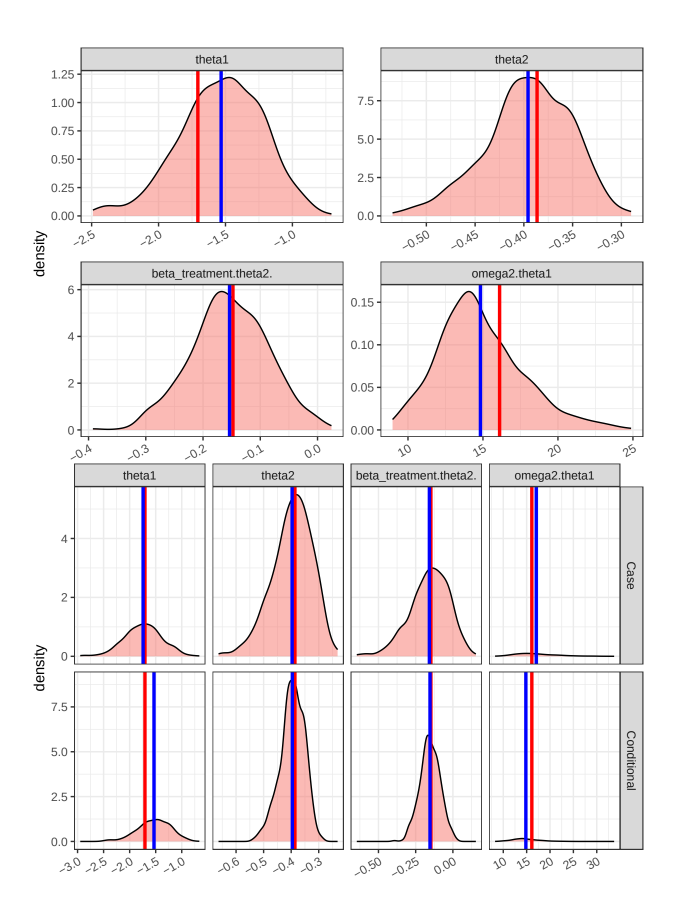
```
if(!runBootstrap) {
  cond.bin <- read.table(file.path(saemixDir, "bootstrap", "results", "toenail_condBootstrap.res"), header
  nboot<-dim(cond.bin)[1]
  } else
    cond.bin <- saemix.bootstrap(binary.fit, method="conditional", nboot=nboot)
summary(cond.bin)</pre>
```

```
##
      Replicate
                         theta1
                                            theta2
                                                           beta_treatment.theta2.
##
           : 1.0
                    Min.
                            :-2.4906
                                       Min.
                                               :-0.5347
                                                          Min.
                                                                  :-0.39237
    1st Qu.:125.8
                                       1st Qu.:-0.4202
                                                           1st Qu.:-0.19598
                    1st Qu.:-1.7290
##
   Median :250.5
                                                          Median :-0.15489
                    Median :-1.5067
                                       Median :-0.3928
           :250.5
                            :-1.5325
                                               :-0.3957
                                                                  :-0.15369
##
    Mean
                    Mean
                                       Mean
                                                          Mean
##
    3rd Qu.:375.2
                    3rd Qu.:-1.2970
                                        3rd Qu.:-0.3621
                                                           3rd Qu.:-0.10827
##
    Max.
           :500.0
                    Max.
                            :-0.7082
                                       Max.
                                               :-0.2900
                                                          Max.
                                                                 : 0.02433
    omega2.theta1
##
##
    Min.
          : 8.971
##
    1st Qu.:12.851
   Median :14.390
##
    Mean
           :14.831
##
    3rd Qu.:16.433
           :24.879
    Max.
```

Bootstrap distributions

if(nboot<200) cat("The number of bootstrap samples is too low to provide good estimates of the confiden
 resboot1<-cond.bin</pre>

```
ypd2<-NULL
for(icol in 1:4) {
  ypd2<-rbind(ypd2,data.frame(rep=resboot1[,1],Param=colnames(resboot1)[(icol+1)],value=resboot1[,(icol+1)]</pre>
ypd2$Param<-factor(ypd2$Param, levels = unique(ypd2$Param))</pre>
ypd2.fix<-ypd2[ypd2$Param %in% unique(ypd2$Param)[1:3],]</pre>
ypd2.iiv<-ypd2[ypd2$Param %in% unique(ypd2$Param)[4],]</pre>
ypd <- rbind(ypd,ypd2)</pre>
par.estim<-c(binary.fit@results@fixed.effects,diag(binary.fit@results@omega)[binary.fit@results@indx.
mean.bootDist<-apply(resboot1, 2, mean)[-c(1)]</pre>
df2<-data.frame(Param=unique(ypd2$Param), mean.boot=mean.bootDist, est.saemix=par.estim, Bootstrap="C
df<-rbind(df,df2)</pre>
  plot.density2<-ggplot(data=ypd2) + geom_density(aes(value,fill="red4"), alpha=0.5) +</pre>
  geom_vline(data=df2,aes(xintercept=est.saemix),colour="red",size=1.2) +
  geom_vline(data=df2,aes(xintercept=mean.boot),colour="blue",size=1.2) +
  theme_bw() + theme(axis.title.x = element_blank(),axis.text.x = element_text(size=9, angle=30, hjus
  facet_wrap(~Param, ncol=2, scales = 'free')
print(plot.density2)
plot.density3<-ggplot(data=ypd) + geom_density(aes(value,fill="red4"), alpha=0.5) +
  geom_vline(data=df,aes(xintercept=est.saemix),colour="red",size=1.2) +
  geom_vline(data=df,aes(xintercept=mean.boot),colour="blue",size=1.2) +
  theme_bw() + theme(axis.title.x = element_blank(),axis.text.x = element_text(size=9, angle=30, hjus
  facet_grid(Bootstrap~Param, scales = 'free')
   facet_wrap(Bootstrap~Param, nrow=2, scales = 'free')
print(plot.density3)
```



Bootstrap results Here we produce a table showing the parameters estimated on the original dataset along with the bootstrap estimates (mean (SD) of the bootstrap distribution) and the 95% CI. The table is produced when the number of bootstrap is higher than 200 in the code below.

```
if(nboot<200) cat("The number of bootstrap samples is too low to provide good estimates of the confiden
  par.estim<-c(binary.fit@results@fixed.effects,diag(binary.fit@results@omega)[binary.fit@results@indx.
  namsd<-paste0("SD.",colnames(binary.fit@results@omega)[binary.fit@results@indx.omega])
  df2<-data.frame(parameter=c(colnames(case.bin)[-c(1)],namsd), saemix=par.estim)
  for(i in 1:2) {
    if(i==1) {
      resboot1<-case.bin
      namboot<-"case"
      } else {
      resboot1<-cond.bin
      namboot <-"cNP"
      }
    ncol<-dim(resboot1)[2]</pre>
    resboot1<-cbind(resboot1, sqrt(resboot1[,ncol]))</pre>
    mean.bootDist<-apply(resboot1, 2, mean)[-c(1)]</pre>
    sd.bootDist<-apply(resboot1, 2, sd)[-c(1)]</pre>
    quant.bootDist<-apply(resboot1[-c(1)], 2, quantile, c(0.025, 0.975))
    11<-paste0(format(mean.bootDist, digits=2)," (",format(sd.bootDist,digits=2, trim=T),")")</pre>
    12<-paste0("[",format(quant.bootDist[1,], digits=2),", ",format(quant.bootDist[2,],digits=2, trim=T
    df2<-cbind(df2, 11, 12)
    i1 < -3 + 2 * (i-1)
    colnames(df2)[i1:(i1+1)] <-paste0(namboot, ".", c("estimate", "CI"))</pre>
  }
  print(df2)
}
##
                                                              case.CI cNP.estimate
                  parameter
                                 saemix case.estimate
## 1
                     theta1 -1.7063992 -1.74 (0.360) [-2.44, -1.077] -1.53 (0.312)
## 2
                     theta2 -0.3864426 -0.40 (0.073) [-0.56, -0.279] -0.40 (0.044)
## 3 beta_treatment.theta2. -0.1477510 -0.16 (0.132) [-0.44, 0.066] -0.15 (0.067)
## 4
              omega2.theta1 16.1090732 17.06 (4.500) [10.36, 28.229] 14.83 (2.816)
## 5
                  SD.theta1 4.0136110 4.10 (0.530) [ 3.22, 5.313] 3.83 (0.360)
             cNP.CI
## 1 [-2.20, -0.98]
## 2 [-0.49, -0.32]
## 3 [-0.29, -0.02]
## 4 [10.01, 21.50]
## 5 [ 3.16, 4.64]
print(xtable(df2), only.contents=TRUE, include.rownames=FALSE)
## % latex table generated in R 4.2.2 by xtable 1.8-4 package
## % Thu Sep 21 09:53:34 2023
## parameter & saemix & case.estimate & case.CI & cNP.estimate & cNP.CI \\
##
     \hline
## theta1 & -1.71 & -1.74 (0.360) & [-2.44, -1.077] & -1.53 (0.312) & [-2.20, -0.98] \\
     theta2 & -0.39 & -0.40 (0.073) & [-0.56, -0.279] & -0.40 (0.044) & [-0.49, -0.32] \\
     beta\_treatment.theta2. & -0.15 & -0.16 (0.132) & [-0.44, 0.066] & -0.15 (0.067) & [-0.29, -0.02]
##
     omega2.theta1 & 16.11 & 17.06 (4.500) & [10.36, 28.229] & 14.83 (2.816) & [10.01, 21.50] \\
##
     SD.theta1 & 4.01 & 4.10 (0.530) & [3.22, 5.313] & 3.83 (0.360) & [3.16, 4.64] \\
##
##
      \hline
```

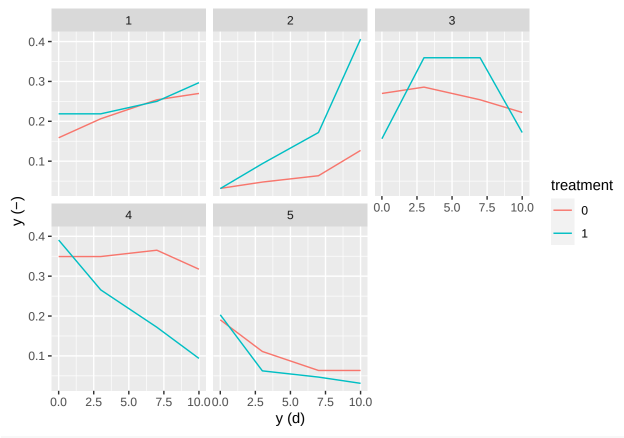
```
# Bootstrap SE on omega instead of omega2
# sd(sqrt(case.bin$omega2.theta1))
# sd(sqrt(cond.bin$omega2.theta1))
```

Categorical response model

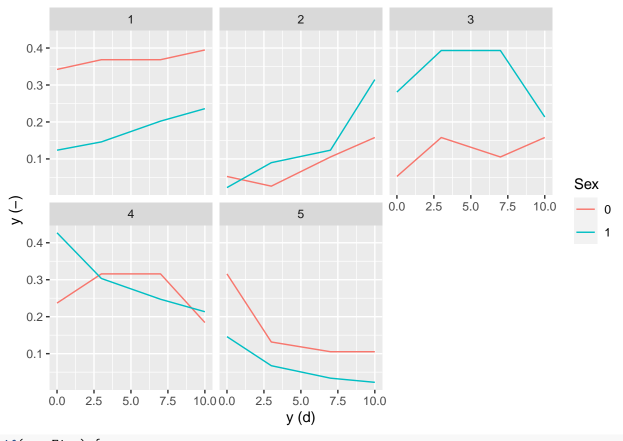
Data The *knee.saemix* data represents pain scores recorded in a clinical study in 127 patients with sport related injuries treated with two different therapies. The pain occurring during knee movement was observed after 3,7 and 10 days of treatment. It was taken from the **catdata** package in R (Schauberger and Tutz 2020) (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)
- Age2 represents the square of centered Age.

We can visualise the evolution of the proportion of each score over time using the *plotDiscreteData()* function with the outcome set to *categorical*, stratifying by treatment. In both treatment groups, the probability of lower scores (1 or 2) increases with time while the probability to obtain a high (4 or 5) pain score decreases.



plotDiscreteData(ordknee.data, outcome="categorical", which.cov="Sex")



```
if(saveFigs) {
  namfig<-"knee_rawDataPropTime.eps"
  cairo_ps(file = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widelight = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widelight = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widelight = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widelight = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widelight = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widelight = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widelight = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widelight = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widelight = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widelight = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widelight = file.path(figDir, namfig), onefile = file.path(file.path(file.path(file.path(file.path(file.path(file.path(file.
```

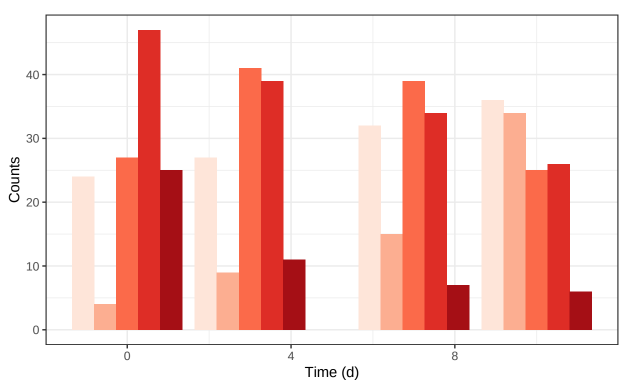
The following R code represents the data as barplots of the different pain scores as a function of time in study, also illustrating a recovery as the proportion of lower pain scores increases. Other graphs can be created to explore this data, such as the evolution of mean scores over time.

```
gtab <- knee.saemix %>%
  group_by(time, y) %>%
  summarise(n=length(y)) %>%
  mutate(y=as.factor(y))

## `summarise()` has grouped output by 'time'. You can override using the
## `.groups` argument.

ggplot(data = gtab, aes(x = time, y=n, group=y, fill=y)) +
  geom_bar(stat="identity", position = "dodge") + theme_bw() +
  scale_fill_brewer(palette = "Reds") + theme(legend.position = "top") +
  labs(fill = "Score") + xlab("Time (d)") + ylab("Counts")
```





```
# Gender effect ?
if(FALSE) {
  gtab2 <- knee.saemix %>%
    group_by(time, Sex) %>%
    summarise(meanscore=mean(y))

ggplot(data = gtab2, aes(x = time, y=meanscore, group=Sex, fill=Sex)) +
    geom_line() + theme_bw() + theme(legend.position = "top") +
    labs(fill = "Score") + xlab("Time (d)") + ylab("Mean score")
}
```

Model The dataset is part of the datasets analysed in (Tutz 2012) with various methods described in the vignettes in the documentation of the *knee* dataset, but mainly as logistic regression on the response after 10 days, or as mixed binary regression after dichotomising the response. Here, we fit a 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:

$$logit(P(Y_{ij} = 1|\psi_{i})) = \theta_{1,i} + \beta_{i}t_{ij}$$

$$logit(P(Y_{ij} = 2|\psi_{i})) = \theta_{1,i} + \theta_{2}$$

$$logit(P(Y_{ij} = 3|\psi_{i})) = \theta_{1,i} + \theta_{2} + \theta_{3}$$

$$logit(P(Y_{ij} = 4|\psi_{i})) = \theta_{1,i} + \theta_{2} + \theta_{3} + \theta_{4}$$

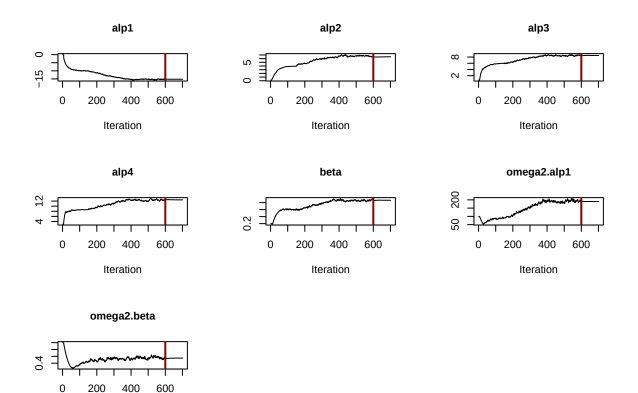
$$P(Y_{ij} = 4|\psi_{i}) = 1 - \sum_{k=1}^{4} P(Y_{ij} = k|\psi_{i})$$
(1)

where θ_1 and β are assumed to have interindividual variability and to follow a normal distribution. β is the effect of time, θ_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.

The following segment of code defines the ordinal model, computing the different logits for the different categories and deriving the corresponding probability given the observed data passed in *xidep*. We first fit a base model without covariate.

```
# Model for ordinal responses
ordinal.model<-function(psi,id,xidep) {</pre>
  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</pre>
  logit3<-logit2+alp3</pre>
  logit4<-logit3+alp4</pre>
  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)</pre>
  return(logpdf)
}
# simulate function
simulateOrdinal<-function(psi,id,xidep) {</pre>
  y < -xidep[,1]
  time<-xidep[,2]
  alp1<-psi[id,1]
  alp2<-psi[id,2]
  alp3<-psi[id,3]</pre>
  alp4<-psi[id,4]
  beta <- psi[id, 5]
  logit1<-alp1 + beta*time</pre>
  logit2<-logit1+alp2</pre>
  logit3<-logit2+alp3</pre>
  logit4<-logit3+alp4</pre>
  pge1<-exp(logit1)/(1+exp(logit1))</pre>
  pge2<-exp(logit2)/(1+exp(logit2))
  pge3<-exp(logit3)/(1+exp(logit3))
  pge4<-exp(logit4)/(1+exp(logit4))
  x<-runif(length(time))</pre>
  ysim<-1+as.integer(x>pge1)+as.integer(x>pge2)+as.integer(x>pge3)+as.integer(x>pge4)
  return(ysim)
}
# Saemix model
saemix.model<-saemixModel(model=ordinal.model,description="Ordinal categorical model",modeltype="likeli")</pre>
                 simulate.function=simulateOrdinal, 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),
```

```
omega.init=diag(c(100, 1, 1, 1, 1)), covariance.model = diag(c(1,0,0,0,1)), verbose=FAL
# Fitting
saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, fim=FALSE, nb.chains=10, nbiter.saemix=c</pre>
\#saemix.options < -list(seed = 632545, save = FALSE, save.graphs = FALSE, \ nb.\ chains = 10, \ fim = FALSE)
ord.fit<-saemix(saemix.model,ordknee.data,saemix.options)</pre>
summary(ord.fit)
## -----
## ----- Fixed effects -----
## -----
   Parameter Estimate
## 1
       alp1
           -15.21
       alp2
## 2
              6.51
      alp3
              8.49
## 4
       alp4
            12.48
## 5
       beta
              0.87
## ----- Variance of random effects -----
       Parameter Estimate
## alp1 omega2.alp1
                189.79
## beta omega2.beta
                 0.55
## -----
## ----- Correlation matrix of random effects -----
## -----
           omega2.alp1 omega2.beta
##
## omega2.alp1 1.00
                    0.00
## omega2.beta 0.00
                     1.00
## ----- Statistical criteria -----
  _____
##
## Likelihood computed by importance sampling
##
      -2LL= 859.7992
      AIC = 875.7992
##
##
      BIC = 898.5527
plot(ord.fit, plot.type="convergence")
```



Iteration

Note: comparable estimates obtained with Monolix (not same, but within CI)
quite a lot of sensitivity to distributions (when using eg normal distributions in Monolix the param

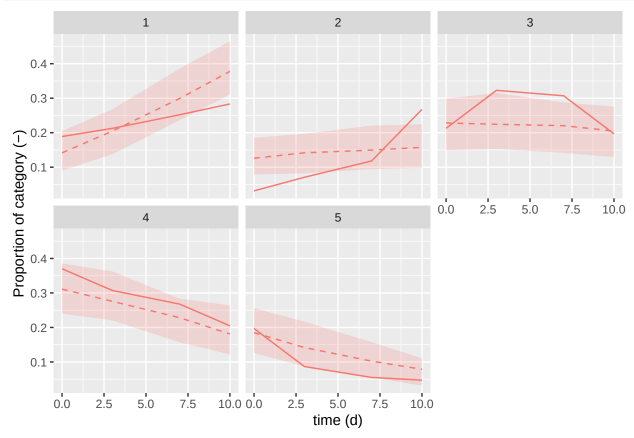
We then applied the stepwise algorithm proposed by Delattre et al. to simultaneously investigate the covariate model and variability structure. This was run outside Rstudio because of memory issues. The resulting model has variability on all the α parameters, and included treatment on α_2 and β and age squared on α_1 . Comparing the two models through the *compare.saemix()* function shows about 30 points improvement in the BIC.

```
# Do not run, Rstudio fails (ran in a script as "R CMD BATCH paper_kneeCovModel.R paper_kneeCovModel.ou
#if(runCovKnee) cov.ordfit <- step.saemix(ord.fit, trace=TRUE, direction='both')</pre>
# Resulting model
## IIV: all alphas, none on beta :-/
## Covariates:
                 alp1(Age2)alp2(treatment)beta(treatment)
covariate.model <- matrix(data=0, nrow=4, ncol=5)</pre>
covariate.model[3,2] < -covariate.model[3,5] < -covariate.model[4,1] < -1
ordmodel.cov<-saemixModel(model=ordinal.model,description="Ordinal categorical model",modeltype="likeli
                simulate.function=simulateOrdinal, 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),
                omega.init=diag(c(100, 1, 1, 1, 1)), covariate.model=covariate.model, covariance.model
# Fitting
saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, fim=FALSE, nb.chains=10, nbiter.saemix=c</pre>
#saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, nb.chains=10, fim=FALSE)
ord.fit.cov<-saemix(ordmodel.cov,ordknee.data,saemix.options)
summary(ord.fit.cov)
```

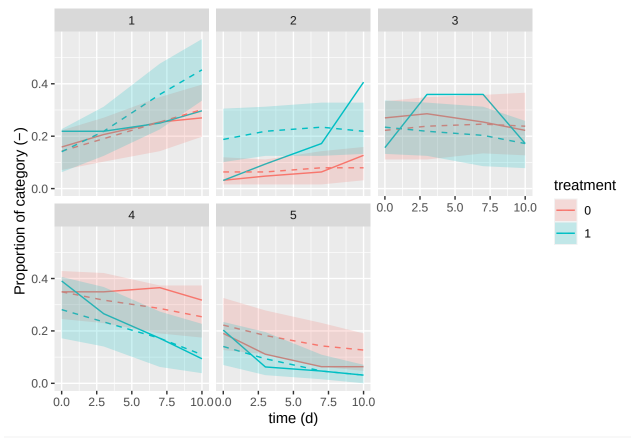
```
----- Fixed effects ------
  _____
##
             Parameter Estimate
## 1
                 alp1
                      -23.349
## 2
        beta_Age2(alp1)
                       0.062
                 alp2
                        3.499
## 4 beta_treatment(alp2)
                       0.977
## 5
                 alp3
                       8.538
## 6
                 alp4
                       16.436
## 7
                 beta
                       0.884
## 8 beta_treatment(beta)
                       0.557
  ----- Variance of random effects -----
##
        Parameter Estimate
## alp1 omega2.alp1
                  244.71
## alp2 omega2.alp2
                   0.18
## alp3 omega2.alp3
                   0.55
## alp4 omega2.alp4
                   0.47
  ----- Correlation matrix of random effects -----
  _____
##
            omega2.alp1 omega2.alp2 omega2.alp3 omega2.alp4
## omega2.alp1 1.00
                      0.00
                                0.00
                                          0.00
## omega2.alp2 0.00
                      1.00
                                0.00
                                          0.00
## omega2.alp3 0.00
                      0.00
                                1.00
                                          0.00
  omega2.alp4 0.00
                      0.00
                                0.00
                                          1.00
   ----- Statistical criteria -----
  ______
##
  Likelihood computed by importance sampling
       -2LL= 807.0315
##
##
       AIC = 833.0315
##
       BIC = 870.0059
  ______
# Compare the base and covariate model
compare.saemix(ord.fit, ord.fit.cov)
## Likelihoods calculated by importance sampling
##
        AIC
               BIC
## 1 875.7992 898.5527
## 2 833.0315 870.0059
```

Model evaluation In the code below we produce VPC plots with the discreteVPC() function (see details above for the binary example) for the model with covariates. The VPC show some model misspecification, especially in the intermediate pain scores, as well as a tendency to overestimate the improvement, driven by the increase in the lowest pain score. This suggests the impact of time and treatment are not well taken into account in the current model. A second plot stratified by treatment also suggests treatment effect is not properly accounted for the three lowest pain categories and across time. Finally, although Sex was not retained in the covariate model, VPC stratified on Sex seem to indicate different distributions of the pain scores between men and women, especially in the first and third categories (no or middle pain levels).

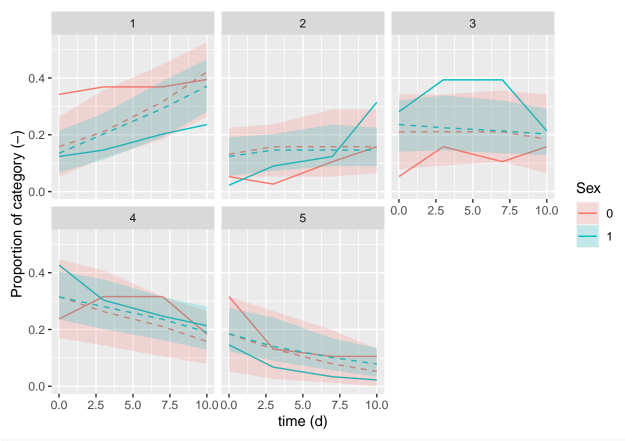
```
### Simulations for VPC
nsim<-100
yfit<-ord.fit.cov
yfit<-simulateDiscreteSaemix(yfit, nsim=nsim)
discreteVPC(yfit, outcome="categorical")</pre>
```



discreteVPC(yfit, outcome='categorical',covsplit=TRUE, which.cov="treatment")



discreteVPC(yfit, outcome='categorical',covsplit=TRUE, which.cov="Sex")



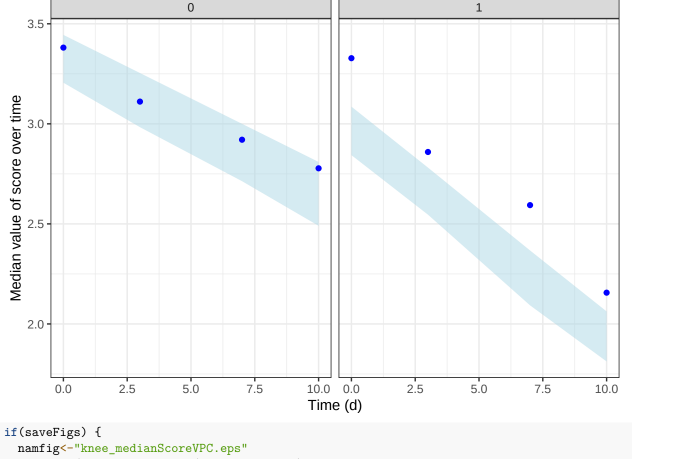
```
if(saveFigs) {
  namfig<-"knee_VPCbytreatment.eps"</pre>
  cairo_ps(file = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, wi
  discreteVPC(yfit, outcome='categorical',covsplit=TRUE, which.cov="treatment")
  dev.off()
  namfig<-"knee_VPCbySex.eps"
  cairo_ps(file = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, wi
  discreteVPC(yfit, outcome='categorical',covsplit=TRUE, which.cov="Sex")
  dev.off()
}
if(FALSE) {
  ### Simulations for VPC - model without covariates
  nsim<-100
  yfit<-ord.fit
  yfit<-simulateDiscreteSaemix(yfit, nsim=nsim)</pre>
  discreteVPC(yfit, outcome="categorical")
}
```

We can also look at the VPC for the median score in each treatment group to find that the model tends to underpredict the pain scores, especially in the group receiving therapy 1.

```
# VPC for median score in each group
knee3 <- knee.saemix %>%
  group_by(time, treatment) %>%
  summarise(mean=mean(y))
```

`summarise()` has grouped output by 'time'. You can override using the

```
## `.groups` argument.
simdat <-yfit@sim.data@datasim</pre>
simdat$time<-rep(yfit@data@data$time,nsim)</pre>
simdat$treatment<-rep(yfit@data@data$treatment,nsim)</pre>
ytab<-NULL
for(irep in 1:nsim) {
  xtab<-simdat[simdat$irep==irep,]</pre>
  suppressMessages(
 xtab1 <- xtab %>%
    group_by(time, treatment) %>%
    summarise(mean=mean(ysim))
 ytab<-rbind(ytab,xtab1[,c("time","treatment","mean")])</pre>
gtab <- ytab %>%
  group_by(time, treatment) %>%
  summarise (lower=quantile (mean, c(0.05)), mean=median (mean), upper=quantile (mean, c(0.95)))
## `summarise()` has grouped output by 'time'. You can override using the
## `.groups` argument.
kneeMedvpc <- ggplot(data = knee3, aes(x = time, y=mean, group=treatment)) +</pre>
  geom_ribbon(data=gtab, aes(x=time, ymin=lower, ymax=upper), alpha=0.5, fill="lightblue") +
  geom point(colour='blue') + theme bw() +
  scale_fill_brewer(palette = "Blues") + theme(legend.position = "top") +
  labs(fill = "Score") + xlab("Time (d)") + ylab("Median value of score over time") + facet_wrap(.~trea
print(kneeMedvpc)
```



```
namfig<-"knee_medianScoreVPC.eps"
cairo_ps(file = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, wiplot(kneeMedvpc)
dev.off()
}</pre>
```

Taken together, these results suggest that the proportional odds model may not be adequate to describe this data. We could investigate multinomial models to allow for different time trends within each value of score, or consider different types of models such as Markov models, but these are currently out of scope in **saemix**.

Estimation errors

Boostrap methods As previously, we can assess parameters uncertainty using bootstrap approaches. Here we load the results from the two bootstrap files prepared beforehand by running the *saemix.bootstrap()* function with 500 simulations. We compute the bootstrap quantiles for the 95% CI, as well as the SD of the bootstrap distribution, corresponding to a normal approximation of the SE.

```
if(runBootstrap) {
   case.ordinal <- saemix.bootstrap(ord.fit.cov, method="case", nboot=nboot)
   cond.ordinal <- saemix.bootstrap(ord.fit.cov, method="conditional", nboot=nboot)
} else {
   case.ordinal <- read.table(file.path(saemixDir, "bootstrap", "results", "knee_caseBootstrap.res"), header
   cond.ordinal <- read.table(file.path(saemixDir, "bootstrap", "results", "knee_condBootstrap.res"), header
   nboot<-dim(case.ordinal)[1]
}
case.ordinal <- case.ordinal[!is.na(case.ordinal[,2]),]</pre>
```

```
cond.ordinal <- cond.ordinal[!is.na(cond.ordinal[,2]),]</pre>
par.estim<-format(c(ord.fit@results@fixed.effects,diag(ord.fit@results@omega)[ord.fit@results@indx.omeg
df2<-data.frame(parameter=colnames(case.ordinal)[-c(1)], saemix=par.estim)
for(i in 1:2) {
     if(i==1) {
          resboot1<-case.ordinal
          namboot<-"case"
     } else {
          resboot1<-cond.ordinal
          namboot <-"cNP"
     }
     mean.bootDist<-apply(resboot1, 2, mean, na.rm=T)[-c(1)]</pre>
     sd.bootDist<-apply(resboot1, 2, sd, na.rm=T)[-c(1)]</pre>
     quant.bootDist<-apply(resboot1[-c(1)], 2, quantile, c(0.025, 0.975), na.rm=T)
     11<-paste0(format(mean.bootDist, digits=2)," (",format(sd.bootDist,digits=2, trim=T),")")</pre>
     12<-paste0("[",format(quant.bootDist[1,], digits=2),", ",format(quant.bootDist[2,],digits=2, trim=T),</pre>
     df2<-cbind(df2, 11, 12)
     i1 < -3 + 2 * (i-1)
     colnames(df2)[i1:(i1+1)] <-paste0(namboot, ".", c("estimate", "CI"))</pre>
print(df2)
##
                  parameter saemix case.estimate
                                                                                                                               case.CI
                                                                                                                                                         cNP.estimate
## 1
                                alp1 -15.21 -16.12 (2.28) [-20.80, -11.70]
                                                                                                                                                    -14.68 (2.01)
## 2
                                                                       7.07 (1.01)
                                                                                                         [ 5.25, 9.22]
                                                                                                                                                            5.97 (0.90)
                               alp2
                                                  6.51
## 3
                                alp3
                                                 8.49
                                                                       8.96 (1.38) [ 6.62, 11.87]
                                                                                                                                                            8.40 (1.06)
## 4
                                alp4 12.48
                                                                                                        [ 9.46, 18.64]
                                                                                                                                                          12.86 (1.91)
                                                                     13.26 (2.42)
## 5
                                                 0.87
                                                                       0.91 (0.11)
                                                                                                          [ 0.70, 1.14]
                                                                                                                                                            0.82 (0.11)
## 6 omega2.alp1 189.79 221.01 (64.76) [117.37, 363.58] 177.78 (48.53)
## 7 omega2.beta
                                                  0.55
                                                                       0.57 (0.18)
                                                                                                        [ 0.26, 0.97]
                                                                                                                                                      0.52 (0.16)
##
                                       cNP.CI
## 1 [-18.55, -11.09]
## 2
              [ 4.49, 7.77]
## 3 [ 6.67, 10.69]
## 4 [ 9.85, 17.34]
               [ 0.63, 1.03]
## 5
## 6 [ 99.82, 289.20]
## 7
                  [0.25, 0.86]
# pretty print for paper
df3 < -df2[,c(2,4)] # case bootstrap
 rownames(df3) < -c("\$\\alpha_1\$", "\$\\alpha_2\$", "\$\\alpha_3\$", "\$\\alpha_4\$", "\$\\beta_4\", "\$\\alpha_3\$", "\$\\alpha_4\$", "\$\\alpha_3\$", "\$\\alpha
print(xtable(df3), only.contents=TRUE, include.rownames=TRUE, floating=F, sanitize.rownames.function =
## \% latex table generated in R 4.2.2 by xtable 1.8-4 package
## % Thu Sep 21 09:55:30 2023
## & saemix & case.CI \\
##
            \hline
## $\alpha_1$ & -15.21 & [-20.80, -11.70] \\
##
             $\alpha_2$ &
                                                    6.51 & [ 5.25, 9.22] \\
##
             $\alpha_3$ &
                                                8.49 & [ 6.62, 11.87] \\
##
             $\alpha_4$ & 12.48 & [ 9.46, 18.64] \\
##
             $\beta$ & 0.87 & [ 0.70, 1.14] \\
```

```
##
    $\omega_{\alpha_1}$ & 189.79 & [117.37, 363.58] \\
##
    $\omega_{\beta}$ &
                    0.55 & [ 0.26, 0.97] \\
##
     \hline
df3<-df2[,c(2,5)] # conditional bootstrap
print(xtable(df3), only.contents=TRUE, include.rownames=TRUE, floating=F, sanitize.rownames.function =
## % latex table generated in R 4.2.2 by xtable 1.8-4 package
## % Thu Sep 21 09:55:30 2023
## & saemix & cNP.estimate \\
##
    \hline
## $\alpha_1$ & -15.21 & -14.68 (2.01) \\
##
    $\alpha_2$ &
                6.51 &
                        5.97 (0.90) \\
##
    $\alpha_3$ &
                8.49 &
                       8.40 (1.06) \\
    $\alpha_4$ & 12.48 & 12.86 (1.91) \\
##
    $\beta$ &
##
              0.87 &
                     0.82 (0.11) \\
##
    $\omega_{\alpha_1}$ & 189.79 & 177.78 (48.53) \\
    \sigma_{\star} \ 0.55 & 0.52 (0.16) \
##
     \hline
##
```

References

Comets E, Rodrigues C, Jullien V, Ursino M (2021). Conditional non-parametric bootstrap for non-linear mixed effect models. *Pharmaceutical Research*, 38: 1057-66.

Delattre M, Lavielle M, Poursat MA (2014) A note on BIC in mixed effects models. *Electronic Journal of Statistics* 8(1) p. 456-475

Delattre M, Poursat MA (2017) BIC strategies for model choice in a population approach. (arXiv:1612.02405)

Schauberger G, Tutz G (2020). catdata: Categorical Data. R package version 1.2.2. https://CRAN.R-project.org/package=catdata