Testing examples in saemix 3.0 - discrete models

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Objective

Check saemix for discrete data models

Setup

- set up work directories
- two versions toggled by testMode
 - if testMode is FALSE, load the functions in R
 - if testMode is TRUE, load the library in a dev_mode environment
- aim: check the examples used in the online documentation
 - all examples must run without error

```
if(testMode) cat("Testing package\n") else cat("Loading functions\n")
```

Loading functions

Testing library

Binary response model

- Toenail data
 - using the full model with 2 random effects (better than with only random effect on intercept according to AIC/BIC)
 - quick diagnostics using a simulation function
- TODO
 - add diagnostics (npd-categorical?)
 - maybe check SE's with package by S. Ueckert

```
## [1] "treatment"
##
```

The following SaemixData object was successfully created:

```
##
## Object of class SaemixData
       longitudinal data for use with the SAEM algorithm
## Dataset toenail.saemix
##
       Structured data: y ~ time + y | id
       X variable for graphs: time ()
##
       covariates: treatment (-)
##
         reference class for covariate treatment : 0
##
# Explore 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(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"))</pre>
plot1<-ggplot(toe1, aes(x=visit, y=freq, group=treatment)) + geom_line(aes(colour=treatment)) +</pre>
  geom_point(aes(colour=treatment)) +
  geom_ribbon(aes(ymin=lower, ymax=upper, fill=treatment), alpha=0.2) +
  vlim(c(0,1)) + theme bw() + theme(legend.position = "top") +
  xlab("Visit number") + ylab("Observed frequency of infection")
# 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)
}
saemix.model<-saemixModel(model=binary.model,description="Binary model",</pre>
                           modeltype="likelihood",
                           psi0=matrix(c(0,-.5,0,0.5),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1","t.
                           transform.par=c(0,0), covariate.model=c(0,1),covariance.model=matrix(c(1,0,0,
##
##
## The following SaemixModel object was successfully created:
## Nonlinear mixed-effects model
##
     Model function: Binary model
##
     Model type: likelihood
## function(psi,id,xidep) {
    tim<-xidep[,1]
```

```
##
    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)
## }
##
    Nb of parameters: 2
##
        parameter names: theta1 theta2
##
        distribution:
       Parameter Distribution Estimated
##
## [1,] theta1 normal
                           Estimated
              normal
## [2,] theta2
                            Estimated
##
   Variance-covariance matrix:
##
        theta1 theta2
            1 0
## theta1
## theta2
             0
##
   Covariate model:
      theta1 theta2
## [1,]
       0
##
      Initial values
##
            theta1 theta2
## Pop.CondInit 0 -0.5
## Cov.CondInit
                        0.5
                   0
saemix.options<-list(seed=1234567,save=FALSE,save.graphs=FALSE, displayProgress=FALSE, nb.chains=10, fix</pre>
# saemix fit
binary.fit <- saemix (saemix.model, saemix.data, saemix.options)
## Nonlinear mixed-effects model fit by the SAEM algorithm
## -----
## ----
           Data
## -----
## Object of class SaemixData
      longitudinal data for use with the SAEM algorithm
## Dataset toenail.saemix
      Structured data: y ~ time + y | id
##
##
      X variable for graphs: time ()
##
      covariates: treatment (-)
##
        reference class for covariate treatment : 0
## Dataset characteristics:
##
      number of subjects:
                            294
##
      number of observations: 1908
      average/min/max nb obs: 6.49 / 1 / 7
## First 10 lines of data:
##
     id
             time y y.1 treatment mdv cens occ ytype
## 1 1 0.0000000 1 1
                           1 0 0
## 2 1 0.8571429 1 1
                              1 0
                                        0 1
## 3
     1 3.5357143 1 1
                               1 0
                                      0
                                          1
## 4 1 4.5357143 0 0
                              1 0 0 1
```

1 0 0 1

5 1 7.5357143 0 0

```
1 10.0357143 0 0
                            1 0
## 6
                             1
                                   0
                                        1
     1 13.0714286 0 0
                                 0
## 8 2 0.0000000 0 0
                            0 0 0 1
## 9 2 0.9642857 0 0
                            0 0
                                   0 1
                                             1
                             0 0
                                    0 1
## 10 2 2.0000000 1 1
## -----
             Model
## -----
## Nonlinear mixed-effects model
    Model function: Binary model
    Model type: likelihood
## function(psi,id,xidep) {
    tim<-xidep[,1]
##
##
    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)
## }
## <bytecode: 0x561398afb298>
##
    Nb of parameters: 2
##
       parameter names: theta1 theta2
##
       distribution:
      Parameter Distribution Estimated
## [1,] theta1 normal Estimated
## [2,] theta2 normal
                       Estimated
##
   Variance-covariance matrix:
##
        theta1 theta2
## theta1
          1
## theta2
            0
##
  Covariate model:
##
          [,1] [,2]
## treatment 0 1
##
      Initial values
##
             theta1 theta2
## Pop.CondInit 0 -0.5
## Cov.CondInit
                0 0.5
## -----
         Key algorithm options ----
##
  _____
      Estimation of individual parameters (MAP)
      Estimation of log-likelihood by importance sampling
##
##
      Number of iterations: K1=300, K2=100
##
      Number of chains: 10
##
      Seed: 1234567
##
      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 a file: FALSE
##
       save the graphs to files: FALSE
## -----
## ----
                  Results
## -----
## ----- Fixed effects -----
## -----
     Parameter
                       Estimate
## [1,] theta1
                      -2.20
## [2,] theta2
                      -1.25
## [3,] beta_treatment(theta2) -0.47
## -----
## ----- Variance of random effects -----
## -----
       Parameter Estimate
## theta1 omega2.theta1 59.3
## theta2 omega2.theta2 1.1
## -----
## ----- Correlation matrix of random effects -----
## -----
##
           omega2.theta1 omega2.theta2
## omega2.theta1 1
## omega2.theta2 0
## -----
## ----- Statistical criteria -----
## -----
##
## Likelihood computed by importance sampling
      -2LL= 1116.755
      AIC = 1128.755
##
##
      BIC = 1150.856
# simulate from model (nsim=100)
simulBinary<-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<-1/(1+exp(-logit))
 ysim<-rbinom(length(tim),size=1, prob=pevent)</pre>
 return(ysim)
}
yfit<-binary.fit
nsim<-100
yfit <- simulateDiscreteSaemix(yfit, simulBinary, nsim=nsim)</pre>
simdat <-yfit@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
```

```
xtab<-simdat[simdat$irep==irep,]</pre>
  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>
}
  `summarise()` has grouped output by 'visit'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'visit'. You can override using the `.groups` argument.
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```

for(irep in 1:nsim) {

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

```
## `summarise()` has grouped output by 'visit'. You can override using the `.groups` argument.
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"))
## `summarise()` has grouped output by 'visit'. You can override using the `.groups` argument.
gtab$freq<-1
plot2 <- ggplot(toe1, aes(x=visit, y=freq, group=treatment)) + geom_line(aes(colour=treatment)) +</pre>
  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)
                                                                       В
                           Α
   0.5
   0.4 -
Frequency of infection
   0.3
   0.2
   0.1
   0.0
                                           Visit number
```

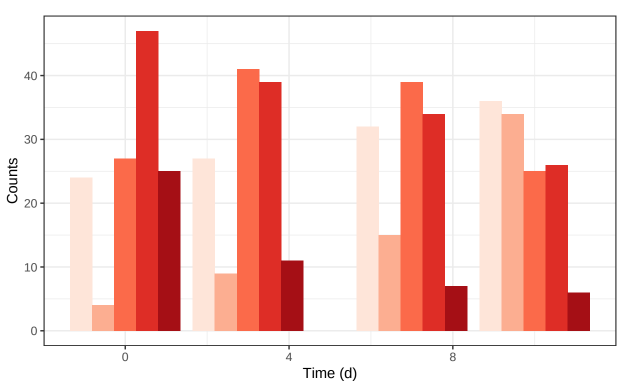
```
if(saveForDocs) {
  namfig<-"toenail_barplotData.eps"
  cairo_ps(file = file.path(figDir, namfig), onefile = TRUE, fallback_resolution = 600, height=8.27, widelight=8.27)
  plot(plot2)
  dev.off()
}</pre>
```

Categorical response model

- Knee pain after 3, 7 and 10 days of treatment compared to baseline (time=0)
 - longitudinal ordinal model with 5 categories
 - similar results to Monolix in terms of parameter estimates
 - SE don't seem so off, but still higher than Monolix (but computed with linearisation anyway)
- Comparing the 3 covariate models model with Age on alp1 and treatment on beta best

```
if(testMode)
  data(knee.saemix) else
   knee.saemix<-read.table(file.path(datDir, "knee.saemix.tab"), header=TRUE)</pre>
# Data
saemix.data<-saemixData(name.data=knee.saemix,name.group=c("id"),</pre>
                        name.predictors=c("y", "time"), name.X=c("time"),
                        name.covariates = c("Age", "Sex", "treatment", "Age2"),
                        units=list(x="d",y="", covariates=c("yr","-","-","yr2")))
## Column name(s) do(es) not exist in the dataset, please check
## Remove columns 1 ( )
## No valid name given, attempting automatic recognition
## Automatic recognition of columns y successful
## [1] "Age"
                   "Sex"
                               "treatment" "Age2"
##
##
## The following SaemixData object was successfully created:
##
## Object of class SaemixData
       longitudinal data for use with the SAEM algorithm
##
## Dataset knee.saemix
##
       Structured data: y ~ y + time | id
       X variable for graphs: time (d)
##
       covariates: Age (yr), Sex (-), treatment (-), Age2 (yr2)
##
         reference class for covariate Sex: 0
##
         reference class for covariate treatment : 0
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")
```





```
# Model for ordinal responses
ordinal.model<-function(psi,id,xidep) {</pre>
  y<-xidep[,1]
  time<-xidep[,2]</pre>
  alp1<-psi[id,1]</pre>
  alp2<-psi[id,2]</pre>
  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))</pre>
  pge3<-exp(logit3)/(1+exp(logit3))</pre>
  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)
}
# Fitting
covmodel2<-covmodel1<-matrix(data=0,ncol=5,nrow=4)</pre>
covmodel1[,1] < -1
```

```
covmodel1[,5]<-1
covmodel2[3,5] < -covmodel2[4,1] < -1
saemix.model<-saemixModel(model=ordinal.model,description="Ordinal categorical model",modeltype="likeli")</pre>
                           psi0=matrix(c(0,0.2, 0.6, 3, 0.2),ncol=5,byrow=TRUE,dimnames=list(NULL,c("alp
                           transform.par=c(0,1,1,1,1),omega.init=diag(rep(1,5)), covariance.model = diag
##
##
## The following SaemixModel object was successfully created:
## Nonlinear mixed-effects model
     Model function: Ordinal categorical model
##
##
     Model type: likelihood
## 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)</pre>
##
##
     return(logpdf)
## }
##
    Nb of parameters: 5
##
         parameter names: alp1 alp2 alp3 alp4 beta
##
         distribution:
##
        Parameter Distribution Estimated
## [1,] alp1
                normal
                               Estimated
## [2,] alp2
                              Estimated
                 log-normal
## [3,] alp3
                  log-normal
                               Estimated
## [4,] alp4
                  log-normal
                               Estimated
                               Estimated
                  log-normal
## [5,] beta
    Variance-covariance matrix:
##
        alp1 alp2 alp3 alp4 beta
## alp1
          1
## alp2
           0
                0
                     0
                          0
                                0
## alp3
           0
                     0
                          0
                                0
## alp4
           0
                     0
                          0
                                0
                0
## beta
##
       No covariate in the model.
```

##

Initial values

```
##
                alp1 alp2 alp3 alp4 beta
                   0 0.2 0.6
                                  3 0.2
## Pop.CondInit
saemix.model.cov1<-saemixModel(model=ordinal.model,description="Ordinal categorical model",modeltype="1</pre>
                              psi0=matrix(c(0,0.2, 0.6, 3, 0.2),ncol=5,byrow=TRUE,dimnames=list(NULL,c(
                              transform.par=c(0,1,1,1,1),omega.init=diag(rep(1,5)), covariance.model =
                               covariate.model = covmodel1)
##
##
## The following SaemixModel object was successfully created:
## Nonlinear mixed-effects model
    Model function: Ordinal categorical model
##
    Model type: likelihood
## 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</pre>
##
     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)</pre>
##
##
##
     return(logpdf)
## }
##
    Nb of parameters: 5
##
         parameter names: alp1 alp2 alp3 alp4 beta
##
         distribution:
##
       Parameter Distribution Estimated
## [1,] alp1
                  normal
                               Estimated
## [2,] alp2
                  log-normal Estimated
## [3,] alp3
                              Estimated
                  log-normal
## [4,] alp4
                  log-normal
                               Estimated
## [5,] beta
                  log-normal
                               Estimated
     Variance-covariance matrix:
##
##
        alp1 alp2 alp3 alp4 beta
## alp1
           1
                     0
## alp2
           0
                     0
                               0
           0
                     0
                          0
                               0
## alp3
                0
## alp4
           0
                0
                     0
## beta
           0
                0
                     0
##
    Covariate model:
##
        alp1 alp2 alp3 alp4 beta
```

```
## [2,]
## [3,]
           1
                          0
                                1
## [4,]
           1
                0
                     0
                           0
                                1
##
       Initial values
##
                alp1 alp2 alp3 alp4 beta
                   0 0.2 0.6
## Pop.CondInit
                                   0.0
## Cov.CondInit
                   0 0.0 0.0
saemix.model.cov2<-saemixModel(model=ordinal.model,description="Ordinal categorical model",modeltype="1
                                psi0=matrix(c(0,0.2, 0.6, 3, 0.2),ncol=5,byrow=TRUE,dimnames=list(NULL,c
                                transform.par=c(0,1,1,1,1),omega.init=diag(rep(1,5)), covariance.model =
                                covariate.model = covmodel2)
##
##
## The following SaemixModel object was successfully created:
##
## Nonlinear mixed-effects model
##
     Model function: Ordinal categorical model
##
     Model type: likelihood
## 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</pre>
##
     logit2<-logit1+alp2
##
     logit3<-logit2+alp3
##
     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)
## }
##
     Nb of parameters: 5
##
         parameter names: alp1 alp2 alp3 alp4 beta
##
         distribution:
##
        Parameter Distribution Estimated
## [1,] alp1
                 normal
                               Estimated
## [2,] alp2
                               Estimated
                  log-normal
## [3,] alp3
                  log-normal
                               Estimated
## [4,] alp4
                  log-normal
                               Estimated
## [5,] beta
                  log-normal
     Variance-covariance matrix:
##
        alp1 alp2 alp3 alp4 beta
##
## alp1
          1 0 0
```

[1,]

1

1

0

0

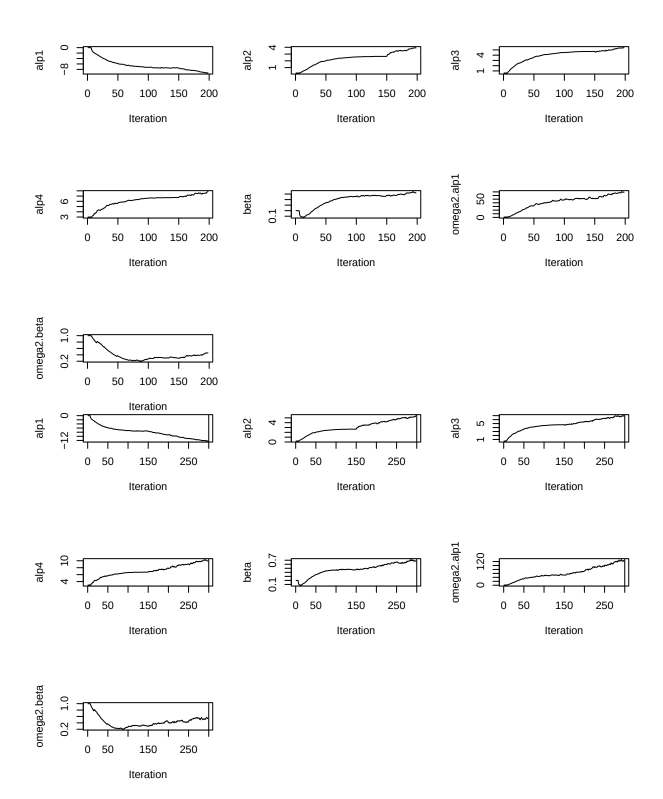
0

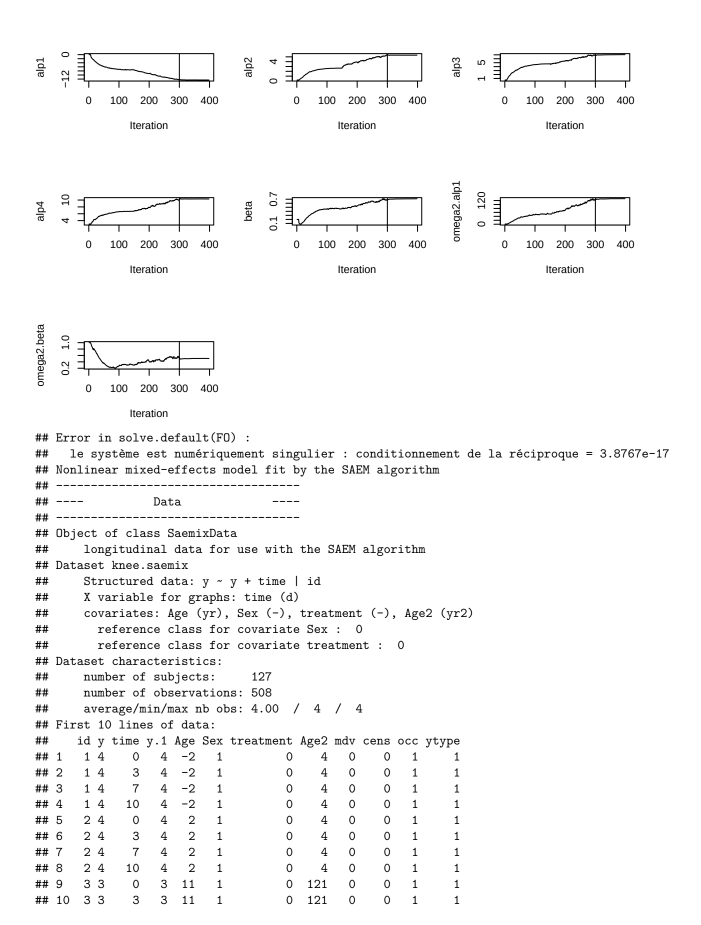
0

0

```
## alp2
                  0
## alp3
            0
                  0
                        0
                              0
                                    0
## alp4
            0
                                    0
## beta
            0
                  0
                        0
                              0
                                    1
##
      Covariate model:
##
         alp1 alp2 alp3 alp4 beta
## [1,]
            0
                  0
                        0
   [2,]
            0
                  0
                        0
                              0
                                    0
##
## [3,]
            0
                  0
                        0
                              0
                                    1
##
   [4,]
            1
                  0
                              0
                                    0
##
        Initial values
##
                  alp1 alp2 alp3 alp4 beta
## Pop.CondInit
                      0 0.2 0.6
## Cov.CondInit
                      0.0 0.0
                                        0.0
saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, nb.chains=10)</pre>
\#saemix.options <-list(seed=632545, save=FALSE, save.graphs=FALSE, nb.chains=10, fim=FALSE)
ord.fit<-saemix(saemix.model,saemix.data,saemix.options)</pre>
    0
alp1
                                       0.5
    ဖှ
        0
            20
                40
                    60
                        80 100
                                           0
                                               20
                                                   40
                                                     60
                                                           80 100
                                                                              0
                                                                                 20
                                                                                      40
                                                                                          60
                                                                                              80 100
                Iteration
                                                  Iteration
                                                                                      Iteration
                                                                     omega2.alp1
                                                                          40
                                   beta
                                                                          0
        0
           20
                40
                    60
                        80 100
                                           0
                                               20
                                                   40
                                                      60
                                                           80 100
                                                                              0
                                                                                 20
                                                                                      40
                                                                                          60
                                                                                              80 100
                Iteration
                                                  Iteration
                                                                                      Iteration
omega2.beta
    1.0
    0.2
        0
            20
                40
                   60
                        80 100
```

Iteration





```
Model
## -----
## Nonlinear mixed-effects model
    Model function: Ordinal categorical model
    Model type: likelihood
##
## 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</pre>
##
    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)</pre>
##
##
    return(logpdf)
## }
## <bytecode: 0x56139956ccb0>
    Nb of parameters: 5
##
        parameter names: alp1 alp2 alp3 alp4 beta
##
        distribution:
##
       Parameter Distribution Estimated
            normal Estimated
## [1,] alp1
             log-normal Estimated log-normal Estimated
## [2,] alp2
## [3,] alp3
## [4,] alp4
               log-normal Estimated
             log-normal
## [5,] beta
                           Estimated
##
   Variance-covariance matrix:
       alp1 alp2 alp3 alp4 beta
##
## alp1 1 0 0 0
        0
                 0
             0
                        0
## alp2
                        0
## alp3
        0
             0
                   0
## alp4 0
                   0
                        0
                             0
              0
                   0
## beta
##
      No covariate in the model.
##
      Initial values
##
              alp1 alp2 alp3 alp4 beta
## Pop.CondInit 0 0.2 0.6 3 0.2
## -----
## ---- Key algorithm options ----
##
      Estimation of individual parameters (MAP)
##
      Estimation of standard errors and linearised log-likelihood
```

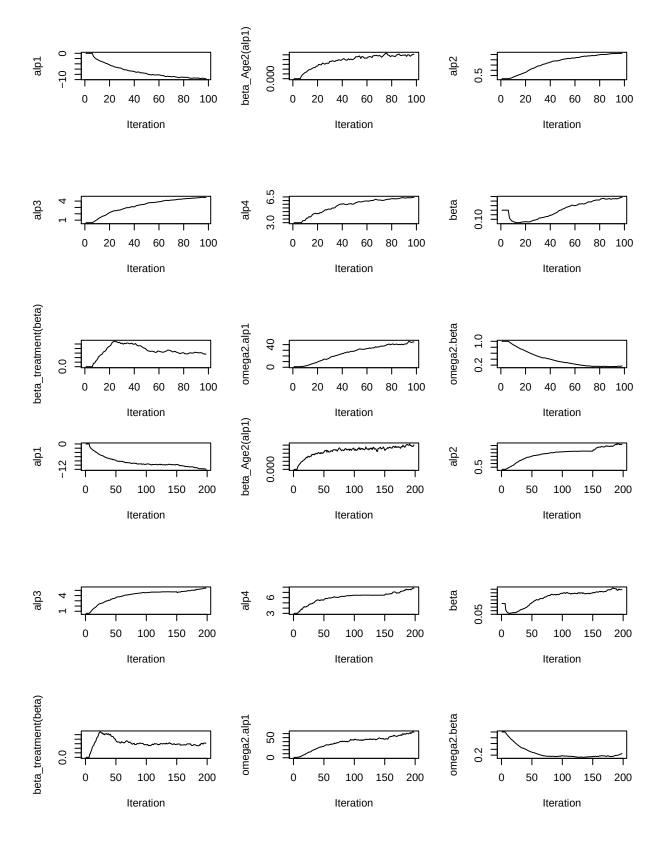
```
##
     Estimation of log-likelihood by importance sampling
##
     Number of iterations: K1=300, K2=100
     Number of chains: 10
##
##
     Seed: 632545
##
     Number of MCMC iterations for IS: 5000
##
     Simulations:
        nb of simulated datasets used for npde: 1000
##
        nb of simulated datasets used for VPC: 100
##
##
     Input/output
##
        save the results to a file: FALSE
        save the graphs to files: FALSE
## -----
                 Results
## -----
## ----- Fixed effects -----
## -----
##
     Parameter Estimate SE CV(%)
## [1,] alp1 -12.47 1.96 16
              5.34 1.93 36
## [2,] alp2
              7.05 1.56 22
## [3,] alp3
## [4,] alp4
             10.31 3.03 29
## [5,] beta
             0.71 0.15 22
## -----
## ----- Variance of random effects -----
## -----
     Parameter Estimate SE CV(%)
## alp1 omega2.alp1 129.61 NA NA
## beta omega2.beta 0.51 NA NA
## ----- Correlation matrix of random effects -----
## -----
##
           omega2.alp1 omega2.beta
## omega2.alp1 1
                    0
## omega2.beta 0
                    1
## ----- Statistical criteria -----
## -----
## Likelihood computed by linearisation
##
      -2LL= 5970.576
##
      AIC = 5986.576
##
      BIC = 6009.33
## Likelihood computed by importance sampling
##
      -2LL= 864.4609
      AIC = 880.4609
##
      BIC = 903.2144
##
ord.fit.cov1<-saemix(saemix.model.cov1, saemix.data, saemix.options)
## Error in plot.new() : figure margins too large
## Error in plot.new() : figure margins too large
## Error in plot.new() : figure margins too large
```

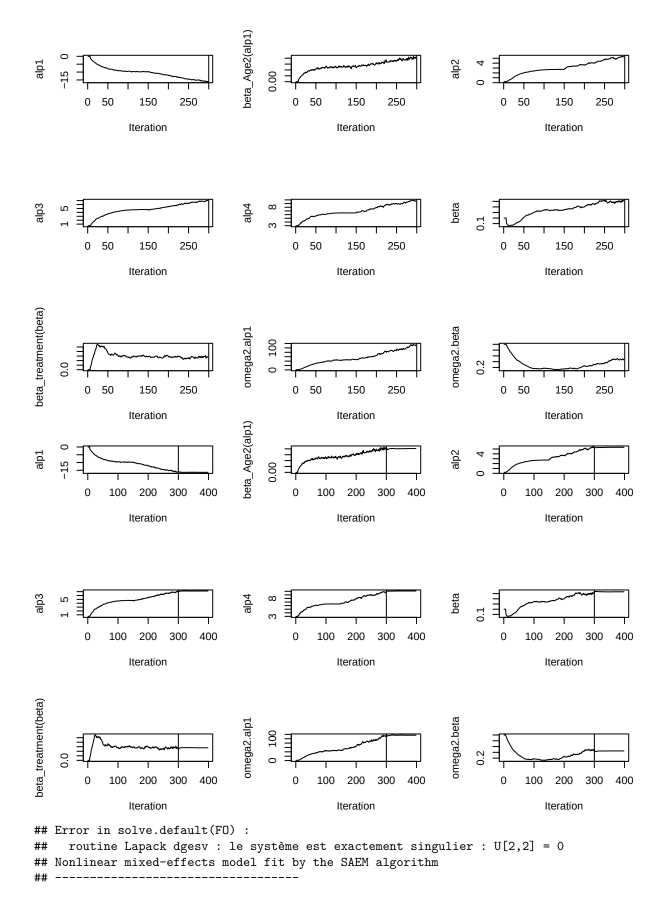
```
## Error in plot.new() : figure margins too large
## Error in solve.default(F0) :
## routine Lapack dgesv : le système est exactement singulier : U[2,2] = 0
## Nonlinear mixed-effects model fit by the SAEM algorithm
## -----
## ----
             Data
## -----
## Object of class SaemixData
##
     longitudinal data for use with the SAEM algorithm
## Dataset knee.saemix
     Structured data: y ~ y + time | id
##
      X variable for graphs: time (d)
      covariates: Age (yr), Sex (-), treatment (-), Age2 (yr2)
##
##
       reference class for covariate Sex : 0
##
       reference class for covariate treatment : 0
## Dataset characteristics:
##
     number of subjects:
                          127
##
      number of observations: 508
##
     average/min/max nb obs: 4.00 / 4 / 4
## First 10 lines of data:
##
     id y time y.1 Age Sex treatment Age2 mdv cens occ ytype
    1 4
           0 4 -2 1
                              0
                                  4 0
## 2
    1 4
           3 4 -2 1
                                      0
                               0
                                           Ω
                                              1
## 3
     1 4
           7 4 -2
                     1
                              0
                                     0
                                              1
## 4
     1 4 10 4 -2 1
                             0
                                 4 0
                                           0 1
## 5 2 4 0 4 2 1
                             0
                                 4 0
## 6 2 4 3 4 2
                     1
                             0
                                 4 0
                                           0 1
                                                   1
## 7
     2 4
          7 4 2
                             0
                                  4
                                     0
                                         0 1
                     1
                                                  1
## 8 2 4 10 4 2 1
                             0
                                 4 0
                                        0 1
                                                  1
## 9 3 3 0 3 11 1
                             0 121
                                      0 0 1
                                                  1
                            0 121
## 10 3 3 3 11 1
                                      0 0 1
                                                    1
          Model
## -----
## Nonlinear mixed-effects model
   Model function: Ordinal categorical model
    Model type: likelihood
## 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))</pre>
##
    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)*(pge3 - pge3)*(pge3 - pge3
##
          logpdf <- log(pobs)</pre>
##
##
          return(logpdf)
## }
## <bytecode: 0x56139956ccb0>
          Nb of parameters: 5
##
                   parameter names: alp1 alp2 alp3 alp4 beta
##
                  distribution:
##
                 Parameter Distribution Estimated
## [1,] alp1
                                  normal Estimated
                                     log-normal Estimated
## [2,] alp2
                                                              Estimated
## [3,] alp3
                                     log-normal
## [4,] alp4
                                     log-normal
                                                                  Estimated
                              log-normal
## [5,] beta
                                                               Estimated
##
          Variance-covariance matrix:
##
                 alp1 alp2 alp3 alp4 beta
## alp1
                 1
                                           0
## alp2
                       0
                                  0
                                             0
                                                       0
## alp3
                      0
                                  0
                                             0
                                                       0
                                                                  0
## alp4
                   0
                                  0
                                             Λ
                                                       0
                                                                  Λ
## beta
                  0
                                  0
##
         Covariate model:
                       [,1] [,2] [,3] [,4] [,5]
##
## Age
                              1
                                             0
                                                       0
## Sex
                                  1
                                             0
                                                       0
                                                                             1
                                             0
                                                       0
                                                                 0
## treatment
                                  1
                                                                             1
                                                                             1
## Age2
                                  1
                                             0
##
              Initial values
##
                                 alp1 alp2 alp3 alp4 beta
## Pop.CondInit
                                       0 0.2 0.6
## Cov.CondInit
                                       0 0.0 0.0
                                                                        0.0
## psi1
                                      0 0.0 0.0
                                                                        0.0
                                       0.0 0.0
                                                                         0.0
## psi1
## psi1
                                       0 0.0 0.0
                                                                        0.0
## -----
## ---- Key algorithm options ----
## -----
##
              Estimation of individual parameters (MAP)
##
              Estimation of standard errors and linearised log-likelihood
              Estimation of log-likelihood by importance sampling
##
##
              Number of iterations: K1=300, K2=100
              Number of chains: 10
##
##
              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 a file: FALSE
##
                       save the graphs to files: FALSE
                                                    Results
## -----
```

```
## ------ Fixed effects ------
## -----
##
      Parameter
                     Estimate SE
                                 CV(%) p-value
## [1,] alp1
                     -1.6e+01 3.4778 22 -
## [2,] beta_Age(alp1)
                     1.2e-01 0.1171
                                  95 0.146
                  -6.9e-01 2.5495 370 0.394
## [3,] beta Sex(alp1)
## [4,] beta_treatment(alp1) 1.7e+00 2.1382 124 0.210
## [5,] beta_Age2(alp1)
                     3.4e-02 0.0166 49 0.021
## [6,] alp2
                      5.2e+00 1.8170 35 -
## [7,] alp3
                     6.9e+00 1.5416 22 -
## [8,] alp4
                      9.8e+00 2.8741 29 -
## [9,] beta
                      4.8e-01 0.2935 61 -
## [10,] beta_Age(beta)
                     -1.6e-02 0.0229 142 0.240
                  3.4e-02 0.4957 1446 0.472
## [11,] beta_Sex(beta)
## [12,] beta_treatment(beta) 5.1e-01 0.4301 85 0.119
## [13,] beta_Age2(beta) 7.4e-04 0.0028 381 0.397
## ----- Variance of random effects -----
## -----
     Parameter Estimate SE CV(%)
## alp1 omega2.alp1 108.43 NA NA
## beta omega2.beta 0.41 NA NA
## -----
## ----- Correlation matrix of random effects -----
## -----
          omega2.alp1 omega2.beta
## omega2.alp1 1
                   0
## omega2.beta 0
                   1
## ------ Statistical criteria ------
## -----
## Likelihood computed by linearisation
##
     -2LL= 5958.536
##
      AIC = 5990.536
##
      BIC = 6036.043
##
## Likelihood computed by importance sampling
##
      -2LL= 840.4144
##
      AIC = 872.4144
##
      BIC = 917.9213
## -----
```

ord.fit.cov2<-saemix(saemix.model.cov2,saemix.data,saemix.options)





```
Data
## -----
## Object of class SaemixData
              longitudinal data for use with the SAEM algorithm
## Dataset knee.saemix
##
              Structured data: y ~ y + time | id
              X variable for graphs: time (d)
              covariates: Age (yr), Sex (-), treatment (-), Age2 (yr2)
##
##
                  reference class for covariate Sex : 0
##
                  reference class for covariate treatment : 0
## Dataset characteristics:
##
              number of subjects:
                                                               127
##
              number of observations: 508
              average/min/max nb obs: 4.00 / 4 / 4
##
## First 10 lines of data:
            id y time y.1 Age Sex treatment Age2 mdv cens occ ytype
            1 4
                        0 4 -2 1
## 1
                                                              0
                                                                                  4 0
## 2
            1 4
                       3 4 -2 1
                                                                          0
## 3
            1 4
                          7 4 -2 1
                                                                      0
                                                                                4 0
                                                                                                       0
                                                                                                            1
            1 4 10 4 -2 1
                                                                                4 0
## 4
                                                                        0
                                                                                                      0
                                                                     0
          2 4
## 5
                       0 4 2 1
                                                                               4 0
                                                                                                  0 1
## 6 2 4 3 4 2 1
                                                                      0
## 7 2 4 7 4 2 1
                                                                                4 0 0 1
                                                                       0
           2 4 10 4 2 1
## 8
                                                                        0
                                                                                 4 0
                                                                                                  0 1
## 9 3 3 0 3 11 1
                                                                      0 121 0 0 1
## 10 3 3 3 11 1
                                                                      0 121 0 0 1
                       Model
## Nonlinear mixed-effects model
##
          Model function: Ordinal categorical model
##
         Model type: likelihood
## 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</pre>
          logit2<-logit1+alp2
##
##
          logit3<-logit2+alp3
          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)+(pge4 - pge3)+(pge4 - pge3)+(pge4 - pge4)+(pge4 - pge4
##
##
          logpdf <- log(pobs)</pre>
##
##
         return(logpdf)
## }
```

```
## <bytecode: 0x56139956ccb0>
##
   Nb of parameters: 5
##
      parameter names: alp1 alp2 alp3 alp4 beta
##
      distribution:
      Parameter Distribution Estimated
## [1,] alp1
          normal Estimated
## [2,] alp2
            log-normal Estimated
            log-normal Estimated
## [3,] alp3
           log-normal Estimated
## [4,] alp4
## [5,] beta
             log-normal Estimated
## Variance-covariance matrix:
      alp1 alp2 alp3 alp4 beta
## alp1 1 0 0 0
      0
          0 0 0 0
## alp2
## alp3 0 0 0 0 0
## alp4
      0
           0
                0
                    0
## beta 0
           0
               0 0
## Covariate model:
    [,1] [,2] [,3] [,4] [,5]
## treatment 0 0 0 0 1
## Age2
           1
                0 0 0 0
## Initial values
##
            alp1 alp2 alp3 alp4 beta
## Pop.CondInit 0 0.2 0.6 3 0.2
## Cov.CondInit 0 0.0 0.0
                         0 0.0
## psi1
         0 0.0 0.0 0 0.0
## -----
## ---- Key algorithm options ----
     Estimation of individual parameters (MAP)
##
     Estimation of standard errors and linearised log-likelihood
##
     Estimation of log-likelihood by importance sampling
##
     Number of iterations: K1=300, K2=100
##
     Number of chains: 10
     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 a file: FALSE
        save the graphs to files: FALSE
## -----
## ----
                  Results
## -----
## ----- Fixed effects -----
## -----
## Parameter
## [1,] alp1
                      Estimate SE CV(%) p-value
                     -16.300 2.406 15
                      0.041 0.014 33
## [2,] beta_Age2(alp1)
                                       0.0014
## [3,] alp2
                       5.340 1.814 34
                       7.173 1.587 22
## [4,] alp3
                      10.079 3.010 30
## [5,] alp4
                       0.535 0.176 33
## [6,] beta
```

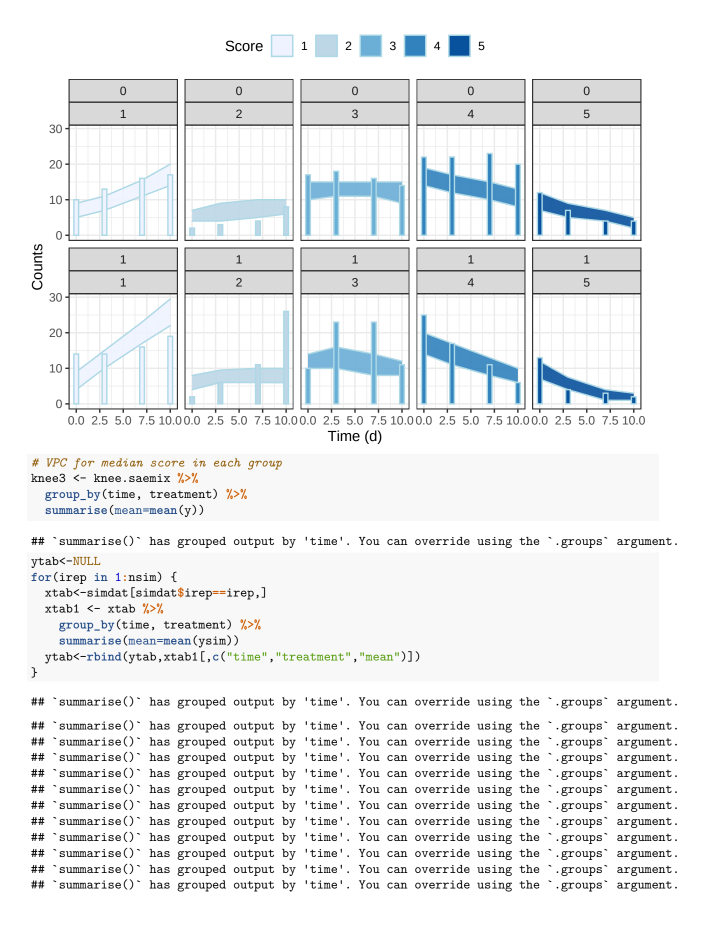
```
## [7,] beta_treatment(beta) 0.554 0.347 63 0.0552
## -----
## ----- Variance of random effects -----
## -----
      Parameter Estimate SE CV(%)
## alp1 omega2.alp1 116.22 NA NA
## beta omega2.beta 0.45 NA NA
## -----
## ----- Correlation matrix of random effects -----
## -----
           omega2.alp1 omega2.beta
              0
## omega2.alp1 1
## omega2.beta 0
                     1
## -----
## ------ Statistical criteria -----
## -----
## Likelihood computed by linearisation
      -2LL= 5980.122
##
      AIC = 6000.122
      BIC = 6028.564
##
##
## Likelihood computed by importance sampling
##
      -2LL= 843.57
      AIC = 863.57
##
      BIC = 892.0119
BIC(ord.fit)
## [1] 903.2144
BIC(ord.fit.cov1)
## [1] 917.9213
BIC(ord.fit.cov2)
## [1] 892.0119
# Comparing the 3 covariate models - model with Age2 on alp1 and treatment on beta best
compare.saemix(ord.fit, ord.fit.cov1, ord.fit.cov2)
## Likelihoods calculated by importance sampling
              BIC BIC.cov
       AIC
## 1 880.4609 903.2144 892.8407
## 2 872.4144 917.9213 907.5477
## 3 863.5700 892.0119 881.6382
###################
# But VPC not good
### Simulations for VPC
simulateOrdinal<-function(psi,id,xidep) {</pre>
 y < -xidep[,1]
 time<-xidep[,2]
 alp1<-psi[id,1]
 alp2<-psi[id,2]
 alp3<-psi[id,3]
```

```
logit1<-alp1 + beta*time</pre>
  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))
  x<-runif(length(time))</pre>
  ysim<-1+as.integer(x>pge1)+as.integer(x>pge2)+as.integer(x>pge3)+as.integer(x>pge4)
  return(vsim)
nsim<-100
yfit<-ord.fit.cov2
yfit<-simulateDiscreteSaemix(yfit, simulateOrdinal, nsim=nsim)
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>
  xtab1 <- xtab %>%
    group_by(time, treatment, ysim) %>%
    summarise(n=length(ysim))
  ytab<-rbind(ytab,xtab1[,c("time","ysim","n","treatment")])</pre>
}
## `summarise()` has grouped output by 'time', 'treatment'. You can override using the `.groups` argume
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```

alp4<-psi[id,4]
beta<-psi[id,5]</pre>

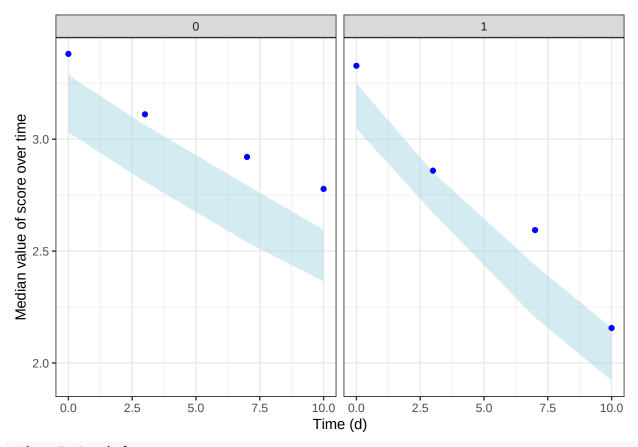
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gtab <- ytab %>%
  group_by(time, treatment, ysim) %>%
  summarise(lower=quantile(n, c(0.05)), n=quantile(n, c(0.5)), upper=quantile(n, c(0.95))) %>%
  mutate(y=as.factor(ysim))
## `summarise()` has grouped output by 'time', 'treatment'. You can override using the `.groups` argume
knee2 <- knee.saemix %>%
  group_by(time, treatment, y) %>%
  summarise(n=length(y)) %>%
  mutate(y=as.factor(y))
## `summarise()` has grouped output by 'time', 'treatment'. You can override using the `.groups` argume
kneevpc <- ggplot(data = knee2, aes(x = time, y=n, fill=y, group=treatment)) +</pre>
  geom_ribbon(data=gtab, aes(x=time, ymin=lower, ymax=upper), alpha=0.9, colour="lightblue") +
  geom_col(position = "dodge", width=0.5, colour="lightblue") + theme_bw() +
  scale fill brewer(palette = "Blues") + theme(legend.position = "top") +
  labs(fill = "Score") + xlab("Time (d)") + ylab("Counts") + facet wrap(treatment~y, nrow=2)
print(kneevpc)
```



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



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

Count data model

- Epilepsy
 - dataset epil from MASS
 - very basic model with only one parameter
- Drinking patterns amongst students (David Atkins from tutorial)
 - dataset rapi.saemix
 - lambda parameter from Poisson model with a time-effect, gender effects on both intercept and slope
 - different models can be adjusted to the data, accounting for overdispersion

Epilepsy data

```
## [1] "trt" "base" "age"
##
##
## The following SaemixData object was successfully created:
##
## Object of class SaemixData
       longitudinal data for use with the SAEM algorithm
##
## Dataset epilepsy
##
       Structured data: y ~ period + y | subject
##
       X variable for graphs: period (2-week)
##
       covariates: trt (), base (), age (yr)
##
          reference class for covariate trt : placebo
## Poisson model with one parameter
countmodel.poisson<-function(psi,id,xidep) {</pre>
  y < -xidep[,2]
  lambda<-psi[id,1]</pre>
  logp <- -lambda + y*log(lambda) - log(factorial(y))</pre>
  return(logp)
}
# Adding a period effect
countmodel.periodpoi<-function(psi,id,xidep) {</pre>
  tim <- xidep[,1]</pre>
  y < -xidep[,2]
  lam<-psi[id,1]</pre>
  betaT<-psi[id,2]
  lambda<-lam*exp(beta*log(tim))</pre>
  logp <- -lambda + y*log(lambda) - log(factorial(y))</pre>
  return(logp)
}
## Generalised Poisson model
countmodel.genpoisson<-function(psi,id,xidep) {</pre>
  y < -xidep[,1]
  delta<-psi[id,1]
  lambda<-psi[id,2]</pre>
  logp <- -lambda
  pos.ind <- which(y>0)
  lp1 <-log(lambda) + (y-1)*log(lambda+y*delta) - (lambda+y*delta) - log(factorial(y))</pre>
  logp[pos.ind] <- lp1[pos.ind]</pre>
  return(logp)
}
## Poisson model wtih Zero-Inflation
countmodel.zip<-function(psi,id,xidep) {</pre>
  y < -xidep[,2]
  lambda<-psi[id,1]</pre>
  p0<-psi[id,2]
  logp <- log(1-p0) -lambda + y*log(lambda) - log(factorial(y))</pre>
  logp0 \leftarrow log(p0+(1-p0)*exp(-lambda))
  logp[y==0] < -logp0[y==0]
  return(logp)
}
```

```
saemix.model.poi<-saemixModel(model=countmodel.poisson,description="count model Poisson",modeltype="lik</pre>
                           psi0=matrix(c(0.5),ncol=1,byrow=TRUE,dimnames=list(NULL, c("lambda"))),
                           transform.par=c(1))
##
##
## The following SaemixModel object was successfully created:
##
## Nonlinear mixed-effects model
     Model function: count model Poisson
##
     Model type: likelihood
##
## function(psi,id,xidep) {
     y < -xidep[,2]
##
##
     lambda<-psi[id,1]</pre>
##
     logp <- -lambda + y*log(lambda) - log(factorial(y))</pre>
##
     return(logp)
## }
##
     Nb of parameters: 1
##
         parameter names: lambda
##
         distribution:
        Parameter Distribution Estimated
##
## [1,] lambda
                  log-normal
                                Estimated
     Variance-covariance matrix:
##
##
          lambda
## lambda
##
       No covariate in the model.
##
       Initial values
##
                lambda
## Pop.CondInit
saemix.model.zip<-saemixModel(model=countmodel.zip,description="count model ZIP",modeltype="likelihood"</pre>
                               psi0=matrix(c(0.5,0.2),ncol=2,byrow=TRUE,dimnames=list(NULL, c("lambda","
                               transform.par=c(1,3), \#omega.init=matrix(c(0.5,0,0,0.3),ncol=2,byrow=TRUE)
                               covariance.model=matrix(c(1,0,0,0),ncol=2,byrow=TRUE))
##
##
## The following SaemixModel object was successfully created:
##
## Nonlinear mixed-effects model
     Model function: count model ZIP
##
     Model type: likelihood
##
## function(psi,id,xidep) {
##
     y < -xidep[,2]
##
     lambda<-psi[id,1]</pre>
##
     p0<-psi[id,2]
##
     logp <- log(1-p0) -lambda + y*log(lambda) - log(factorial(y))</pre>
     logp0 \leftarrow log(p0+(1-p0)*exp(-lambda))
##
##
     logp[y==0] < -logp0[y==0]
##
     return(logp)
## }
##
     Nb of parameters: 2
##
         parameter names: lambda p0
```

##

distribution:

```
Parameter Distribution Estimated
## [1,] lambda
                log-normal Estimated
## [2,] p0
                  logit
                                Estimated
     Variance-covariance matrix:
##
##
          lambda p0
## lambda
              1 0
               0 0
## p0
##
       No covariate in the model.
##
       Initial values
##
                lambda p0
## Pop.CondInit
                   0.5 0.2
saemix.model.gp<-saemixModel(model=countmodel.zip,description="Generalised Poisson model",modeltype="li</pre>
                               psi0=matrix(c(0.5,0.2),ncol=2,byrow=TRUE,dimnames=list(NULL, c("delta","1
                               {\tt transform.par=c(1,1),\ \textit{\#omega.init=matrix}(c(0.5,0,0,0.3),\textit{ncol=2},\textit{byrow=TRUE})}
                               covariance.model=matrix(c(1,0,0,0),ncol=2,byrow=TRUE))
##
##
## The following SaemixModel object was successfully created:
## Nonlinear mixed-effects model
##
     Model function: Generalised Poisson model
     Model type: likelihood
##
## function(psi,id,xidep) {
     y<-xidep[,2]
##
##
     lambda<-psi[id,1]</pre>
##
     p0<-psi[id,2]
     logp <- log(1-p0) -lambda + y*log(lambda) - log(factorial(y))</pre>
##
##
     logp0 \leftarrow log(p0+(1-p0)*exp(-lambda))
     logp[y==0] < -logp0[y==0]
##
##
     return(logp)
## }
##
     Nb of parameters: 2
##
         parameter names: delta lambda
##
         distribution:
##
        Parameter Distribution Estimated
## [1,] delta
                  log-normal
                                Estimated
## [2,] lambda
                  log-normal
                                Estimated
     Variance-covariance matrix:
##
          delta lambda
## delta
              1
## lambda
              0
                      0
##
       No covariate in the model.
##
       Initial values
##
                delta lambda
## Pop.CondInit 0.5
                          0.2
saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, displayProgress=FALSE)</pre>
poisson.fit<-saemix(saemix.model.poi,saemix.data,saemix.options)</pre>
## Nonlinear mixed-effects model fit by the SAEM algorithm
## ----
```

Data

```
## Object of class SaemixData
      longitudinal data for use with the SAEM algorithm
## Dataset epilepsy
      Structured data: y ~ period + y | subject
##
      X variable for graphs: period (2-week)
      covariates: trt (), base (), age (yr)
       reference class for covariate trt : placebo
##
## Dataset characteristics:
##
      number of subjects:
                           59
      number of observations: 236
      average/min/max nb obs: 4.00 / 4 / 4
##
## First 10 lines of data:
     subject period y y.1
                         trt base age mdv cens occ ytype
          1
               1 5 5 placebo
                               11 31
               2 3 3 placebo
## 2
          1
                                11 31
## 3
         1
              3 3 3 placebo
                               11 31
                                       0
                                              1
## 4
         1
               4 3 3 placebo
                               11 31 0
## 5
         2
              1 3 3 placebo
                               11 30
                                       0 0 1
         2 2 5 5 placebo
2 3 3 3 placebo
## 6
                               11 30
                                       0
         2
## 7
                               11 30
                                       0 0 1
## 8
         2
               4 3 3 placebo
                                11 30
                                       0 0 1
## 9
         3
               1 2 2 placebo
                               6 25
                                        0 0 1
                                                    1
       3
              2 4 4 placebo
                               6 25 0 0 1
## -----
             Model
## -----
## Nonlinear mixed-effects model
    Model function: count model Poisson
    Model type: likelihood
## function(psi,id,xidep) {
##
    y < -xidep[,2]
##
    lambda<-psi[id,1]</pre>
##
    logp <- -lambda + y*log(lambda) - log(factorial(y))</pre>
##
    return(logp)
## }
## <bytecode: 0x561396fc8028>
##
    Nb of parameters: 1
##
       parameter names: lambda
##
       distribution:
      Parameter Distribution Estimated
## [1,] lambda log-normal Estimated
   Variance-covariance matrix:
##
        lambda
## lambda
      No covariate in the model.
##
##
      Initial values
##
             lambda
## Pop.CondInit 0.5
## -----
## ---- Key algorithm options ----
## -----
##
     Estimation of individual parameters (MAP)
##
     Estimation of standard errors and linearised log-likelihood
```

```
##
     Estimation of log-likelihood by importance sampling
##
    Number of iterations: K1=300, K2=100
##
    Number of chains: 1
    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 a file: FALSE
       save the graphs to files: FALSE
## -----
                Results
## -----
## ----- Fixed effects -----
## -----
##
     Parameter Estimate SE CV(%)
## [1,] lambda 5.1 0.71 14
## -----
## ----- Variance of random effects -----
## -----
                Estimate SE CV(%)
      Parameter
## lambda omega2.lambda 0.9 0.21 23
## -----
## ----- Correlation matrix of random effects -----
## -----
##
            omega2.lambda
## omega2.lambda 1
## ----- Statistical criteria ------
## -----
## Likelihood computed by linearisation
##
     -2LL= 60096.92
##
     AIC = 60102.92
##
     BIC = 60109.15
##
## Likelihood computed by importance sampling
      -2LL= 1402.095
##
##
      AIC = 1408.095
##
      BIC = 1414.327
## -----
genpoisson.fit<-saemix(saemix.model.gp,saemix.data,saemix.options)</pre>
## Nonlinear mixed-effects model fit by the SAEM algorithm
## -----
## ----
           Data
## -----
## Object of class SaemixData
##
     longitudinal data for use with the SAEM algorithm
## Dataset epilepsy
     Structured data: y ~ period + y | subject
##
     X variable for graphs: period (2-week)
##
##
     covariates: trt (), base (), age (yr)
      reference class for covariate trt : placebo
```

```
## Dataset characteristics:
##
     number of subjects:
                          59
##
      number of observations: 236
      average/min/max nb obs: 4.00 / 4 / 4
##
## First 10 lines of data:
     subject period y y.1 trt base age mdv cens occ ytype
         1
               1 5 5 placebo
                               11 31
               2 3
## 2
          1
                     3 placebo
                                11 31
                                        0
                                            0
                                                1
               3 3 3 placebo
## 3
         1
                                11 31
                                       0
                                            0
                                                1
               4 3 3 placebo
## 4
         1
                                11 31 0 0 1
## 5
         2
               1 3 3 placebo
                                11 30 0 0 1
         2 2 5 5 placebo
2 3 3 3 placebo
                                11 30
                                            0 1
## 6
                                       0
                                                    1
## 7
                                11 30
                                       0 0 1
               4 3 3 placebo
## 8
         2
                                11 30 0 0 1
## 9
         3
               1 2
                     2 placebo
                               6 25
                                       0 0 1
                                                    1
             2 4 4 placebo
       3
## 10
                                 6 25
                                        0 0 1
                                                     1
            Model
## -----
## Nonlinear mixed-effects model
##
    Model function: Generalised Poisson model
    Model type: likelihood
## function(psi,id,xidep) {
    y<-xidep[,2]
##
##
    lambda<-psi[id,1]
    p0<-psi[id,2]
##
    logp <- log(1-p0) -lambda + y*log(lambda) - log(factorial(y))</pre>
    logp0 \leftarrow log(p0+(1-p0)*exp(-lambda))
    logp[y==0] < -logp0[y==0]
##
##
    return(logp)
## }
## <bytecode: 0x56139d720350>
##
    Nb of parameters: 2
##
       parameter names: delta lambda
##
       distribution:
      Parameter Distribution Estimated
## [1,] delta log-normal Estimated
## [2,] lambda log-normal
                         Estimated
    Variance-covariance matrix:
##
        delta lambda
## delta 1 0
## lambda
           0
     No covariate in the model.
##
      Initial values
             delta lambda
## Pop.CondInit 0.5 0.2
## -----
         Key algorithm options ----
## -----
##
      Estimation of individual parameters (MAP)
##
      Estimation of standard errors and linearised log-likelihood
##
      Estimation of log-likelihood by importance sampling
##
     Number of iterations: K1=300, K2=100
     Number of chains: 1
##
```

```
##
    Seed: 632545
##
    Number of MCMC iterations for IS: 5000
##
##
       nb of simulated datasets used for npde: 1000
##
       nb of simulated datasets used for VPC: 100
    Input/output
##
##
       save the results to a file: FALSE
##
       save the graphs to files: FALSE
## -----
                 Results
 -----
## ------ Fixed effects ------
## -----
     Parameter Estimate SE CV(%)
## [1,] delta
          5.314 0.747 14
## [2,] lambda
           0.041
                 0.024 58
## ----- Variance of random effects -----
## -----
      Parameter
               Estimate SE CV(%)
## delta omega2.delta 0.86 0.21 24
## -----
## ----- Correlation matrix of random effects -----
## -----
##
           omega2.delta
## omega2.delta 1
## -----
## ----- Statistical criteria -----
## -----
## Likelihood computed by linearisation
##
      -2LL= 60647.88
##
      AIC = 60655.88
##
     BIC = 60664.19
##
## Likelihood computed by importance sampling
      -2LL= 1381.329
##
      AIC = 1389.329
##
      BIC = 1397.639
## -----
zippoisson.fit<-saemix(saemix.model.zip,saemix.data,saemix.options)</pre>
## Nonlinear mixed-effects model fit by the SAEM algorithm
## -----
            Data
## -----
## Object of class SaemixData
    longitudinal data for use with the SAEM algorithm
## Dataset epilepsy
##
     Structured data: y ~ period + y | subject
##
     X variable for graphs: period (2-week)
##
     covariates: trt (), base (), age (yr)
      reference class for covariate trt : placebo
## Dataset characteristics:
    number of subjects:
```

```
##
      number of observations: 236
##
      average/min/max nb obs: 4.00 / 4 / 4
## First 10 lines of data:
     subject period y y.1
                          trt base age mdv cens occ ytype
## 1
          1
               1 5 5 placebo
                                11 31
                                         0
## 2
                2 3 3 placebo
                                 11 31
          1
                                          0
                                                  1
## 3
                3 3
                      3 placebo
                                 11 31
          1
                                         0
                                 11 31
          1
                4 3
## 4
                      3 placebo
                                          0
                                              0
                                                  1
               1 3
## 5
          2
                      3 placebo
                                 11 30
                                          0
                                              0
                                                  1
                                                       1
         2
               2 5 5 placebo
## 6
                                 11 30
                                         0 0 1
## 7
         2
               3 3 3 placebo
                                 11 30 0 0 1
          2
                4 3 3 placebo
                                         0 0 1
## 8
                                 11 30
                                                       1
                                                      1
## 9
          3
               1 2 2 placebo
                                  6 25
                                         0 0 1
          3 2 4 4 placebo
## 10
                                  6 25
                                        0 0 1 1
          Model
## Nonlinear mixed-effects model
    Model function: count model ZIP
##
    Model type: likelihood
## function(psi,id,xidep) {
    y < -xidep[,2]
##
    lambda<-psi[id,1]</pre>
##
    p0<-psi[id,2]
##
    logp <- log(1-p0) -lambda + y*log(lambda) - log(factorial(y))</pre>
    logp0 \leftarrow log(p0+(1-p0)*exp(-lambda))
##
    logp[y==0] < -logp0[y==0]
##
    return(logp)
## }
## <bytecode: 0x56139d720350>
##
    Nb of parameters: 2
##
        parameter names: lambda p0
##
        distribution:
##
       Parameter Distribution Estimated
## [1,] lambda log-normal Estimated
## [2,] p0
                logit
                            Estimated
   Variance-covariance matrix:
##
       lambda p0
## lambda
         1 0
             0 0
## p0
      No covariate in the model.
##
      Initial values
             lambda p0
## Pop.CondInit 0.5 0.2
## ---- Key algorithm options ----
## -----
##
      Estimation of individual parameters (MAP)
##
      Estimation of standard errors and linearised log-likelihood
##
      Estimation of log-likelihood by importance sampling
##
      Number of iterations: K1=300, K2=100
##
      Number of chains: 1
##
      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 a file: FALSE
        save the graphs to files: FALSE
##
## -----
                 Results
## -----
## ------ Fixed effects ------
## -----
     Parameter Estimate SE CV(%)
## [1,] lambda 5.320 0.748 14
             0.041 0.024 58
## [2,] p0
## ----- Variance of random effects -----
      Parameter Estimate SE CV(%)
##
## lambda omega2.lambda 0.86 0.21 24
## -----
## ----- Correlation matrix of random effects -----
## -----
##
            omega2.lambda
## omega2.lambda 1
## -----
## ----- Statistical criteria -----
## -----
## Likelihood computed by linearisation
      -2LL= 61045.94
      AIC = 61053.94
##
##
      BIC = 61062.25
## Likelihood computed by importance sampling
      -2LL= 1381.314
##
##
      AIC = 1389.314
      BIC = 1397.624
## -----
RAPI
if(testMode)
 data(rapi.saemix) else
  rapi.saemix<-read.table(file.path(datDir, "rapi.saemix.tab"), header=TRUE)
saemix.data<-saemixData(name.data=rapi.saemix, name.group=c("id"),</pre>
                 name.predictors=c("time","rapi"),name.response=c("rapi"),
                 name.covariates=c("gender"),
                 units=list(x="months",y="",covariates=c("")))
## [1] "gender"
##
##
```

The following SaemixData object was successfully created:

```
##
## Object of class SaemixData
##
       longitudinal data for use with the SAEM algorithm
## Dataset rapi.saemix
##
       Structured data: rapi ~ time + rapi | id
##
       X variable for graphs: time (months)
##
       covariates: gender ()
##
         reference class for covariate gender : Men
hist(rapi.saemix$rapi, main="", xlab="RAPI score", breaks=30)
     500
     1000
Frequency
                               20
             0
                                                   40
                                                                      60
```

```
## Models
# Poisson with a time effect
count.poisson<-function(psi,id,xidep) {</pre>
  time<-xidep[,1]
  y < -xidep[,2]
  intercept<-psi[id,1]</pre>
  slope<-psi[id,2]</pre>
  lambda<- exp(intercept + slope*time)</pre>
  logp <- -lambda + y*log(lambda) - log(factorial(y))</pre>
  return(logp)
}
# Fits
## Poisson
### Model without covariate
saemix.model.poi<-saemixModel(model=count.poisson,description="Count model Poisson",modeltype="likeliho")</pre>
                                 psi0=matrix(c(log(5),0.01),ncol=2,byrow=TRUE,dimnames=list(NULL, c("inter
                                 transform.par=c(0,0), omega.init=diag(c(0.5, 0.5)))
```

RAPI score

##
The following SaemixModel object was successfully created:

##

```
##
## Nonlinear mixed-effects model
##
     Model function: Count model Poisson
     Model type: likelihood
##
## function(psi,id,xidep) {
     time<-xidep[,1]
##
     y < -xidep[,2]
##
     intercept<-psi[id,1]</pre>
##
##
     slope<-psi[id,2]</pre>
##
     lambda<- exp(intercept + slope*time)</pre>
     logp <- -lambda + y*log(lambda) - log(factorial(y))</pre>
##
     return(logp)
## }
##
     Nb of parameters: 2
##
         parameter names: intercept slope
##
         distribution:
##
        Parameter Distribution Estimated
## [1,] intercept normal
                                Estimated
## [2,] slope
                  normal
                                Estimated
   Variance-covariance matrix:
##
             intercept slope
## intercept
## slope
                      0
       No covariate in the model.
##
##
       Initial values
                intercept slope
## Pop.CondInit 1.609438 0.01
### Gender effect on intercept and slope
saemix.model.poi.cov2<-saemixModel(model=count.poisson,description="Count model Poisson",modeltype="lik
                                     psi0=matrix(c(log(5),0.01),ncol=2,byrow=TRUE,dimnames=list(NULL, c("
                                     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))
##
##
## The following SaemixModel object was successfully created:
## Nonlinear mixed-effects model
     Model function: Count model Poisson
##
##
     Model type: likelihood
## function(psi,id,xidep) {
##
     time<-xidep[,1]
     y < -xidep[,2]
##
##
     intercept<-psi[id,1]</pre>
##
     slope<-psi[id,2]</pre>
##
     lambda<- exp(intercept + slope*time)</pre>
     logp <- -lambda + y*log(lambda) - log(factorial(y))</pre>
##
     return(logp)
##
## }
     Nb of parameters: 2
##
##
         parameter names: intercept slope
##
         distribution:
##
        Parameter Distribution Estimated
```

```
## [1,] intercept normal Estimated
## [2] slope normal Estimated
## [2,] slope normal
                          Estimated
## Variance-covariance matrix:
##
     intercept slope
## intercept 1
## slope
                 1
   Covariate model:
##
      intercept slope
## [1,]
         1
##
     Initial values
##
            intercept slope
## Pop.CondInit 1.609438 0.01
## Cov.CondInit 0.000000 0.00
saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, displayProgress=FALSE, fim=FALSE)</pre>
### Fit with saemix
poisson.fit<-saemix(saemix.model.poi,saemix.data,saemix.options)</pre>
## Nonlinear mixed-effects model fit by the SAEM algorithm
         Data
## ----
## -----
## Object of class SaemixData
     longitudinal data for use with the SAEM algorithm
## Dataset rapi.saemix
##
     Structured data: rapi ~ time + rapi | id
     X variable for graphs: time (months)
##
     covariates: gender ()
##
##
       reference class for covariate gender : Men
## Dataset characteristics:
##
     number of subjects:
                          818
     number of observations: 3616
##
     average/min/max nb obs: 4.42 / 1 / 5
## First 10 lines of data:
     id time rapi rapi.1 gender mdv cens occ ytype
                                 0
        0 0 0
                                    1
## 1
                       Men 0
     1
            0
## 2
     1
                    0
                        Men 0
                                 0
         6
                                     1
## 3
     1 18 0
                  0 Men
                            0
                               0 1
## 4 2 0 3
                   3 Women 0
        6 6
                   6 Women 0
                                 0 1
## 5 2
                  5 Women 0
## 6 2 12 5
                                0
                                    1
## 7 2 18 4
                               0 1
                  4 Women O
## 8 2 24 5
                  5 Women 0 0 1
## 9 3
         0 9
                                    1
                    9 Men 0
                                 0
            1
                  1 Men O
                                0 1
## 10 3 12
## -----
             Model
## -----
## Nonlinear mixed-effects model
    Model function: Count model Poisson
    Model type: likelihood
##
## function(psi,id,xidep) {
## time<-xidep[,1]</pre>
```

y < -xidep[,2]

```
##
    intercept<-psi[id,1]</pre>
##
    slope<-psi[id,2]</pre>
##
    lambda<- exp(intercept + slope*time)</pre>
    logp <- -lambda + y*log(lambda) - log(factorial(y))</pre>
##
##
    return(logp)
## }
## <bytecode: 0x56139959ffc0>
   Nb of parameters: 2
##
##
       parameter names: intercept slope
##
       distribution:
      Parameter Distribution Estimated
## [1,] intercept normal Estimated
              normal
## [2,] slope
                        Estimated
  Variance-covariance matrix:
##
##
         intercept slope
## intercept 1 0
## slope
                 0
                      1
##
     No covariate in the model.
##
     Initial values
##
            intercept slope
## Pop.CondInit 1.609438 0.01
## -----
## ---- Key algorithm options ----
## -----
##
     Estimation of individual parameters (MAP)
     Estimation of log-likelihood by importance sampling
##
     Number of iterations: K1=300, K2=100
##
     Number of chains: 1
##
     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 a file: FALSE
##
         save the graphs to files: FALSE
## -----
                   Results
## -----
## ----- Fixed effects -----
## -----
      Parameter Estimate
## [1,] intercept 1.577
## [2,] slope -0.033
## ----- Variance of random effects -----
## -----
         Parameter Estimate
## intercept omega2.intercept 0.9039
       omega2.slope 0.0039
## ----- Correlation matrix of random effects -----
##
                omega2.intercept omega2.slope
```

```
## omega2.intercept 1
## omega2.slope 0
## -----
## ------ Statistical criteria -----
## -----
##
## Likelihood computed by importance sampling
      -2LL= 21486.75
##
      AIC = 21496.75
##
      BIC = 21520.29
## -----
poisson.fit.cov2<-saemix(saemix.model.poi.cov2,saemix.data,saemix.options)</pre>
## Nonlinear mixed-effects model fit by the SAEM algorithm
            Data
## -----
## Object of class SaemixData
     longitudinal data for use with the SAEM algorithm
## Dataset rapi.saemix
##
     Structured data: rapi ~ time + rapi | id
     X variable for graphs: time (months)
##
     covariates: gender ()
      reference class for covariate gender : Men
## Dataset characteristics:
##
     number of subjects:
                         818
##
     number of observations: 3616
     average/min/max nb obs: 4.42 / 1 / 5
##
## First 10 lines of data:
     id time rapi rapi.1 gender mdv cens occ ytype
##
## 1
    1 0 0 0 Men 0
    1 6 0
## 2
                  0 Men 0
                                0 1
     1 18 0
## 3
                   0
                     Men O
                               0
## 4 2 0 3
                  3 Women 0 0 1
## 5 2 6 6
                  6 Women 0 0 1
## 6 2 12 5
                  5 Women 0
                              0 1
## 7 2 18 4 4 Women 0 0 1
## 8 2 24 5 5 Women 0 0 1
## 9 3 0 9
                  9 Men 0 0 1
## 10 3 12 1 1 Men 0
                                0 1
            Model
## Nonlinear mixed-effects model
##
   Model function: Count model Poisson
    Model type: likelihood
## function(psi,id,xidep) {
##
    time<-xidep[,1]
##
    y < -xidep[,2]
##
    intercept<-psi[id,1]</pre>
##
    slope<-psi[id,2]</pre>
##
    lambda<- exp(intercept + slope*time)</pre>
##
    logp <- -lambda + y*log(lambda) - log(factorial(y))</pre>
```

##

return(logp)

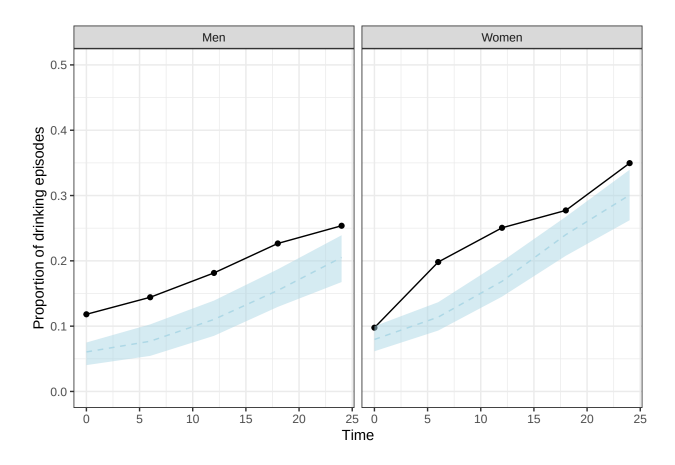
```
## }
## <bytecode: 0x56139959ffc0>
  Nb of parameters: 2
      parameter names: intercept slope
##
      distribution:
##
      Parameter Distribution Estimated
## [1,] intercept normal Estimated
## [2,] slope
                      Estimated
            normal
  Variance-covariance matrix:
##
      intercept slope
## intercept 1
              1
## slope
  Covariate model:
## [,1] [,2]
## gender 1 1
## Initial values
##
           intercept slope
## Pop.CondInit 1.609438 0.01
## Cov.CondInit 0.000000 0.00
## -----
## ---- Key algorithm options ----
## -----
##
     Estimation of individual parameters (MAP)
     Estimation of log-likelihood by importance sampling
     Number of iterations: K1=300, K2=100
##
##
     Number of chains: 1
     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 a file: FALSE
##
        save the graphs to files: FALSE
## -----
                 Results
## -----
## ----- Fixed effects -----
## -----
##
      Parameter
                      Estimate
## [1,] intercept
## [2,] beta_gender(intercept) -0.196
## [3,] slope
                       -0.022
## [4,] beta_gender(slope)
                       -0.017
## -----
## ----- Variance of random effects -----
## -----
        Parameter Estimate
## intercept omega2.intercept 0.9179
## slope omega2.slope 0.0039
## ----- Correlation matrix of random effects -----
##
               omega2.intercept omega2.slope
```

```
## omega2.intercept 1.00
                                       -0.14
## omega2.slope
                                        1.00
                     -0.14
          ----- Statistical criteria ------
##
## Likelihood computed by importance sampling
         -2LL= 21454.94
##
##
         AIC = 21470.94
##
         BIC = 21508.59
exp(poisson.fit@results@fixed.effects)
## [1] 4.8394604 0.9673886
exp(poisson.fit.cov2@results@fixed.effects)
## [1] 5.3842360 0.8217414 0.9780800 0.9833256
### Simulations
saemix.simulatePoisson<-function(psi, id, xidep) {</pre>
  time<-xidep[,1]</pre>
  y < -xidep[,2]
  intercept<-psi[id,1]</pre>
  slope<-psi[id,2]</pre>
  lambda<- exp(intercept + slope*time)</pre>
  y<-rpois(length(time), lambda=lambda)</pre>
  return(y)
}
nsim<-100
yfit1<-simulateDiscreteSaemix(poisson.fit.cov2, saemix.simulatePoisson, nsim=nsim)</pre>
hist(yfit1@data@data$rapi, xlim=c(0,50), freq=F, breaks=30, xlab="Observed counts", main="")
lines(density(yfit1@sim.data@datasim$ysim[yfit1@sim.data@datasim$ysim<50]), lwd = 2, col = 'red')
     0.20
     0.15
Density
     0.10
             0
                           10
                                         20
                                                                      40
                                                        30
                                                                                    50
                                         Observed counts
```

```
cat("Observed proportion of 0's", length(yfit1@data@data$rapi[yfit1@data@data$rapi==0])/yfit1@data@ntot
## Observed proportion of 0's 0.2090708
           Poisson model, p=",length(yfit1@sim.data@datasim$ysim[yfit1@sim.data@datasim$ysim==0])/length
cat("
##
         Poisson model, p = 0.1518501
# Checking proportion of zeroes
vfit<-vfit1
simdat <-yfit@sim.data@datasim</pre>
simdat$time<-rep(yfit@data@data$time,nsim)</pre>
simdat$gender<-rep(yfit@data@data$gender,nsim)</pre>
ytab<-NULL
for(irep in 1:nsim) {
  xtab<-simdat[simdat$irep==irep,]</pre>
  xtab1 <- xtab %>%
    group_by(time, gender) %>%
    summarise(nev = sum(ysim==0), n=n()) %>%
    mutate(freq = nev/n)
  ytab<-rbind(ytab,xtab1[,c("time","gender","freq")])</pre>
}
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
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## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
```

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

```
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
gtab <- ytab %>%
    group_by(time, gender) %>%
  summarise(lower=quantile(freq, c(0.05)), median=quantile(freq, c(0.5)), upper=quantile(freq, c(0.95))
  mutate(gender=ifelse(gender==0, "Men", "Women"))
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
gtab$freq<-1
gtab1<-cbind(gtab, model="Poisson")</pre>
rapipl <- rapi.saemix %>%
    group_by(time, gender) %>%
  summarise(nev = sum(rapi==0), n=n()) %>%
  mutate(freq = nev/n, sd=sqrt((1-nev/n)/nev)) %>%
  mutate(lower=freq-1.96*sd, upper=freq+1.96*sd)
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
rapipl$lower[rapipl$lower<0] <-0 # we should use a better approximation for CI
plot2 <- ggplot(rapip1, aes(x=time, y=freq, group=gender)) + geom_line() +</pre>
  geom_point() +
  geom_line(data=gtab, aes(x=time, y=median, group=gender), linetype=2, colour='lightblue') +
  geom_ribbon(data=gtab,aes(ymin=lower, ymax=upper, group=gender), alpha=0.5, fill='lightblue') +
  ylim(c(0,0.5)) + theme_bw() + theme(legend.position = "none") + facet_wrap(.~gender) +
  xlab("Time") + ylab("Proportion of drinking episodes")
print(plot2)
```



Overdispersion

```
## ZIP Poisson model with time effect
count.poissonzip<-function(psi,id,xidep) {</pre>
  time<-xidep[,1]
  y < -xidep[,2]
  intercept<-psi[id,1]</pre>
  slope<-psi[id,2]</pre>
  p0<-psi[id,3] # Probability of zero's</pre>
  lambda<- exp(intercept + slope*time)</pre>
  logp \leftarrow log(1-p0) - lambda + y*log(lambda) - log(factorial(y)) # Poisson
  logp0 \leftarrow log(p0+(1-p0)*exp(-lambda)) # Zeroes
  logp[y==0] < -logp0[y==0]
  return(logp)
}
## Generalized Poisson model with time effect
count.genpoisson<-function(psi,id,xidep) {</pre>
  time<-xidep[,1]</pre>
  y < -xidep[,2]
  intercept<-psi[id,1]</pre>
  slope<-psi[id,2]</pre>
  lambda<- exp(intercept + slope*time)</pre>
  delta<-psi[id,3]</pre>
  logp <- log(lambda) + (y-1)*log(lambda+y*delta) - lambda - y*delta - log(factorial(y))</pre>
  return(logp)
```

```
## Negative binomial model with time effect
count.NB<-function(psi,id,xidep) {</pre>
  time<-xidep[,1]
  y < -xidep[,2]
  intercept<-psi[id,1]</pre>
  slope<-psi[id,2]</pre>
  k < -psi[id,3]
  lambda<- exp(intercept + slope*time)</pre>
  logp <- log(factorial(y+k-1)) - log(factorial(y)) - log(factorial(k-1)) + y*log(lambda) - y*log(lambd
  return(logp)
}
## ZIP
### base model
saemix.model.zip<-saemixModel(model=count.poissonzip,description="count model ZIP",modeltype="likelihoo")</pre>
                                psi0=matrix(c(1.5, 0.01, 0.2),ncol=3,byrow=TRUE,dimnames=list(NULL, c("in
                                transform.par=c(0,0,3), covariance.model=diag(c(1,1,0)), omega.init=diag(
##
##
## The following SaemixModel object was successfully created:
## Nonlinear mixed-effects model
##
     Model function: count model ZIP
##
     Model type: likelihood
## function(psi,id,xidep) {
##
     time<-xidep[,1]
     y<-xidep[,2]
##
##
     intercept<-psi[id,1]</pre>
     slope<-psi[id,2]</pre>
##
##
     p0<-psi[id,3] # Probability of zero's
##
     lambda<- exp(intercept + slope*time)</pre>
##
     logp \leftarrow log(1-p0) - lambda + y*log(lambda) - log(factorial(y)) # Poisson
     logp0 \leftarrow log(p0+(1-p0)*exp(-lambda)) # Zeroes
##
     logp[y==0] < -logp0[y==0]
##
##
     return(logp)
## }
##
     Nb of parameters: 3
##
         parameter names: intercept slope p0
##
         distribution:
##
        Parameter Distribution Estimated
## [1,] intercept normal
                                 Estimated
## [2,] slope
                   normal
                                Estimated
## [3,] p0
                   logit
                                Estimated
##
     Variance-covariance matrix:
##
             intercept slope p0
## intercept
                            0 0
                      1
## slope
                      0
                            1 0
## p0
                      0
                            0 0
##
       No covariate in the model.
##
       Initial values
##
                 intercept slope p0
```

```
## Pop.CondInit
                      1.5 0.01 0.2
### ZIP Poisson with gender on both intercept
saemix.model.zip.cov1<-saemixModel(model=count.poissonzip,description="count model ZIP",modeltype="like
                                    psi0=matrix(c(1.5, 0.01, 0.2),ncol=3,byrow=TRUE,dimnames=list(NULL,
                                    transform.par=c(0,0,3), covariance.model=diag(c(1,1,0)), omega.init=
                                    covariate.model = matrix(c(1,0,0),ncol=3, byrow=TRUE))
##
##
## The following SaemixModel object was successfully created:
## Nonlinear mixed-effects model
     Model function: count model ZIP
     Model type: likelihood
##
## function(psi,id,xidep) {
##
    time<-xidep[,1]
##
     y < -xidep[,2]
##
     intercept <- psi[id,1]
##
     slope<-psi[id,2]</pre>
##
    pO<-psi[id,3] # Probability of zero's
##
     lambda<- exp(intercept + slope*time)</pre>
##
     logp <- log(1-p0) -lambda + y*log(lambda) - log(factorial(y)) # Poisson</pre>
##
     logp0 \leftarrow log(p0+(1-p0)*exp(-lambda)) # Zeroes
##
     logp[y==0] < -logp0[y==0]
##
     return(logp)
## }
##
    Nb of parameters: 3
##
         parameter names: intercept slope p0
##
         distribution:
##
        Parameter Distribution Estimated
                               Estimated
## [1,] intercept normal
## [2,] slope
                                Estimated
                  normal
## [3,] p0
                  logit
                                Estimated
##
    Variance-covariance matrix:
##
             intercept slope p0
## intercept
                     1
## slope
                     0
                            1 0
                     0
## p0
##
     Covariate model:
##
        intercept slope p0
## [1,]
##
       Initial values
##
                intercept slope p0
## Pop.CondInit
                      1.5 0.01 0.2
## Cov.CondInit
                      0.0 0.00 0.0
### ZIP Poisson with gender on both intercept and slope
saemix.model.zip.cov2<-saemixModel(model=count.poissonzip,description="count model ZIP",modeltype="like
                                    psi0=matrix(c(1.5, 0.01, 0.2),ncol=3,byrow=TRUE,dimnames=list(NULL,
                                    transform.par=c(0,0,3), covariance.model=diag(c(1,1,0)), omega.init=
                                    covariate.model = matrix(c(1,1,0),ncol=3, byrow=TRUE))
##
##
```

The following SaemixModel object was successfully created:

```
##
## Nonlinear mixed-effects model
##
    Model function: count model ZIP
    Model type: likelihood
##
## function(psi,id,xidep) {
    time<-xidep[,1]
##
    y < -xidep[,2]
##
    intercept<-psi[id,1]</pre>
##
    slope<-psi[id,2]</pre>
##
##
    p0<-psi[id,3] # Probability of zero's
    lambda<- exp(intercept + slope*time)</pre>
    logp <- log(1-p0) -lambda + y*log(lambda) - log(factorial(y)) # Poisson</pre>
##
    logp0 \leftarrow log(p0+(1-p0)*exp(-lambda)) # Zeroes
##
##
    logp[y==0] < -logp0[y==0]
##
    return(logp)
## }
##
    Nb of parameters: 3
##
        parameter names: intercept slope p0
##
        distribution:
##
       Parameter Distribution Estimated
## [1,] intercept normal Estimated
## [2,] slope
              normal
                             Estimated
## [3,] p0
                logit
                            Estimated
   Variance-covariance matrix:
##
##
       intercept slope p0
## intercept
                1
                        0 0
                         1 0
                    0
## slope
                          0 0
## p0
                    0
##
    Covariate model:
##
       intercept slope p0
## [1,]
              1
##
      Initial values
##
              intercept slope p0
## Pop.CondInit
                    1.5 0.01 0.2
                     0.0 0.00 0.0
## Cov.CondInit
zippoisson.fit<-saemix(saemix.model.zip,saemix.data,saemix.options)
## Nonlinear mixed-effects model fit by the SAEM algorithm
## -----
                Data
## -----
## Object of class SaemixData
      longitudinal data for use with the SAEM algorithm
## Dataset rapi.saemix
##
      Structured data: rapi ~ time + rapi | id
##
      X variable for graphs: time (months)
##
      covariates: gender ()
##
        reference class for covariate gender : Men
## Dataset characteristics:
##
      number of subjects:
##
      number of observations: 3616
      average/min/max nb obs: 4.42 / 1 / 5
## First 10 lines of data:
     id time rapi rapi.1 gender mdv cens occ ytype
```

```
## 1
          0
               0
                     0
                       Men
                        Men
## 2
               0
                     0
                               0
                                   0
      1
          6
                                       1
## 3
         18
                        Men
## 4
          0
                                   0 1
      2
              3
                     3 Women 0
                                            1
## 5
      2
          6
              6
                     6 Women
                                   0
## 6
     2 12 5
                             0
                                  0
                                      1
                     5 Women
## 7
      2 18 4
                     4 Women 0
                                           1
      2
         24 5
                     5 Women 0
                                   0 1
## 8
## 9
      3
        0 9
                     9
                        Men 0
                                   0
                                      1
                                            1
                       Men O
                                   0 1
## 10 3 12 1
                     1
## ----
         Model
## -----
## Nonlinear mixed-effects model
    Model function: count model ZIP
##
    Model type: likelihood
## function(psi,id,xidep) {
    time<-xidep[,1]
##
##
    y < -xidep[,2]
##
    intercept <- psi[id,1]
##
    slope<-psi[id,2]</pre>
##
    pO<-psi[id,3] # Probability of zero's
##
    lambda<- exp(intercept + slope*time)</pre>
##
    logp <- log(1-p0) -lambda + y*log(lambda) - log(factorial(y)) # Poisson</pre>
##
    logp0 \leftarrow log(p0+(1-p0)*exp(-lambda)) # Zeroes
##
    logp[y==0] < -logp0[y==0]
##
    return(logp)
## }
## <bytecode: 0x56139a088e30>
    Nb of parameters: 3
##
        parameter names: intercept slope p0
##
        distribution:
##
       Parameter Distribution Estimated
## [1,] intercept normal Estimated
## [2,] slope normal
                           Estimated
## [3,] p0
                           Estimated
               logit
##
   Variance-covariance matrix:
##
          intercept slope p0
## intercept 1
                       0 0
## slope
                  0
                        1 0
## p0
                 0
                        0 0
##
      No covariate in the model.
##
      Initial values
##
             intercept slope p0
## Pop.CondInit 1.5 0.01 0.2
## -----
         Key algorithm options ----
## -----
##
      Estimation of individual parameters (MAP)
##
      Estimation of log-likelihood by importance sampling
##
      Number of iterations: K1=300, K2=100
##
      Number of chains: 1
##
      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 a file: FALSE
        save the graphs to files: FALSE
##
## -----
                 Results
## -----
## ----- Fixed effects -----
## -----
     Parameter Estimate
## [1,] intercept 1.657
## [2,] slope -0.029
## [3,] p0
             0.076
## ----- Variance of random effects -----
## -----
        Parameter
                     Estimate
## intercept omega2.intercept 0.7977
## slope omega2.slope 0.0032
## -----
## ----- Correlation matrix of random effects -----
## -----
##
              omega2.intercept omega2.slope
## omega2.intercept 1
                   0
                           1
## omega2.slope
## ----- Statistical criteria -----
## -----
## Likelihood computed by importance sampling
     -2LL= 20479.88
##
##
      AIC = 20491.88
      BIC = 20520.12
zippoisson.fit.cov1<-saemix(saemix.model.zip.cov1,saemix.data,saemix.options)
## Nonlinear mixed-effects model fit by the SAEM algorithm
## -----
           Data
## -----
## Object of class SaemixData
##
     longitudinal data for use with the SAEM algorithm
## Dataset rapi.saemix
     Structured data: rapi ~ time + rapi | id
##
     X variable for graphs: time (months)
##
     covariates: gender ()
##
      reference class for covariate gender : Men
## Dataset characteristics:
##
     number of subjects:
     number of observations: 3616
     average/min/max nb obs: 4.42 / 1 / 5
## First 10 lines of data:
```

```
id time rapi rapi.1 gender mdv cens occ ytype
## 1
      1
          0
               0
                      0
                          Men
                                0
                                     0
                                         1
                          Men
## 2
      1
               0
## 3
          18
               0
                          Men
                                     0
                      0
                              0
                                        1
      1
## 4
      2
          0
               3
                      3 Women
                               0
                                     0
## 5
      2
          6
               6
                      6 Women
                               0
                                    0
## 6
      2 12
               5
                      5 Women 0
      2 18
## 7
               4
                      4 Women O
                                     0
          24
## 8
      2
               5
                      5 Women 0
                                     0
        0
## 9
      3
               9
                              0
                                     0
                      9
                        Men
## 10 3
        12
             1
                      1
                        Men
               Model
## -----
## Nonlinear mixed-effects model
##
    Model function: count model ZIP
##
    Model type: likelihood
## function(psi,id,xidep) {
##
    time<-xidep[,1]
##
    y < -xidep[,2]
##
    intercept<-psi[id,1]</pre>
##
    slope<-psi[id,2]</pre>
##
    p0<-psi[id,3] # Probability of zero's
##
    lambda<- exp(intercept + slope*time)</pre>
##
    logp <- log(1-p0) -lambda + y*log(lambda) - log(factorial(y)) # Poisson</pre>
    logp0 \leftarrow log(p0+(1-p0)*exp(-lambda)) # Zeroes
##
    logp[y==0] < -logp0[y==0]
##
    return(logp)
## }
## <bytecode: 0x56139a088e30>
##
    Nb of parameters: 3
##
        parameter names: intercept slope p0
##
        distribution:
##
       Parameter Distribution Estimated
## [1,] intercept normal Estimated
## [2,] slope normal
                           Estimated
## [3,] p0
               logit
                            Estimated
## Variance-covariance matrix:
##
    intercept slope p0
               1 0 0
## intercept
## slope
                   0
                        1 0
## p0
                   0
                        0 0
   Covariate model:
       [,1] [,2] [,3]
## gender 1 0
##
      Initial values
              intercept slope p0
##
## Pop.CondInit 1.5 0.01 0.2
## Cov.CondInit
                  0.0 0.00 0.0
## -----
## ---- Key algorithm options ----
##
      Estimation of individual parameters (MAP)
##
      Estimation of log-likelihood by importance sampling
```

```
##
     Number of iterations: K1=300, K2=100
##
     Number of chains: 1
##
     Seed: 632545
     Number of MCMC iterations for IS: 5000
##
##
##
        nb of simulated datasets used for npde: 1000
##
        nb of simulated datasets used for VPC: 100
##
     Input/output
##
        save the results to a file: FALSE
##
        save the graphs to files: FALSE
                  Results
## -----
## ----- Fixed effects -----
##
     Parameter
                       Estimate
## [1,] intercept
                       1.786
## [2,] beta_gender(intercept) -0.226
## [3,] slope
                     -0.029
## [4,] p0
                      0.076
## -----
## ----- Variance of random effects -----
## -----
                      Estimate
         Parameter
## intercept omega2.intercept 0.7849
## slope omega2.slope 0.0033
## ----- Correlation matrix of random effects -----
    omega2.intercept omega2.slope
## omega2.intercept 1
## omega2.slope 0
                           1
## -----
## ----- Statistical criteria -----
## -----
## Likelihood computed by importance sampling
##
      -2LL= 20469.41
##
      AIC = 20483.41
##
     BIC = 20516.35
## -----
zippoisson.fit.cov2<-saemix(saemix.model.zip.cov2,saemix.data,saemix.options)</pre>
## Nonlinear mixed-effects model fit by the SAEM algorithm
## -----
           Data
## -----
## Object of class SaemixData
##
     longitudinal data for use with the SAEM algorithm
## Dataset rapi.saemix
##
     Structured data: rapi ~ time + rapi | id
##
     X variable for graphs: time (months)
##
     covariates: gender ()
      reference class for covariate gender : Men
```

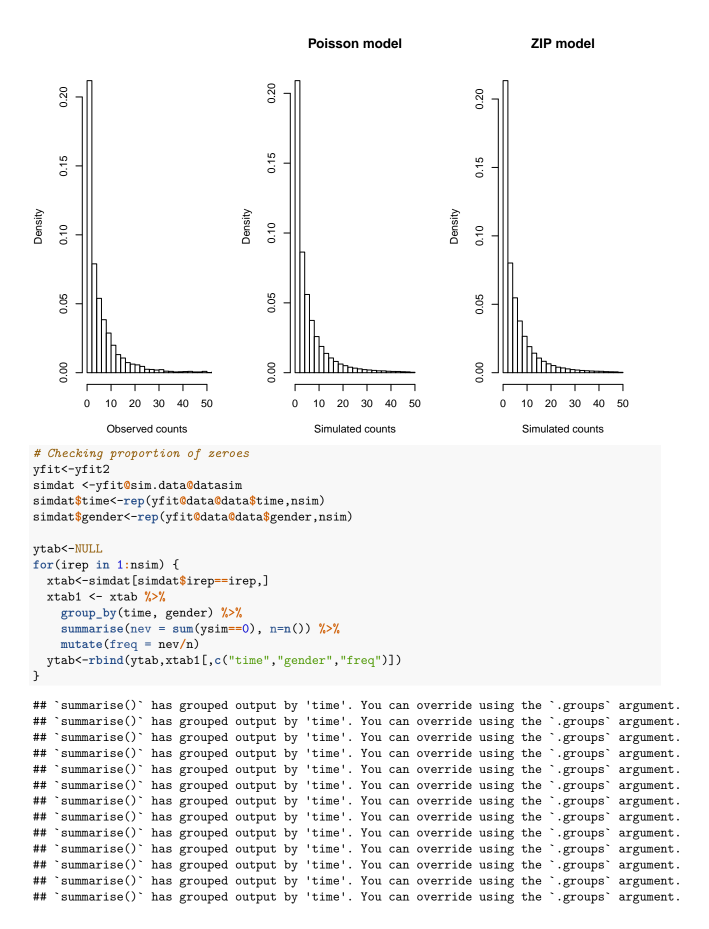
```
## Dataset characteristics:
##
      number of subjects:
                            818
      number of observations: 3616
##
      average/min/max nb obs: 4.42 / 1 / 5
##
## First 10 lines of data:
##
     id time rapi rapi.1 gender mdv cens occ ytype
          0
               0
                      0
                          Men
## 2
          6
               0
                      0
                           Men
      1
                                0
                                     0
                                         1
## 3
      1
          18
               0
                      0
                           Men
                                0
                                     0
## 4
      2
        0 3
                                0
                                     0
                      3 Women
## 5
      2 6 6
                      6 Women
      2 12 5
## 6
                      5 Women
                               0
                                     0
## 7
      2 18 4
                      4 Women
                               0
                                     0
                                        1
                                             1
## 8
      2 24 5
                      5 Women
                                   0
## 9
      3 0 9
                        Men
                                0
                                     0
                      9
                                       1
                                             1
## 10 3
          12
             1
                      1
                           Men
                                0
                                     0
              Model
## -----
## Nonlinear mixed-effects model
    Model function: count model ZIP
##
    Model type: likelihood
## function(psi,id,xidep) {
##
    time<-xidep[,1]
##
    y<-xidep[,2]
##
    intercept<-psi[id,1]</pre>
##
    slope<-psi[id,2]</pre>
##
    p0<-psi[id,3] # Probability of zero's
##
    lambda<- exp(intercept + slope*time)</pre>
    logp <- log(1-p0) -lambda + y*log(lambda) - log(factorial(y)) # Poisson</pre>
##
    logp0 \leftarrow log(p0+(1-p0)*exp(-lambda)) # Zeroes
    logp[y==0] < -logp0[y==0]
##
##
    return(logp)
## }
## <bytecode: 0x56139a088e30>
##
    Nb of parameters: 3
        parameter names: intercept slope p0
##
##
        distribution:
##
       Parameter Distribution Estimated
## [1,] intercept normal Estimated
## [2,] slope normal
                           Estimated
## [3,] p0
                logit
                            Estimated
   Variance-covariance matrix:
           intercept slope p0
## intercept
                 1
                         1 0
                   0
## slope
## p0
                   0
    Covariate model:
        [,1] [,2] [,3]
## gender 1 1
##
      Initial values
##
              intercept slope p0
## Pop.CondInit 1.5 0.01 0.2
## Cov.CondInit
                  0.0 0.00 0.0
```

```
## ----
      Key algorithm options ----
## -----
##
     Estimation of individual parameters (MAP)
     Estimation of log-likelihood by importance sampling
##
     Number of iterations: K1=300, K2=100
##
    Number of chains: 1
    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 a file: FALSE
##
       save the graphs to files: FALSE
## -----
                 Results
  -----
## ----- Fixed effects -----
## -----
##
     Parameter
                     Estimate
## [1,] intercept
## [2,] beta_gender(intercept) -0.197
## [3,] slope
                      -0.020
## [4,] beta_gender(slope)
                     -0.016
## [5,] p0
                      0.075
## ----- Variance of random effects -----
   Parameter Estimate
## intercept omega2.intercept 0.7826
                   0.0033
## slope omega2.slope
## -----
## ----- Correlation matrix of random effects -----
## -----
##
             omega2.intercept omega2.slope
## omega2.intercept 1 0
            0
## omega2.slope
## -----
## ----- Statistical criteria -----
## -----
## Likelihood computed by importance sampling
    -2LL= 20459.27
     AIC = 20475.27
##
     BIC = 20512.93
exp(zippoisson.fit@results@fixed.effects)
## [1] 5.2450012 0.9714983 1.0793068
exp(zippoisson.fit.cov1@results@fixed.effects)
```

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[1] 5.9656256 0.7975888 0.9714754 1.0793259

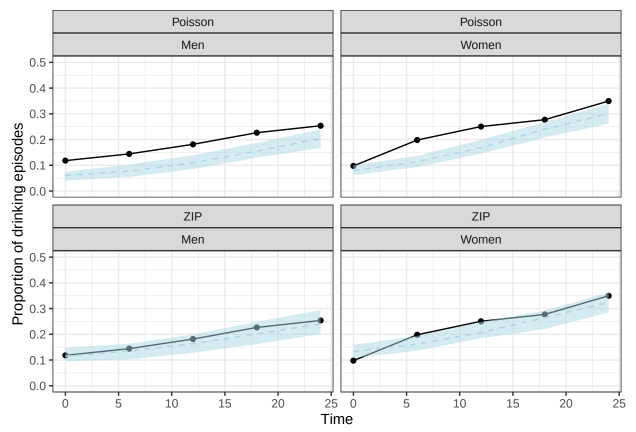
```
exp(zippoisson.fit.cov2@results@fixed.effects)
## [1] 5.8872267 0.8213610 0.9797720 0.9844017 1.0783237
saemix.simulatePoissonZIP<-function(psi, id, xidep) {</pre>
 time<-xidep[,1]
  y < -xidep[,2]
  intercept<-psi[id,1]</pre>
  slope<-psi[id,2]</pre>
  p0<-psi[id,3] # Probability of zero's
  lambda<- exp(intercept + slope*time)</pre>
  prob0<-rbinom(length(time), size=1, prob=p0)</pre>
  y<-rpois(length(time), lambda=lambda)</pre>
 y[prob0==1]<-0
  return(y)
yfit2<-simulateDiscreteSaemix(zippoisson.fit.cov2, saemix.simulatePoissonZIP, 100)
cat("Observed proportion of 0's", length(yfit1@data@data$rapi[yfit1@data@data$rapi==0])/yfit1@data@ntot
## Observed proportion of 0's 0.2090708
cat("
           Poisson model, p=",length(yfit1@sim.data@datasim$ysim[yfit1@sim.data@datasim$ysim==0])/lengti
##
         Poisson model, p= 0.1518501
cat(" ZI-Poisson model, p=",length(yfit2@sim.data@datasim$ysim[yfit2@sim.data@datasim$ysim==0])/length
     ZI-Poisson model, p= 0.1957329
##
par(mfrow=c(1,3))
hist(yfit1@data@data$rapi, xlim=c(0,50), freq=F, breaks=30, xlab="Observed counts", main="")
hist(yfit10sim.dataodatasim$ysim[yfit10sim.dataodatasim$ysim<50], xlim=c(0,50), freq=F, breaks=20, xlab
hist(yfit20sim.data0datasim$ysim[yfit20sim.data0datasim$ysim<50], xlim=c(0,50), freq=F, breaks=20, xlab
```



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

```
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `
                                                                            .groups argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `
                                                                            .groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `
                                                                            .groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `
                                                                            .groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
gtab <- ytab %>%
    group_by(time, gender) %>%
  summarise(lower=quantile(freq, c(0.05)), median=quantile(freq, c(0.5)), upper=quantile(freq, c(0.95))
  mutate(gender=ifelse(gender==0, "Men", "Women"))
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
gtab$freq<-1
gtab2<-cbind(gtab, model="ZIP")</pre>
gtab<-rbind(gtab1, gtab2)
rapipl <- rapi.saemix %>%
    group by(time, gender) %>%
  summarise(nev = sum(rapi==0), n=n()) %>%
  mutate(freq = nev/n, sd=sqrt((1-nev/n)/nev)) %>%
  mutate(lower=freq-1.96*sd, upper=freq+1.96*sd)
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
rapipl$lower[rapipl$lower<0] <-0 # we should use a better approximation for CI
plot2 <- ggplot(rapipl, aes(x=time, y=freq, group=gender)) + geom_line() +</pre>
```

```
geom_point() +
geom_line(data=gtab, aes(x=time, y=median, group=gender), linetype=2, colour='lightblue') +
geom_ribbon(data=gtab,aes(ymin=lower, ymax=upper, group=gender), alpha=0.5, fill='lightblue') +
ylim(c(0,0.5)) + theme_bw() + theme(legend.position = "none") + facet_wrap(model~gender) +
xlab("Time") + ylab("Proportion of drinking episodes")
print(plot2)
```



Hurdle model

##

covariates: gender ()

```
## Hurdle - 2 models
saemix.data1<-saemixData(name.data=rapi.saemix[rapi.saemix$rapi>0,], name.group=c("id"),
                         name.predictors=c("time", "rapi"), name.response=c("rapi"),
                         name.covariates=c("gender"),
                         units=list(x="week",y="",covariates=c("")))
## [1] "gender"
##
##
##
  The following SaemixData object was successfully created:
##
  Object of class SaemixData
##
       longitudinal data for use with the SAEM algorithm
##
## Dataset rapi.saemix[rapi.saemix$rapi > 0, ]
       Structured data: rapi ~ time + rapi | id
##
##
       X variable for graphs: time (week)
```

```
##
                     reference class for covariate gender : Men
rapi.saemix$y0<-as.integer(rapi.saemix$rapi==0)
saemix.data0<-saemixData(name.data=rapi.saemix, name.group=c("id"),</pre>
                                                          name.predictors=c("time", "y0"), name.response=c("y0"),
                                                          name.covariates=c("gender"),
                                                          units=list(x="week",y="",covariates=c("")))
## [1] "gender"
##
##
## The following SaemixData object was successfully created:
##
## Object of class SaemixData
##
                longitudinal data for use with the SAEM algorithm
## Dataset rapi.saemix
##
                Structured data: y0 ~ time + y0 | id
##
                X variable for graphs: time (week)
                covariates: gender ()
##
##
                     reference class for covariate gender : Men
# Fit Binomial model to saemix.dataO
binary.model<-function(psi,id,xidep) {</pre>
    tim<-xidep[,1]
    y < -xidep[,2]
    inter<-psi[id,1]</pre>
    slope<-psi[id,2]
    logit<-inter+slope*tim</pre>
    pevent<-exp(logit)/(1+exp(logit))</pre>
    pobs = (y==0)*(1-pevent)+(y==1)*pevent
    logpdf <- log(pobs)</pre>
    return(logpdf)
}
saemix.hurdleO<-saemixModel(model=binary.model,description="Binary model",</pre>
                                                                  modeltype="likelihood",
                                                                  psi0=matrix(c(-1.5,-.1,0,0),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=TRUE,dimnames=list(NULL,c("theta1"),ncol=2,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,byrow=1,b
                                                                  transform.par=c(0,0), covariate.model=c(1,1),
                                                                  covariance.model=matrix(c(1,0,0,1),ncol=2), omega.init=diag(c(1,0.3)))
##
##
## The following SaemixModel object was successfully created:
##
## Nonlinear mixed-effects model
           Model function: Binary model
##
           Model type: likelihood
## function(psi,id,xidep) {
##
           tim<-xidep[,1]
           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>
           pobs = (y==0)*(1-pevent)+(y==1)*pevent
##
```

```
##
    logpdf <- log(pobs)</pre>
##
    return(logpdf)
## }
##
    Nb of parameters: 2
##
        parameter names: theta1 theta2
##
        distribution:
       Parameter Distribution Estimated
## [1,] theta1 normal
                            Estimated
## [2,] theta2
                normal
                            Estimated
    Variance-covariance matrix:
        theta1 theta2
## theta1
            1
## theta2
             0
##
   Covariate model:
##
       theta1 theta2
## [1,]
         1
##
      Initial values
##
              theta1 theta2
## Pop.CondInit -1.5
                      -0.1
## Cov.CondInit
                 0.0
                        0.0
saemix.options<-list(seed=1234567,save=FALSE,save.graphs=FALSE, displayProgress=FALSE, nb.chains=10, fix</pre>
hurdlefit0<-saemix(saemix.hurdle0,saemix.data0,saemix.options)
## Nonlinear mixed-effects model fit by the SAEM algorithm
## -----
              Data
## -----
## Object of class SaemixData
##
      longitudinal data for use with the SAEM algorithm
## Dataset rapi.saemix
      Structured data: y0 ~ time + y0 | id
##
##
      X variable for graphs: time (week)
##
      covariates: gender ()
        reference class for covariate gender : Men
## Dataset characteristics:
##
      number of subjects:
                            818
##
      number of observations: 3616
      average/min/max nb obs: 4.42 / 1 / 5
## First 10 lines of data:
##
     id time y0 y0.1 gender mdv cens occ ytype
## 1
          0 1
                      Men
                                 0
     1
                1
                           0
                                    1
## 2
      1
          6 1
                  1
                       Men
                            0
                                 0
                                    1
## 3
      1
          18 1
                  1
                       Men
                            0
                                 0
                                    1
## 4
      2
          0 0
                  0 Women
                           Ο
                                 0
                                    1
## 5
      2
          6 0
                  0 Women
## 6
          12 0
                  0 Women
                                 0
      2
                           Ω
                                    1
## 7
      2
          18 0
                  0 Women
                            0
                                 0
                  0 Women
## 8
      2
          24 0
                           0
                                 0 1
                                          1
## 9
      3
         0 0
                  0
                     Men
                                    1
## 10 3
          12 0
                           0
                                 0
                  0
                      Men
                                     1
                                          1
```

Model

```
## Nonlinear mixed-effects model
##
    Model function: Binary model
    Model type: likelihood
## function(psi,id,xidep) {
##
    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>
##
    pobs = (y==0)*(1-pevent)+(y==1)*pevent
##
    logpdf <- log(pobs)</pre>
    return(logpdf)
##
## }
## <bytecode: 0x56139857d680>
##
    Nb of parameters: 2
##
       parameter names: theta1 theta2
##
       distribution:
##
      Parameter Distribution Estimated
## [1,] theta1 normal Estimated
                         Estimated
## [2,] theta2 normal
   Variance-covariance matrix:
##
        theta1 theta2
## theta1
           1 0
## theta2
           0
## Covariate model:
    [,1] [,2]
##
## gender 1
     Initial values
##
##
            theta1 theta2
## Pop.CondInit -1.5 -0.1
                    0.0
## Cov.CondInit 0.0
## -----
## ---- Key algorithm options ----
## -----
##
     Estimation of individual parameters (MAP)
##
     Estimation of log-likelihood by importance sampling
##
     Number of iterations: K1=300, K2=100
     Number of chains: 10
##
##
     Seed: 1234567
##
     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 a file: FALSE
         save the graphs to files: FALSE
## -----
                    Results
## -----
## ----- Fixed effects -----
## -----
##
      Parameter
                      Estimate
## [1,] theta1
                       -2.796
```

```
## [2,] beta_gender(theta1) 0.132
## [3,] theta2
                         0.036
## [4,] beta_gender(theta2) 0.030
## -----
## ----- Variance of random effects -----
## -----
        Parameter
                   Estimate
## theta1 omega2.theta1 2.4033
## theta2 omega2.theta2 0.0062
## ----- Correlation matrix of random effects -----
## -----
##
              omega2.theta1 omega2.theta2
## omega2.theta1 1
                           Ω
## omega2.theta2 0
## -----
## ----- Statistical criteria -----
## Likelihood computed by importance sampling
##
       -2LL= 3249.132
       AIC = 3263.132
##
##
       BIC = 3296.08
cat("Expected proportion of 0's at time 0:",1/(1+exp(-hurdlefit0@results@fixed.effects[1])),"\n")
## Expected proportion of 0's at time 0: 0.05753853
table(rapi.saemix$rapi[rapi.saemix$time==0] == 0) # 10.6%
##
## FALSE TRUE
   731
          87
# Fit Poisson model to saemix.data1
saemix.hurdle1.cov2<-saemixModel(model=count.poisson,description="Count model Poisson",modeltype="likel
                             psi0=matrix(c(log(5),0.01),ncol=2,byrow=TRUE,dimnames=list(NULL, c("in
                             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))
##
## The following SaemixModel object was successfully created:
## Nonlinear mixed-effects model
    Model function: Count model Poisson
##
    Model type: likelihood
## function(psi,id,xidep) {
    time<-xidep[,1]
##
    y<-xidep[,2]
##
##
    intercept<-psi[id,1]</pre>
    slope<-psi[id,2]</pre>
##
    lambda<- exp(intercept + slope*time)</pre>
##
    logp <- -lambda + y*log(lambda) - log(factorial(y))</pre>
##
```

```
return(logp)
## }
## <bytecode: 0x56139959ffc0>
    Nb of parameters: 2
##
        parameter names: intercept slope
##
        distribution:
       Parameter Distribution Estimated
## [1,] intercept normal
                             Estimated
## [2,] slope
                 normal
                             Estimated
   Variance-covariance matrix:
            intercept slope
## intercept
                   1
## slope
                    1
##
    Covariate model:
##
       intercept slope
## [1,]
             1
##
      Initial values
##
              intercept slope
## Pop.CondInit 1.609438 0.01
## Cov.CondInit 0.000000 0.00
saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, displayProgress=FALSE)</pre>
hurdlefit1<-saemix(saemix.hurdle1.cov2,saemix.data1,saemix.options)
## Error in solve.default(F0) :
    routine Lapack dgesv : le système est exactement singulier : U[2,2] = 0
## Nonlinear mixed-effects model fit by the SAEM algorithm
## -----
                Data
## -----
## Object of class SaemixData
      longitudinal data for use with the SAEM algorithm
## Dataset rapi.saemix[rapi.saemix$rapi > 0, ]
      Structured data: rapi ~ time + rapi | id
##
##
      X variable for graphs: time (week)
##
      covariates: gender ()
        reference class for covariate gender : Men
## Dataset characteristics:
      number of subjects:
      number of observations: 2860
##
      average/min/max nb obs: 3.57 / 1 / 5
## First 10 lines of data:
##
     id time rapi rapi.1 gender mdv cens occ ytype
                3
                                      0
## 4
      2
           0
                      3 Women
                                 0
                                          1
## 5
      2
                6
                       6 Women
                                 0
                                      0
                                          1
          6
## 6
          12
                5
                      5 Women
                                      0
## 7
      2
          18
                4
                       4 Women
                                      0
                                 0
                                          1
## 8
      2
          24
                5
                       5 Women
                                 0
                                      0
## 9
      3
          0
                9
                      9
                         Men
                                 0
                                      0
                                          1
## 10 3
         12
               1
                         Men
                      1
## 12 4
          0
                3
                      3 Women
                                 0
                                      0
                                          1
## 13 4
           6
               2
                      2 Women
                                 0
                                      0
                                          1
## 14 5
                                 0
                                      0
           0 35
                      35 Women
```

```
## ---- Model
## -----
## Nonlinear mixed-effects model
    Model function: Count model Poisson
    Model type: likelihood
## function(psi,id,xidep) {
    time<-xidep[,1]
    y < -xidep[,2]
##
##
    intercept<-psi[id,1]</pre>
##
    slope<-psi[id,2]</pre>
##
    lambda<- exp(intercept + slope*time)</pre>
    logp <- -lambda + y*log(lambda) - log(factorial(y))</pre>
##
##
    return(logp)
## }
## <bytecode: 0x56139959ffc0>
##
    Nb of parameters: 2
##
       parameter names: intercept slope
##
       distribution:
##
      Parameter Distribution Estimated
## [1,] intercept normal Estimated
## [2,] slope normal
                       Estimated
## Variance-covariance matrix:
##
    intercept slope
## Covariate model:
##
    [,1] [,2]
## gender 1
     Initial values
##
            intercept slope
## Pop.CondInit 1.609438 0.01
## Cov.CondInit 0.000000 0.00
## -----
## ---- Key algorithm options ----
## -----
##
     Estimation of individual parameters (MAP)
##
     Estimation of standard errors and linearised log-likelihood
##
     Estimation of log-likelihood by importance sampling
##
     Number of iterations: K1=300, K2=100
##
     Number of chains: 1
##
     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 a file: FALSE
         save the graphs to files: FALSE
## -----
                    Results
## -----
## ----- Fixed effects -----
## -----
                         Estimate SE CV(%) p-value
##
      Parameter
```

```
## [1,] intercept
                  1.8656 0.066 3.5 -
## [2,] beta_gender(intercept) -0.1972  0.089  44.9  0.013
         -0.0059 0.057 955.8 -
## [3,] slope
## [4,] beta_gender(slope) -0.0085 0.075 881.7 0.455
## -----
## ----- Variance of random effects -----
## -----
       Parameter
                   Estimate SE CV(%)
## covar
      cov.intercept.slope -0.0103 NA NA
## -----
## ----- Correlation matrix of random effects -----
## -----
           omega2.intercept omega2.slope
## omega2.intercept 1.00
                -0.32
## omega2.slope -0.32
                     1.00
## -----
## ----- Statistical criteria -----
## -----
## Likelihood computed by linearisation
     -2LL= 437509.5
    AIC = 437525.5
##
     BIC = 437563
##
##
## Likelihood computed by importance sampling
     -2LL= 17628.18
     AIC = 17644.18
     BIC = 17681.67
summary(hurdlefit0)
## -----
## ----- Fixed effects ------
## -----
        Parameter Estimate
          theta1 -2.796
## 2 beta_gender(theta1)
               0.132
          theta2 0.036
## 4 beta_gender(theta2) 0.030
## -----
## ----- Variance of random effects -----
## -----
##
        Parameter Estimate
## theta1 omega2.theta1 2.4033
## theta2 omega2.theta2 0.0062
## -----
## ----- Correlation matrix of random effects -----
## -----
          omega2.theta1 omega2.theta2
## omega2.theta1 1.00 0.00
              1.00
## omega2.theta2 0.00
## -----
## ----- Statistical criteria -----
```

```
##
## Likelihood computed by importance sampling
      -2LL= 3249.132
       AIC = 3263.132
##
       BIC = 3296.08
summary(hurdlefit1)
## -----
## ----- Fixed effects -----
## -----
## Warning in .local(object, ...): NAs introduits lors de la conversion automatique
              Parameter Estimate
                                 SE CV(%) p-value
##
## 1
              intercept 1.8656 0.066 3.53
## 2 beta_gender(intercept) -0.1972 0.089 44.92 0.013
                  slope -0.0059 0.057 955.79
## 4
       beta_gender(slope) -0.0085 0.075 881.67 0.455
## ----- Variance of random effects -----
## -----
##
                Parameter Estimate SE CV(%)
## intercept omega2.intercept 0.6000 NA
## slope omega2.slope 0.0017 NA
## ----- Correlation matrix of random effects -----
##
                omega2.intercept omega2.slope
## omega2.intercept 1.00 -0.32
## omega2.slope -0.32
                              1.00
## ----- Statistical criteria -----
## -----
## Likelihood computed by linearisation
##
       -2LL= 437509.5
       AIC = 437525.5
##
##
      BIC = 437563
## Likelihood computed by importance sampling
       -2LL= 17628.18
##
       AIC = 17644.18
       BIC = 17681.67
# Simulate binary data
simulBinary<-function(psi,id,xidep) {</pre>
 tim<-xidep[,1]
 y < -xidep[,2]
 inter<-psi[id,1]
 slope<-psi[id,2]</pre>
 logit<-inter+slope*tim</pre>
 pevent<-exp(logit)/(1+exp(logit))</pre>
 ysim<-rbinom(length(tim),size=1, prob=pevent)</pre>
```

```
return(ysim)
# proportion of 0's in the data
rapi.tab <- table(rapi.saemix$rapi == 0)</pre>
nsim<-100
ysim.hurdle0 <- simulateDiscreteSaemix(hurdlefit0, simulBinary, nsim=nsim)</pre>
cat("Observed proportion of 0's overall:",rapi.tab[2]/sum(rapi.tab),"\n")
## Observed proportion of 0's overall: 0.2090708
cat("Simulated proportion of 0's overall:",sum(ysim.hurdle0@sim.data@datasim$ysim)/length(ysim.hurdle0@
## Simulated proportion of 0's overall: 0.2069994
# Graph of proportion of O's with time
yfit<-ysim.hurdle0
simdat <-yfit@sim.data@datasim</pre>
simdat$time<-rep(yfit@data@data$time,nsim)</pre>
simdat$gender<-rep(yfit@data@data$gender,nsim)</pre>
vtab<-NULL
for(irep in 1:nsim) {
  xtab<-simdat[simdat$irep==irep,]</pre>
  xtab1 <- xtab %>%
    group_by(time, gender) %>%
    summarise(nev = sum(ysim), n=n()) %>%
    mutate(freq = nev/n)
  ytab<-rbind(ytab,xtab1[,c("time","gender","freq")])</pre>
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
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## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
```

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

```
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
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## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
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## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
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## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
gtab <- ytab %>%
      group by(time, gender) %>%
   summarise(lower=quantile(freq, c(0.05)), median=quantile(freq, c(0.5)), upper=quantile(freq, c(0.95))
  mutate(gender=ifelse(gender==0, "Men", "Women"))
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
gtab$freq<-1
gtab3<-cbind(gtab, model="Hurdle")</pre>
gtab<-rbind(gtab1, gtab2, gtab3)</pre>
gtab <- gtab %>%
   mutate(model=factor(model, levels=c("Poisson", "ZIP", "Hurdle")))
rapipl <- rapi.saemix %>%
      group_by(time, gender) %>%
   summarise(nev = sum(y0), n=n()) %>%
   mutate(freq = nev/n, sd=sqrt((1-nev/n)/nev)) %>%
   mutate(lower=freq-1.96*sd, upper=freq+1.96*sd)
## `summarise()` has grouped output by 'time'. You can override using the `.groups` argument.
rapipl$lower[rapipl$lower<0] <-0 # we should use a better approximation for CI
# Table form - compare to column B in Table 2
yfit0<-hurdlefit0
yfit1<-hurdlefit1
rr.tab<-data.frame(param=c("intercept", "beta.Male.inter", "slope", "beta.Male.slope", "omega.inter", "onega.inter", "onega.in
                              poissonNoZero=c(yfit1@results@fixed.effects, c(sqrt(diag(yfit1@results@omega)))),
                              logistic=c(yfit0@results@fixed.effects, c(sqrt(diag(yfit0@results@omega)))))
print(rr.tab)
```

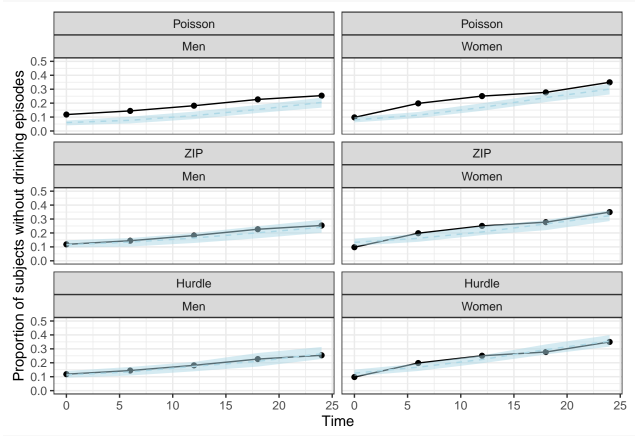
```
##
               param poissonNoZero
                                      logistic
## 1
           intercept
                       1.865583452 -2.79604024
                      -0.197211376 0.13215067
## 2 beta.Male.inter
                      -0.005943599
## 3
                                    0.03642832
               slope
## 4 beta.Male.slope -0.008525854
                                    0.02950090
## 5
         omega.inter
                       0.774608111
                                    1.55026563
## 6
         omega.slope
                       0.041313987
                                    0.07889691
```

Comparing the proportion of 0's for the different models

Clear model misfit for Poisson, much better for the other models with a slight advantage to Hurdle (?).

```
plot2 <- ggplot(rapipl, aes(x=time, y=freq, group=gender)) + geom_line() +
    geom_point() +
    geom_line(data=gtab, aes(x=time, y=median, group=gender), linetype=2, colour='lightblue') +
    geom_ribbon(data=gtab,aes(ymin=lower, ymax=upper, group=gender), alpha=0.5, fill='lightblue') +
    ylim(c(0,0.5)) + theme_bw() + theme(legend.position = "none") + facet_wrap(model~gender, ncol=2) +
    xlab("Time") + ylab("Proportion of subjects without drinking episodes")

print(plot2)</pre>
```



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

Other diagnostics

Plot evolution of median score? VPC on selected scores or on categories (more complicated and not saemix, more using R code with the simulated data)

Summarising all models in a LaTeX table

```
# running fim.saemix to extract the parameters with their name
y1<-fim.saemix(poisson.fit.cov2)
## Error in solve.default(F0) :
     routine Lapack dgesv : le système est exactement singulier : U[2,2] = 0
y2<-fim.saemix(zippoisson.fit.cov2)
## Error in solve.default(F0) :
     routine Lapack dgesv : le système est exactement singulier : U[2,2] = 0
y3<-fim.saemix(hurdlefit1)
## Error in solve.default(F0) :
     routine Lapack dgesv : le système est exactement singulier : U[2,2] = 0
parnam<-as.character(y2@results@conf.int$name)</pre>
estpar<-data.frame(parameter=parnam, poisson=y1@results@conf.int$estimate[match(parnam,y1@results@conf.
           zip=y2@results@conf.int$estimate[match(parnam,y2@results@conf.int$name)],
           hurdle=y3@results@conf.int$estimate[match(parnam,y3@results@conf.int$name)])
estpar[estpar$parnam=="p0", 4]<-hurdlefit0@results@fixed.effects[3]
estpar<-estpar[-c(6:7),]
for(icol in 2:4)
  estpar[,icol] <-format(estpar[,icol], digits=1, ns=1)</pre>
print(estpar)
##
                  parameter poisson
                                      zip hurdle
                               1.68 1.77 1.866
## 1
                  intercept
## 2 beta_gender(intercept)
                              -0.20 -0.20 -0.197
                      slope
                              -0.02 -0.02 -0.006
## 3
## 4
         beta_gender(slope)
                              -0.02 -0.02 -0.009
                                 NA 0.08
## 5
                                               NA
```

Time-to-event

8

9

TTE model - lung cancer

- create dataset for saemix (once) using the lung data from the survival package [see examplesDocumentation.R]
- changes
 - saemix format: added time=0

SD.intercept

SD.slope

- created one column for status (dead or alive, recoded as 1/0) and one for censoring (0/1)
- removed subjects with missing age, institution, sex, or ph scores (ecog and karno)

0.96 0.88 0.775

0.06 0.06 0.041

Checks

- The Surv function from the survival package creates a survival object for use as the response in a model formula.
 - one entry for each subject that is the survival time, which is followed by a + if the subject was censored
 - transform lung.saemix in the Surv format to check the survival function w/r saemix fit
- Weibull model
 - parametric survival function

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

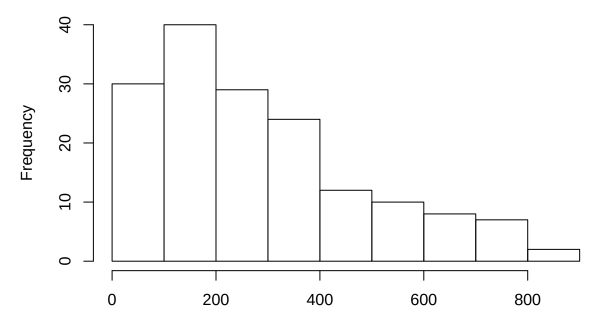
• Also tried computing a SE for S(t) using the delta-method where the vector of derivatives for the survival function of Weibull model are:

$$\begin{pmatrix} \frac{\delta S}{\delta \hat{\lambda}} \\ \frac{\delta S}{\delta \hat{\beta}} \end{pmatrix} = \begin{pmatrix} \frac{\beta}{\lambda} \left(\frac{t}{\lambda} \right)^{\beta} e^{-\left(\frac{t}{\lambda} \right)^{\beta}} \\ -\ln\left(\frac{t}{\lambda} \right) \left(\frac{t}{\lambda} \right)^{\beta} e^{-\left(\frac{t}{\lambda} \right)^{\beta}} \end{pmatrix}$$

• works pretty well compared to the non-parametric KM estimate

```
if(testMode)
  data(lung.saemix) else
    lung.saemix<-read.table(file.path(datDir, "lung.saemix.tab"), header=TRUE)
hist(lung.saemix$time[lung.saemix$status==1])</pre>
```

Histogram of lung.saemix\$time[lung.saemix\$status == 1]



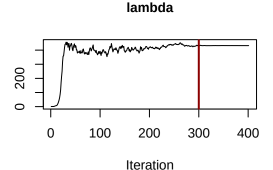
lung.saemix\$time[lung.saemix\$status == 1]

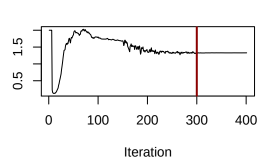
```
name.covariates=c("age", "sex", "ph.ecog", "ph.karno", "pat.karno", "wt.loss", "meal.cal"),
      units=list(x="days",y="",covariates=c("yr","","-","%","%","cal","pounds")))
## [1] "age"
                    "sex"
                                 "ph.ecog"
                                             "ph.karno" "pat.karno" "wt.loss"
## [7] "meal.cal"
##
##
## The following SaemixData object was successfully created:
## Object of class SaemixData
##
       longitudinal data for use with the SAEM algorithm
## Dataset lung.saemix
       Structured data: status ~ time + status + cens | id
##
##
       X variable for graphs: time (days)
       covariates: age (yr), sex (), ph.ecog (-), ph.karno (%), pat.karno (%), wt.loss (cal), meal.cal
##
##
         reference class for covariate sex : 0
weibulltte.model<-function(psi,id,xidep) {</pre>
  T < -xidep[,1]
  y<-xidep[,2] # events (1=event, 0=no event)
  cens<-which(xidep[,3]==1) # censoring times (subject specific)</pre>
  init <- which(T==0)</pre>
  lambda <- psi[id,1] # Parameters of the Weibull model</pre>
  beta <- psi[id,2]
  Nj <- length(T)
  ind <- setdiff(1:Nj, append(init,cens)) # indices of events</pre>
  hazard <- (beta/lambda)*(T/lambda)^(beta-1) # ln(H')
  H \leftarrow (T/lambda)^beta # ln(H)
  logpdf \leftarrow \text{rep}(0,Nj) \# ln(l(T=0))=0
  logpdf[cens] <- -H[cens] + H[cens-1] # ln(l(T=censoring time))</pre>
  logpdf[ind] <- -H[ind] + H[ind-1] + log(hazard[ind]) # ln(l(T=event time))</pre>
  return(logpdf)
}
saemix.model<-saemixModel(model=weibulltte.model,description="time model",modeltype="likelihood",</pre>
  psi0=matrix(c(1,2),ncol=2,byrow=TRUE,dimnames=list(NULL, c("lambda","beta"))),
 transform.par=c(1,1),covariance.model=matrix(c(1,0,0,0),ncol=2, byrow=TRUE))
##
##
## The following SaemixModel object was successfully created:
##
## Nonlinear mixed-effects model
##
     Model function: time model
     Model type: likelihood
##
## 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)</pre>
##
     init <- which(T==0)</pre>
##
     lambda <- psi[id,1] # Parameters of the Weibull model</pre>
##
     beta <- psi[id,2]
     Nj <- length(T)
##
```

```
##
##
     ind <- setdiff(1:Nj, append(init,cens)) # indices of events</pre>
     hazard <- (beta/lambda)*(T/lambda)^(beta-1) # ln(H')
##
     H <- (T/lambda)^beta # ln(H)</pre>
##
##
     logpdf \leftarrow rep(0,Nj) # ln(1(T=0))=0
##
     logpdf[cens] <- -H[cens] + H[cens-1] # ln(1(T=censoring time))</pre>
     logpdf[ind] <- -H[ind] + H[ind-1] + log(hazard[ind]) # ln(l(T=event time))</pre>
##
##
     return(logpdf)
## }
##
     Nb of parameters: 2
##
         parameter names: lambda beta
##
         distribution:
##
        Parameter Distribution Estimated
                  log-normal
## [1,] lambda
                               Estimated
## [2,] beta
                  log-normal
                               Estimated
##
     Variance-covariance matrix:
##
          lambda beta
## lambda
             1
## beta
      No covariate in the model.
##
##
       Initial values
##
               lambda beta
## Pop.CondInit
                     1
saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, displayProgress=FALSE)</pre>
tte.fit<-saemix(saemix.model,saemix.data,saemix.options)
## Nonlinear mixed-effects model fit by the SAEM algorithm
## -----
                Data
## -----
## Object of class SaemixData
       longitudinal data for use with the SAEM algorithm
## Dataset lung.saemix
##
       Structured data: status ~ time + status + cens | id
##
       X variable for graphs: time (days)
##
       covariates: age (yr), sex (), ph.ecog (-), ph.karno (%), pat.karno (%), wt.loss (cal), meal.cal
         reference class for covariate sex : 0
##
## Dataset characteristics:
      number of subjects:
       number of observations: 450
##
       average/min/max nb obs: 2.00 / 2 / 2
##
## First 10 lines of data:
##
      id time status cens status.1 age sex ph.ecog ph.karno pat.karno wt.loss
## 1
      1
           0
                   0
                        0
                                 0
                                    74
                                         0
                                                 1
                                                         90
                                                                  100
                                                                           NA
## 2
      1 306
                        0
                                 1 74
                                                         90
                                                                  100
                                                                           NΑ
                   1
                                         0
                                                 1
## 3
      2
           0
                        0
                                 0
                                    68
                                                 0
                                                         90
                                                                   90
                                                                           15
## 4
      2 455
                                    68
                                                         90
                                                                   90
                                                                           15
                   1
                        0
                                 1
                                         0
                                                 0
## 5
      3
            0
                   0
                        0
                                 0
                                    56
                                         0
                                                 0
                                                         90
                                                                   90
                                                                           15
## 6
      3 1010
                   0
                                 Ω
                                    56
                                         0
                                                 0
                                                         90
                                                                   90
                                                                           15
                        1
## 7
                   0
                        0
                                 0
                                    57
                                                         90
                                                                   60
                                                                           11
           0
                                                 1
## 8
      4
         210
                                    57
                                                                   60
                                                                           11
                   1
                        0
                                 1
                                         0
                                                 1
                                                        90
## 9
      5
                        0
                                 0
                                    60
                                        0
                                                                   90
                                                                            0
          Ο
                   0
                                                 Ω
                                                        100
## 10 5 883
                                                                   90
                   1
                        0
                                 1 60
                                                 0
                                                        100
     meal.cal mdv cens.1 occ ytype
```

```
## 1
                      0 1
      1175 0
                        1
## 2
        1175 0
                      0
## 3
       1225 0
                      0 1
## 4
        1225 0
                      0 1
## 5
         NA O
                      0
                        1
## 6
         NA O
                    0 1
## 7
       1150 0
                    0 1
       1150 0
## 8
                    0 1
        NA O
## 9
                      0 1
## 10
         NA O
                      0 1
             Model
## -----
## Nonlinear mixed-effects model
    Model function: time model
##
    Model type: likelihood
## 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)</pre>
##
##
    init <- which(T==0)</pre>
##
    lambda <- psi[id,1] # Parameters of the Weibull model</pre>
    beta <- psi[id,2]
##
##
    Nj <- length(T)
##
##
    ind <- setdiff(1:Nj, append(init,cens)) # indices of events</pre>
##
    hazard <- (beta/lambda)*(T/lambda)^(beta-1) # ln(H')</pre>
##
    H <- (T/lambda)^beta # ln(H)</pre>
##
    logpdf \leftarrow rep(0,Nj) # ln(1(T=0))=0
    logpdf[cens] <- -H[cens] + H[cens-1] # ln(l(T=censoring time))</pre>
    logpdf[ind] <- -H[ind] + H[ind-1] + log(hazard[ind]) # ln(l(T=event time))</pre>
##
##
    return(logpdf)
## }
## <bytecode: 0x56139cb11568>
##
    Nb of parameters: 2
##
        parameter names: lambda beta
##
        distribution:
##
       Parameter Distribution Estimated
## [1,] lambda log-normal Estimated
## [2,] beta
               log-normal Estimated
    Variance-covariance matrix:
##
        lambda beta
## lambda 1 0
## beta
             0
                  0
      No covariate in the model.
##
      Initial values
##
              lambda beta
## Pop.CondInit 1 2
## -----
         Key algorithm options ----
      Estimation of individual parameters (MAP)
##
##
      Estimation of standard errors and linearised log-likelihood
      Estimation of log-likelihood by importance sampling
##
```

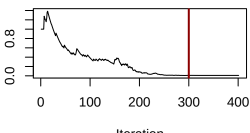
```
##
    Number of iterations: K1=300, K2=100
##
    Number of chains: 1
##
    Seed: 632545
##
    Number of MCMC iterations for IS: 5000
##
##
       nb of simulated datasets used for npde: 1000
##
       nb of simulated datasets used for VPC: 100
##
    Input/output
##
       save the results to a file: FALSE
##
       save the graphs to files: FALSE
    _____
## ----
                 Results
## -----
## ----- Fixed effects -----
## -----
     Parameter Estimate SE CV(%)
##
## [1,] lambda 431.8 51.60 12
## [2,] beta 1.3 0.19 14
## -----
## ----- Variance of random effects -----
               Estimate SE CV(%)
      Parameter
## lambda omega2.lambda 0.009 0.17 1858
## -----
## ----- Correlation matrix of random effects -----
## -----
##
           omega2.lambda
## omega2.lambda 1
## ------ Statistical criteria ------
## -----
## Likelihood computed by linearisation
##
      -2LL= 5189.352
##
      AIC = 5197.352
##
      BIC = 5211.017
##
## Likelihood computed by importance sampling
##
      -2LL= 2269.357
##
      AIC = 2277.357
##
      BIC = 2291.021
plot(tte.fit, plot.type="convergence")
```





beta

omega2.lambda



```
Iteration
```

```
ypred<-predict(tte.fit)</pre>
# Use survival package to assess Survival curve
if(TRUE) {
  library(survival)
  lung.surv<-lung.saemix[lung.saemix$time>0,]
  lung.surv$status<-lung.surv$status+1</pre>
  Surv(lung.surv$time, lung.surv$status) # 1=censored, 2=dead
  f1 <- survfit(Surv(time, status) ~ 1, data = lung.surv)</pre>
  xtim<-seq(0,max(lung.saemix$time), length.out=200)</pre>
  estpar<-tte.fit@results@fixed.effects
  estse<-tte.fit@results@se.fixed
  ypred<-exp(-(xtim/estpar[1])^(estpar[2]))</pre>
# Computing SE for the survival curve based on linearised FIM (probably not a good idea) through the de
  invfim<-solve(tte.fit@results@fim[1:2,1:2])</pre>
  xcal<- (xtim/estpar[1])^estpar[2]</pre>
  dsdbeta<- -log(xtim/estpar[1]) * xcal *exp(-xcal)</pre>
  dsdalpha<- estpar[2]/estpar[1] * xcal *exp(-xcal)</pre>
  xmat<-rbind(dsdalpha, dsdbeta)</pre>
       x1 < -t(xmat[,1:3]) %*% invfim %*% xmat[,1:3]
  sesurv<-rep(0,length(xcal))</pre>
  for(i in 1:length(xcal))
    sesurv[i]<-sqrt(t(xmat[,i]) %*% invfim %*% xmat[,i])</pre>
  if(saveForDocs) {
    namfile<-"lung_compareKM.eps"</pre>
    postscript(file.path(figDir, namfile), horizontal=TRUE)
    plot(f1, xlab = "Days", ylab = "Overall survival probability")
    lines(xtim,ypred, col="red",lwd=2)
```

```
lines(xtim,ypred+1.96*sesurv, col="red",lwd=1, lty=2)
lines(xtim,ypred-1.96*sesurv, col="red",lwd=1, lty=2)
dev.off()
}

# ypred2<-exp(-(xtim/(estpar[1]-1.96*sqrt(invfim[1,1])))^(estpar[2]+1.96*sqrt(invfim[2,2])))
# ypred3<-exp(-(xtim/(estpar[1]+1.96*sqrt(invfim[1,1])))^(estpar[2]+1.96*sqrt(invfim[2,2])))
# lines(xtim,ypred2, col="blue",lwd=1, lty=2)
# lines(xtim,ypred3, col="blue",lwd=1, lty=2)
}</pre>
```

RTTE model

- again difficult to find real data
- simulated data
 - Exemple simulé de Belhal **TODO**
 - data from the Monolix documentation: absolutely no indication where the data comes from (weibull data.txt for the weibullRTTE.mlxtran project in the demo)
- search for real data
 - asked Ulrika Simonsson for the RTTE data on post-operative pain (Pain Medicine 2015)
 - data on events in Gaucher disease used for the ENSAI workshops (but few events)
 - discretised PCA events during warfarin treatment? (from the warfarin PK/PD) (but threshold?)

```
# Simulating RTTE data by simulating from U(0,1) and inverting the cdf
simul.rtte.unif <- function(psi) { # xidep, id not important, we only use psi
  censoringtime <- 3
  maxevents <- 30
  lambda <- psi[,1]</pre>
  beta <- psi[,2]
  simdat<-NULL
  N<-nrow(psi)
  for(i in 1:N) {
    eventTimes<-c(0)
    T<-0
    Vj<-runif(1)</pre>
          T \leftarrow (-\log(Vj)*lambda[i]) \cap (beta[i])
    T \leftarrow lambda[i] * (-log(Vj))^(1/beta[i])
    nev<-0
    while (T < censoringtime & nev<maxevents){</pre>
      eventTimes <- c(eventTimes, T)</pre>
      nev<-nev+1
      Vj<-runif(1)</pre>
              T \leftarrow T + (-\log(Vj) * lambda[i]) \hat{beta[i]}
              T < -(-log(Vj)*lambda[i] + T^(1/beta[i]))^(beta[i])
      T<-lambda[i]*(-log(Vj) + (T/lambda[i])^(beta[i]))^(1/beta[i])
    if(nev==maxevents) {
      message("Reached maximum number of events\n")
    eventTimes<-c(eventTimes, censoringtime)</pre>
    cens<-rep(1,length(eventTimes))</pre>
    cens[1] <-cens[length(cens)] <-0
    simdat<-rbind(simdat,</pre>
```

```
data.frame(id=i, T=eventTimes, status=cens))
  }
  return(simdat)
# Subjects
set.seed(12345)
param < -c(2, 1.5, 0.5)
\# param < -c(4, 1.2, 0.3)
omega < -c(0.25, 0.25)
nsuj<-200
risk<-rep(0,nsuj)
risk[(nsuj/2+1):nsuj]<-1
psiM<-data.frame(lambda=param[1]*exp(rnorm(nsuj,sd=omega[1])), beta=param[2]*exp(param[3]*risk+rnorm(nsuj,sd=omega[1]))
simdat <- simul.rtte.unif(psiM)</pre>
## Reached maximum number of events
simdat$risk<-as.integer(simdat$id>(nsuj/2))
# Simulate T from Weibull (check)
if(FALSE) {
  lambda < -2
  beta<-2
  nsim<-5000
  # By hand
  q1<-runif(nsim)
  # tevent<-lambda*exp(log(q1)/beta)</pre>
  tevent<-lambda*exp(log(-log(q1))/beta)
  tevent<-sort(tevent)</pre>
# plot(tevent, exp(-(tevent/lambda) ^beta))
  tevent2<-sort(rweibull(nsim, shape=beta, scale=lambda))</pre>
  plot(tevent, tevent2)
  abline(0,1)
}
saemix.data<-saemixData(name.data=simdat, name.group=c("id"), name.predictors=c("T"), name.response="st</pre>
## [1] "risk"
##
##
## The following SaemixData object was successfully created:
##
## Object of class SaemixData
##
       longitudinal data for use with the SAEM algorithm
## Dataset simdat
##
       Structured data: status ~ T | id
       Predictor: T ()
##
       covariates: risk (-)
##
##
         reference class for covariate risk: 0
rtte.model<-function(psi,id,xidep) {</pre>
  T < -xidep[,1]
  N <- nrow(psi) # nb of subjects
```

```
Nj <- length(T) # nb of events (including 0 and censoring times)
  # censoringtime = 6
  censoringtime = max(T) # same censoring for everyone
  lambda <- psi[id,1]</pre>
  beta <- psi[id,2]
  tinit <- which(T==0) # indices of beginning of observation period
  tcens <- which (T == censoring time) # indices of censored events
  tevent <- setdiff(1:Nj, append(tinit,tcens)) # indices of non-censored event times
  hazard <- (beta/lambda)*(T/lambda)^(beta-1)
  H <- (T/lambda)^beta</pre>
  logpdf <- rep(0,Nj)</pre>
  logpdf[tcens] <- -H[tcens] + H[tcens-1]</pre>
  logpdf[tevent] <- -H[tevent] + H[tevent-1] + log(hazard[tevent])</pre>
  return(logpdf)
saemix.model.base<-saemixModel(model=rtte.model,description="Repeated TTE model",modeltype="likelihood"
                                psi0=matrix(c(1,2),ncol=2,byrow=TRUE,dimnames=list(NULL, c("lambda","be
                                transform.par=c(1,1),covariance.model=matrix(c(1,0,0,1),ncol=2, byrow=TR
##
##
## The following SaemixModel object was successfully created:
## Nonlinear mixed-effects model
##
    Model function: Repeated TTE model
##
    Model type: likelihood
## function(psi,id,xidep) {
    T < -xidep[,1]
##
    N <- nrow(psi) # nb of subjects
##
##
    Nj <- length(T) # nb of events (including 0 and censoring times)
##
     # censoringtime = 6
##
     censoringtime = max(T) # same censoring for everyone
     lambda <- psi[id,1]</pre>
##
     beta <- psi[id,2]
##
##
     tinit <- which(T==0) # indices of beginning of observation period
     tcens <- which(T==censoringtime) # indices of censored events
##
##
     tevent <- setdiff(1:Nj, append(tinit,tcens)) # indices of non-censored event times
     hazard <- (beta/lambda)*(T/lambda)^(beta-1)</pre>
##
##
    H <- (T/lambda)^beta</pre>
##
     logpdf <- rep(0,Nj)</pre>
     logpdf[tcens] <- -H[tcens] + H[tcens-1]</pre>
##
##
     logpdf[tevent] <- -H[tevent] + H[tevent-1] + log(hazard[tevent])</pre>
##
     return(logpdf)
## }
##
    Nb of parameters: 2
##
         parameter names: lambda beta
##
         distribution:
##
        Parameter Distribution Estimated
## [1,] lambda
                  log-normal
                                Estimated
## [2,] beta
                  log-normal
                                Estimated
     Variance-covariance matrix:
##
##
          lambda beta
## lambda
               1
```

```
## beta
               0
##
       No covariate in the model.
##
       Initial values
                lambda beta
##
## Pop.CondInit
saemix.model<-saemixModel(model=rtte.model,description="Repeated TTE model",modeltype="likelihood",</pre>
                           psi0=matrix(c(1,2),ncol=2,byrow=TRUE,dimnames=list(NULL, c("lambda","beta"))
                           transform.par=c(1,1),covariate.model=matrix(c(0,1),ncol=2),
                           covariance.model=matrix(c(1,0,0,1),ncol=2, byrow=TRUE))
##
##
## The following SaemixModel object was successfully created:
##
## Nonlinear mixed-effects model
     Model function: Repeated TTE model
##
     Model type: likelihood
## function(psi,id,xidep) {
     T<-xidep[,1]
##
     N <- nrow(psi) # nb of subjects
##
     Nj <- length(T) # nb of events (including 0 and censoring times)
##
##
     # censoringtime = 6
##
     censoringtime = max(T) # same censoring for everyone
##
     lambda <- psi[id,1]</pre>
     beta <- psi[id,2]
##
     tinit \leftarrow which(T==0) # indices of beginning of observation period
##
     tcens <- which(T==censoringtime) # indices of censored events
##
##
     tevent <- setdiff(1:Nj, append(tinit,tcens)) # indices of non-censored event times
##
     hazard <- (beta/lambda)*(T/lambda)^(beta-1)
##
    H <- (T/lambda)^beta</pre>
##
     logpdf <- rep(0,Nj)</pre>
##
     logpdf[tcens] <- -H[tcens] + H[tcens-1]</pre>
##
     logpdf[tevent] <- -H[tevent] + H[tevent-1] + log(hazard[tevent])</pre>
##
     return(logpdf)
## }
##
     Nb of parameters: 2
##
         parameter names: lambda beta
##
         distribution:
        Parameter Distribution Estimated
##
## [1,] lambda
                  log-normal
                                Estimated
## [2,] beta
                  log-normal
                                Estimated
     Variance-covariance matrix:
##
##
          lambda beta
## lambda
             1
## beta
     Covariate model:
        lambda beta
##
## [1,]
            0
##
       Initial values
                lambda beta
## Pop.CondInit
                     1
## Cov.CondInit
```

```
saemix.options<-list(seed=632545,save=FALSE,save.graphs=FALSE, fim=FALSE, displayProgress=FALSE)
rtte.fit<-saemix(saemix.model,saemix.data,saemix.options)</pre>
```

```
## Nonlinear mixed-effects model fit by the SAEM algorithm
## -----
         Data
## -----
## Object of class SaemixData
      longitudinal data for use with the SAEM algorithm
## Dataset simdat
##
     Structured data: status ~ T | id
##
     Predictor: T ()
##
     covariates: risk (-)
       reference class for covariate risk : 0
##
## Dataset characteristics:
##
     number of subjects:
                           200
##
      number of observations: 967
      average/min/max nb obs: 4.83 / 2 / 32
## First 10 lines of data:
              T status risk mdv cens occ ytype
## 1
    1 0.0000000 0 0
                            0
                               0
                                    1
## 2
     1 0.7520145
                    1
                         0
                             0
                                 0
                                     1
## 3 1 0.8775847
                   1 0 0
                               0 1
## 4 1 2.4331650
                   1 0 0
                               0 1
                       0 0
                                0 1
## 5
    1 3.0000000
                    0
## 6 2 0.0000000
                     0 0 0
                               0 1 1
## 7 2 1.3712351
                   1 0 0 0 1
## 8 2 3.0000000
                   0 0 0 0 1
## 9 3 0.0000000
                     0
                        0 0
## 10 3 2.8564910
                   1 0 0
                                          1
## ----
             Model
## -----
## Nonlinear mixed-effects model
    Model function: Repeated TTE model
    Model type: likelihood
##
## function(psi,id,xidep) {
##
    T < -xidep[,1]
    N <- nrow(psi) # nb of subjects
##
    Nj <- length(T) # nb of events (including 0 and censoring times)
##
    # censoringtime = 6
##
    censoringtime = max(T) # same censoring for everyone
##
    lambda <- psi[id,1]</pre>
    beta <- psi[id,2]
##
##
    tinit <- which(T==0) # indices of beginning of observation period
##
    tcens <- which(T==censoringtime) # indices of censored events
##
    tevent <- setdiff(1:Nj, append(tinit,tcens)) # indices of non-censored event times
##
    hazard <- (beta/lambda)*(T/lambda)^(beta-1)
##
    H <- (T/lambda)^beta</pre>
##
    logpdf <- rep(0,Nj)</pre>
##
    logpdf[tcens] <- -H[tcens] + H[tcens-1]</pre>
##
    logpdf[tevent] <- -H[tevent] + H[tevent-1] + log(hazard[tevent])</pre>
##
    return(logpdf)
## }
```

```
## <bytecode: 0x561396909538>
##
   Nb of parameters: 2
##
      parameter names: lambda beta
##
      distribution:
     Parameter Distribution Estimated
## [1,] lambda log-normal Estimated
## [2,] beta log-normal Estimated
  Variance-covariance matrix:
##
      lambda beta
## lambda
        1 0
## beta
##
  Covariate model:
     [,1] [,2]
## risk 0 1
##
     Initial values
##
           lambda beta
## Pop.CondInit
             1
## Cov.CondInit
## -----
       Key algorithm options ----
 _____
     Estimation of individual parameters (MAP)
     Estimation of log-likelihood by importance sampling
##
##
     Number of iterations: K1=300, K2=100
    Number of chains: 1
##
##
     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 a file: FALSE
##
        save the graphs to files: FALSE
                  Results
## -----
## ----- Fixed effects -----
## -----
##
     Parameter
                 Estimate
## [1,] lambda
                 2.1
## [2,] beta
## [3,] beta_risk(beta) 0.4
## -----
## ----- Variance of random effects -----
## -----
      Parameter Estimate
##
## lambda omega2.lambda 0.1125
## beta omega2.beta 0.0015
## -----
## ----- Correlation matrix of random effects -----
## -----
            omega2.lambda omega2.beta
## omega2.lambda 1
## omega2.beta 0
```

```
----- Statistical criteria ------
##
## Likelihood computed by importance sampling
          -2LL= 690.2485
##
          AIC = 702.2485
##
          BIC = 722.0384
##
plot(rtte.fit, plot.type="convergence")
           lambda
                                                                            beta_risk(beta)
                                              beta
1.8
4.
1.0
                                                                    0.0
    0
        100
            200
                 300
                      400
                                      0
                                          100
                                              200
                                                   300
                                                        400
                                                                        0
                                                                            100
                                                                                 200
                                                                                     300
                                                                                          400
           Iteration
                                             Iteration
                                                                               Iteration
        omega2.lambda
                                           omega2.beta
9.0
                                  9.7
0.2
                                  0.0
            200
                                              200
    0
        100
                 300
                      400
                                      0
                                          100
                                                   300
                                                        400
```

Exiting

Iteration

```
if(testMode) {
  dev_mode()
}
```

Iteration