Package 'GLMMadaptive'

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Title Generalized Linear Mixed Models using Adaptive Gaussian Quadrature

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BugReports https://github.com/drizopoulos/GLMMadaptive/issues

Description Fits generalized linear mixed models for a single grouping factor under maximum likelihood approximating the integrals over the random effects with an adaptive Gaussian quadrature rule; Jose C. Pinheiro and Douglas M. Bates (1995) <doi:10.1080/10618600.1995.10474663>.

Imports MASS, nlme, parallel, matrixStats

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Continuation Ratio Set-Up

Functions to Set-Up Data for a Continuation Ratio Mixed Model

Description

Data set-up and calculation of marginal probabilities from a continuation ratio model

Usage

cr_setup(y, direction = c("forward", "backward"))

```
cr_marg_probs(eta, direction = c("forward", "backward"))
```

Arguments

У	a numeric vector denoting the ordinal response variable.
direction	character string specifying the direction of the continuation ratio model; "forward" corresponds to a discrete hazard function.
eta	a numeric matrix of the linear predictor, with columns corresponding to the different levels of the ordinal response.

Note

Function cr_setup() is based on the cr.setup() function from package rms.

Author(s)

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Examples

```
n <- 300 # number of subjects</pre>
K <- 8 # number of measurements per subject
t_max <- 15 # maximum follow-up time</pre>
# we constuct a data frame with the design:
# everyone has a baseline measurment, and then measurements at random follow-up times
DF <- data.frame(id = rep(seq_len(n), each = K),</pre>
                 time = c(replicate(n, c(0, sort(runif(K - 1, 0, t_max))))),
                 sex = rep(gl(2, n/2, labels = c("male", "female")), each = K))
# design matrices for the fixed and random effects
X <- model.matrix(~ sex * time, data = DF)[, -1]
Z <- model.matrix(~ 1, data = DF)</pre>
thrs <- c(-1.5, 0, 0.9) # thresholds for the different ordinal categories
betas <- c(-0.25, 0.24, -0.05) # fixed effects coefficients
D11 <- 0.48 # variance of random intercepts
D22 <- 0.1 # variance of random slopes
# we simulate random effects
b <- cbind(rnorm(n, sd = sqrt(D11)), rnorm(n, sd = sqrt(D22)))[, 1, drop = FALSE]</pre>
# linear predictor
eta_y <- drop(X %*% betas + rowSums(Z * b[DF$id, , drop = FALSE]))</pre>
# linear predictor for each category
eta_y <- outer(eta_y, thrs, "+")</pre>
# marginal probabilities per category
mprobs <- cr_marg_probs(eta_y)</pre>
# we simulate ordinal longitudinal data
DF$y <- unname(apply(mprobs, 1, sample, x = ncol(mprobs), size = 1, replace = TRUE))
# If you want to simulate from the backward formulation of the CR model, you need to
# change `eta_y <- outer(eta_y, thrs, "+")` to `eta_y <- outer(eta_y, rev(thrs), "+")`,
# and `mprobs <- cr_marg_probs(eta_y)` to `mprobs <- cr_marg_probs(eta_y, "backward")`
# prepare the data
# If you want to fit the CR model under the backward formulation, you need to change
# `cr_vals <- cr_setup(DF$y)` to `cr_vals <- cr_setup(DF$y, "backward")`</pre>
cr_vals <- cr_setup(DF$y)</pre>
cr_data <- DF[cr_vals$subs, ]</pre>
cr_data$y_new <- cr_vals$y</pre>
cr_data$cohort <- cr_vals$cohort</pre>
# fit the model
fm <- mixed_model(y_new ~ cohort + sex * time, random = ~ 1 | id,</pre>
                  data = cr_data, family = binomial())
summary(fm)
```

effectPlotData

Description

Creates predicted values and their corresponding confidence interval for constructing an effects plot.

Usage

```
effectPlotData(object, newdata, level, ...)
## S3 method for class 'MixMod'
effectPlotData(object, newdata,
    level = 0.95, marginal = FALSE, CR_cohort_varname = NULL,
    direction = NULL, K = 200, seed = 1, sandwich = FALSE,
    ...)
```

Arguments

object	an object inheriting from class "MixMod".
newdata	a data frame base on which predictions will be calculated.
level	a numeric scalar denoting the level of the confidence interval.
marginal	logical; if FALSE predicted values are calculated for the "mean" subject (i.e., the one with random effects values equal to 0). When TRUE marginal predicted values are calculated using function marginal_coefs.
CR_cohort_varna	ame
	a character string denoting the name of the cohort variable when a continuation ratio model is fitted.
direction	the direction argument of cr_marg_probs needs to be provided when CR_cohort_varname is not NULL.
К	numeric scalar denoting the number of Monte Carlo samples from the approxi- mate posterior of the parameters; applicable only for zero-inflated models.
seed	numerical scalar giving the seed to be used in the Monte Carlo scheme.
sandwich	logical; if TRUE robust/sandwich standard errors are used in the calculations.
	additional arguments passed to marginal_coefs, e.g., cores.

Details

The confidence interval is calculated based on a normal approximation.

Value

The data frame newdata with extra columns pred, low and upp.

effectPlotData

Author(s)

Dimitris Rizopoulos <d.rizopoulos@erasmusmc.nl>

See Also

mixed_model, marginal_coefs

Examples

```
# simulate some data
set.seed(123L)
n <- 500
K <- 15
t.max <- 25
betas <- c(-2.13, -0.25, 0.24, -0.05)
D <- matrix(0, 2, 2)
D[1:2, 1:2] <- c(0.48, -0.08, -0.08, 0.18)
times <- c(replicate(n, c(0, sort(runif(K-1, 0, t.max)))))</pre>
group <- sample(rep(0:1, each = n/2))</pre>
DF <- data.frame(year = times, group = factor(rep(group, each = K)))</pre>
X <- model.matrix(~ group * year, data = DF)</pre>
Z <- model.matrix(~ year, data = DF)</pre>
b <- cbind(rnorm(n, sd = sqrt(D[1, 1])), rnorm(n, sd = sqrt(D[2, 2])))</pre>
id <- rep(1:n, each = K)
eta.y <- as.vector(X %*% betas + rowSums(Z * b[id, ]))</pre>
DF$y <- rbinom(n * K, 1, plogis(eta.y))</pre>
DF$id <- factor(id)</pre>
# Fit a model
fm1 <- mixed_model(fixed = y ~ year * group, random = ~ year | id, data = DF,</pre>
                   family = binomial())
# An effects plot for the mean subject (i.e., with random effects equal to 0)
nDF <- with(DF, expand.grid(year = seq(min(year), max(year), length.out = 15),
    group = levels(group)))
plot_data <- effectPlotData(fm1, nDF)</pre>
require("lattice")
xyplot(pred + low + upp ~ year | group, data = plot_data,
       type = "l", lty = c(1, 2, 2), col = c(2, 1, 1), lwd = 2,
       xlab = "Follow-up time", ylab = "log odds")
expit <- function (x) exp(x) / (1 + exp(x))
xyplot(expit(pred) + expit(low) + expit(upp) ~ year | group, data = plot_data,
       type = "l", lty = c(1, 2, 2), col = c(2, 1, 1), lwd = 2,
       xlab = "Follow-up time", ylab = "Probabilities")
```

```
# we put the two groups in the same panel
my.panel.bands <- function(x, y, upper, lower, fill, col, subscripts, ..., font,</pre>
                           fontface) {
    upper <- upper[subscripts]</pre>
    lower <- lower[subscripts]</pre>
   panel.polygon(c(x, rev(x)), c(upper, rev(lower)), col = fill, border = FALSE, ...)
}
xyplot(expit(pred) ~ year, group = group, data = plot_data, upper = expit(plot_data$upp),
       low = expit(plot_data$low), type = "l", col = c("blue", "red"),
       fill = c("#0000FF80", "#FF000080"),
       panel = function (x, y, ...) {
           panel.superpose(x, y, panel.groups = my.panel.bands, ...)
           panel.xyplot(x, y, lwd = 2, ...)
}, xlab = "Follow-up time", ylab = "Probabilities")
# An effects plots for the marginal probabilities
plot_data_m <- effectPlotData(fm1, nDF, marginal = TRUE, cores = 1L)</pre>
expit <- function (x) exp(x) / (1 + exp(x))
xyplot(expit(pred) + expit(low) + expit(upp) ~ year | group, data = plot_data_m,
       type = "l", lty = c(1, 2, 2), col = c(2, 1, 1), lwd = 2,
       xlab = "Follow-up time", ylab = "Probabilities")
```

Extra Family Objects	Family functions for Student's-t, Beta, Zero-Inflated and Hurdle
	Poisson and Negative Binomial, Hurdle Log-Normal, Hurdle Beta,
	Gamma, and Censored Normal Mixed Models

Description

Specifies the information required to fit a Beta, zero-inflated and hurdle Poisson, zero-inflated and hurdle Negative Binomial, a hurdle normal and a hurdle Beta mixed-effects model, using mixed_model().

Usage

```
students.t(df = stop("'df' must be specified"), link = "identity")
beta.fam()
zi.poisson()
zi.binomial()
zi.negative.binomial()
hurdle.poisson()
hurdle.negative.binomial()
hurdle.lognormal()
hurdle.beta.fam()
unit.lindley()
```

Extra Family Objects

```
beta.binomial(link = "logit")
Gamma.fam()
censored.normal()
```

Arguments

link	name of the link function.
df	the degrees of freedom of the Student's t distribution.

Note

Currently only the log-link is implemented for the Poisson, negative binomial and Gamma models, the logit link for the beta and hurdle beta models and the identity link for the log-normal model.

Examples

```
# simulate some data from a negative binomial model
set.seed(102)
dd <- expand.grid(f1 = factor(1:3), f2 = LETTERS[1:2], g = 1:30, rep = 1:15,
                KEEP.OUT.ATTRS = FALSE)
mu <- 5*(-4 + with(dd, as.integer(f1) + 4 * as.numeric(f2)))</pre>
dd$y <- rnbinom(nrow(dd), mu = mu, size = 0.5)
# Fit a zero-inflated Poisson model, with only fixed effects in the
# zero-inflated part
fm1 <- mixed_model(fixed = y ~ f1 * f2, random = ~ 1 | g, data = dd,</pre>
                family = zi.poisson(), zi_fixed = ~ 1)
summary(fm1)
# We extend the previous model allowing also for a random intercept in the
# zero-inflated part
fm2 \le mixed_model(fixed = y \sim f1 * f2, random = \sim 1 | g, data = dd,
               family = zi.poisson(), zi_fixed = ~ 1, zi_random = ~ 1 | g)
# We do a likelihood ratio test between the two models
anova(fm1, fm2)
# The same as above but with a negative binomial model
gm1 <-mixed_model(fixed = y ~ f1 * f2, random = ~ 1 | g, data = dd,
               family = zi.negative.binomial(), zi_fixed = ~ 1)
summary(gm1)
```

We do a likelihood ratio test between the Poisson and negative binomial models anova(fm1, gm1) GLMMadaptive

Description

This package fits generalized linear mixed models for a single grouping factor under maximum likelihood approximating the integrals over the random effects with an adaptive Gaussian quadrature rule.

Details

Package:	GLMMadaptive
Type:	Package
Version:	0.9-7
Date:	2025-03-04
License:	GPL (>=3)

This package fits mixed effects models for grouped / repeated measurements data for which the integral over the random effects in the definition of the marginal likelihood cannot be solved analytically. The package approximates these integrals using the adaptive Gauss-Hermite quadrature rule.

Multiple random effects terms can be included for the grouping factor (e.g., random intercepts, random linear slopes, random quadratic slopes), but currently only a single grouping factor is allowed.

The package also offers several utility functions that can extract useful information from fitted mixed effects models. The most important of those are included in the **See also** Section below.

Author(s)

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References

Fitzmaurice, G., Laird, N. and Ware J. (2011). *Applied Longitudinal Analysis*, 2nd Ed. New York: John Wiley & Sons.

Molenberghs, G. and Verbeke, G. (2005). *Models for Discrete Longitudinal Data*. New York: Springer-Verlag.

See Also

mixed_model, methods.MixMod, effectPlotData, marginal_coefs

marginal_coefs

Description

Calculates marginal coefficients and their standard errors from fitted generalized linear mixed models.

Usage

```
marginal_coefs(object, ...)
```

```
## S3 method for class 'MixMod'
marginal_coefs(object, std_errors = FALSE,
    link_fun = NULL, M = 3000, K = 100, seed = 1,
    cores = max(parallel::detectCores() - 1, 1),
    sandwich = FALSE, ...)
```

Arguments

object	an object inheriting from class "MixMod".
std_errors	logical indicating whether standard errors are to be computed.
link_fun	a function transforming the mean of the repeated measurements outcome to the linear predictor scale. Typically, this derived from the family argument of mixed_model.
Μ	numeric scalar denoting the number of Monte Carlo samples.
К	numeric scalar denoting the number of samples from the sampling distribution of the maximum likelihood estimates.
seed	integer denoting the seed for the random number generation.
cores	integer giving the number of cores to use; applicable only when std_errors = TRUE.
sandwich	logical; if TRUE robust/sandwich standard errors are used in the calculations.
	extra arguments; currently none is used.

Details

It uses the approach of Hedeker et al. (2017) to calculate marginal coefficients from mixed models with nonlinear link functions. The marginal probabilities are calculated using Monte Carlo integration over the random effects with M samples, by sampling from the estimated prior distribution, i.e., a multivariate normal distribution with mean 0 and covariance matrix \hat{D} , where \hat{D} denotes the estimated covariance matrix of the random effects.

To calculate the standard errors, the Monte Carlo integration procedure is repeated K times, where each time instead of the maximum likelihood estimates of the fixed effects and the covariance matrix of the random effects, a realization is used from the sampling distribution of the maximum likelihood estimates. To speed-up this process, package **parallel** is used.

A list of class "m_coefs" with components betas the marginal coefficients, and when std_errors = TRUE, the extra components var_betas the estimated covariance matrix of the marginal coefficients, and coef_table a numeric matrix with the estimated marginal coefficients, their standard errors and corresponding p-values using the normal approximation.

Author(s)

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References

Hedeker, D., du Toit, S. H., Demirtas, H. and Gibbons, R. D. (2018), A note on marginalization of regression parameters from mixed models of binary outcomes. *Biometrics* **74**, 354–361. doi:10.1111/biom.12707

See Also

mixed_model

Examples

```
# simulate some data
set.seed(123L)
n <- 500
K <- 15
t.max <- 25
betas <- c(-2.13, -0.25, 0.24, -0.05)
D <- matrix(0, 2, 2)
D[1:2, 1:2] <- c(0.48, -0.08, -0.08, 0.18)
times <- c(replicate(n, c(0, sort(runif(K-1, 0, t.max)))))</pre>
group <- sample(rep(0:1, each = n/2))
DF <- data.frame(year = times, group = factor(rep(group, each = K)))</pre>
X <- model.matrix(~ group * year, data = DF)</pre>
Z <- model.matrix(~ year, data = DF)</pre>
b <- cbind(rnorm(n, sd = sqrt(D[1, 1])), rnorm(n, sd = sqrt(D[2, 2])))</pre>
id <- rep(1:n, each = K)
eta.y <- as.vector(X %*% betas + rowSums(Z * b[id, ]))</pre>
DF$y <- rbinom(n * K, 1, plogis(eta.y))</pre>
DF$id <- factor(id)</pre>
****
fm1 <- mixed_model(fixed = y \sim year * group, random = \sim 1 \mid id, data = DF,
                   family = binomial())
```

fixef(fm1)
marginal_coefs(fm1)

mixed_model

```
marginal_coefs(fm1, std_errors = TRUE, cores = 1L)
```

mixed_model

Generalized Linear Mixed Effects Models

Description

Fits generalized linear mixed effects models under maximum likelihood using adaptive Gaussian quadrature.

Usage

```
mixed_model(fixed, random, data, family, weights = NULL,
  na.action = na.exclude, zi_fixed = NULL, zi_random = NULL,
  penalized = FALSE, n_phis = NULL, initial_values = NULL,
  control = list(), ...)
```

Arguments

fixed	a formula for the fixed-effects part of the model, including the outcome.
random	a formula for the random-effects part of the model. This should only contain the right-hand side part, e.g., ~ time id, where time is a variable, and id the grouping factor. When the symbol is used in the definition of this argument (instead of), then the covariance matrix of the random effects is assumed to be diagonal.
data	a data.frame containing the variables required in fixed and random.
family	a family object specifying the type of the repeatedly measured response vari- able, e.g., binomial() or poisson(). The function also allows for user-defined family objects, but with specific extra components; see the example is negative.binomial for more details. Contrary to the standard practice in model fitting R functions with a family argument (e.g., glm) in which the default family is gaussian(), in mixed_model() no default is provided. If the users wish to fit a mixed model for a Gaussian outcome, this could be done with function lme() from the nlme package or function lmer() from the lme4 package.
weights	a numeric vector of weights. These are simple multipliers on the log-likelihood contributions of each group/cluster, i.e., we presume that there are multiple replicates of each group/cluster denoted by the weights. The length of 'weights' need to be equal to the number of independent groups/clusters in the data.
na.action	what to do with missing values in data.
<pre>zi_fixed, zi_ra</pre>	ndom
	formulas for the fixed and random effects of the zero inflated part.
penalized	logical or a list. If logical and equal to FALSE, then no penalty is used. If logical and equal to TRUE, for the fixed effects a Student's-t penalty/prior with mean 0, scale equal to 1 and 3 degrees of freedom is used. If a list, then it is expected to have the components pen_mu, pen_sigma and pen_df, denoting the mean, scale and degrees of freedom of the Student's-t penalty/prior for the fixed effects.

- n_phis a numeric scalar; in case the family corresponds to a distribution that has extra (dispersion/shape) parameters, you need to specify how many extra parameters are needed.
- initial_values a list of initial values. This can have up to three components, namely,
 - **betas** a numeric vector of fixed effects. This can also be family object. In this case initial values for the fixed effects will be calculated by using glm to the data ignoring the correlations in the repeated measurements. For example, for a negative binomial response outcome, we could set betas = poisson().
 - **D** a numeric matrix denoting the covariance matrix of the random effects.
 - **phis** a numeric vector for the extra (dispersion/shape) parameters.
- control a list with the following components:
 - iter_EM numeric scalar denoting the number of EM iterations; default is 30.
 - **iter_qN_outer** numeric scalar denoting the number of outer iterations during the quasi-Newton phase; default is 15. In each outer iteration the locations of the quadrature points are updated.
 - iter_qN numeric scalar denoting the starting number of iterations for the quasi-Newton; default is 10.
 - iter_qN_incr numeric scalar denoting the increment in iter_qN for each outer iteration; default is 10.
 - **optimizer** character string denoting the optimizer to be used; available options are "optim" (default), "nlminb" and "optimParallel", the last option implemented in the **optimParallel** package.
 - **optim_method** character string denoting the type of **optim** algorithm to be used when optimizer = "optim"; default is the BFGS algorithm.
 - **parscale_betas** the control argument parscale of optim for the fixed-effects; default is 0.1.
 - **parscale_D** the control argument parscale of optim for the unique element of the covariance matrix of the random effects; default is 0.01.
 - **parscale_phis** the control argument parscale of optim for the extra (dispersion/shape) parameters; default is 0.01.
 - tol1, tol2, tol3 numeric scalars controlling tolerances for declaring convergence; tol1 and tol2 are for checking convergence in successive parameter values; tol3 is similar to reltop of optim; default values are 1e-03, 1e-04, and 1e-08, respectively.
 - **numeric_deriv** character string denoting the type of numerical derivatives to be used. Options are "fd" for forward differences, and cd for central difference; default is "fd".
 - **nAGQ** numeric scalar denoting the number of quadrature points; default is 11 when the number of random effects is one or two, and 7 otherwise.
 - **update_GH_every** numeric scalar denoting every how many iterations to update the quadrature points during the EM-phase; default is 10.
 - **max_coef_value** numeric scalar denoting the maximum allowable value for the fixed effects coefficients during the optimization; default is 10.

- **max_phis_value** numeric scalar denoting the maximum allowable value for the shape/dispersion parameter of the negative binomial distribution during the optimization; default is exp(10).
- **verbose** logical; print information during the optimization phase; default is FALSE.

arguments passed to control.

Details

. . .

General: The mixed_model() function fits mixed effects models in which the integrals over the random effects in the definition of the marginal log-likelihood cannot be solved analytically and need to be approximated. The function works under the assumption of normally distributed random effects with mean zero and variance-covariance matrix *D*. These integrals are approximated numerically using an adaptive Gauss-Hermite quadrature rule. Using the control argument nAGQ, the user can specify the number of quadrature points used in the approximation.

User-defined family: The user can define its own family object; for an example, see the help page of negative.binomial.

Optimization: A hybrid approach is used, starting with iter_EM iterations and unless convergence was achieved it continuous with a direct optimization of the log-likelihood using function optim and the algorithm specified by optim_method. For stability and speed, the derivative of the log-likelihood with respect to the parameters are internally programmed.

Value

An object of class "MixMod" with components:

coefficients	a numeric vector with the estimated fixed effects.
phis	a numeric vector with the estimated extra parameters.
D	a numeric matrix denoting the estimated covariance matrix of the random effects.
post_modes	a numeric matrix with the empirical Bayes estimates of the random effects.
post_vars	a list of numeric matrices with the posterior variances of the random effects.
logLik	a numeric scalar denoting the log-likelihood value at the end of the optimization procedure.
Hessian	a numeric matrix denoting the Hessian matrix at the end of the optimization procedure.
converged	a logical indicating whether convergence was attained.
data	a copy of the data argument.
id	a copy of the grouping variable from data.
id_name	a character string with the name of the grouping variable.
Terms	a list with two terms components, termsX derived from the fixed argument, and termsZ derived from the random argument.
model_frames	a list with two model.frame components, mfX derived from the fixed argument, and mfZ derived from the random argument.

control	a copy of the (user-specific) control argument.
Funs	a list of functions used in the optimization procedure.
family	a copy of the family argument.
call	the matched call.

Author(s)

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See Also

methods.MixMod, effectPlotData, marginal_coefs

Examples

```
# simulate some data
set.seed(123L)
n <- 200
K <- 15
t.max <- 25
betas <- c(-2.13, -0.25, 0.24, -0.05)
D <- matrix(0, 2, 2)
D[1:2, 1:2] <- c(0.48, -0.08, -0.08, 0.18)
times <- c(replicate(n, c(0, sort(runif(K-1, 0, t.max)))))</pre>
group <- sample(rep(0:1, each = n/2))
DF <- data.frame(year = times, group = factor(rep(group, each = K)))</pre>
X <- model.matrix(~ group * year, data = DF)</pre>
Z <- model.matrix(~ year, data = DF)</pre>
b <- cbind(rnorm(n, sd = sqrt(D[1, 1])), rnorm(n, sd = sqrt(D[2, 2])))</pre>
id <- rep(1:n, each = K)
eta.y <- as.vector(X %*% betas + rowSums(Z * b[id, ]))</pre>
DF$y <- rbinom(n * K, 1, plogis(eta.y))</pre>
DF$id <- factor(id)</pre>
*****
fm1 <- mixed_model(fixed = y \sim year + group, random = \sim 1 \mid id, data = DF,
                   family = binomial())
# fixed effects
fixef(fm1)
# random effects
head(ranef(fm1))
# detailed output
summary(fm1)
```

mixed_model

```
# fitted values for the 'mean subject', i.e., with
# random effects values equal to 0
head(fitted(fm1, type = "mean_subject"))
# fitted values for the conditioning on the estimated random effects
head(fitted(fm1, type = "subject_specific"))
################
fm2 <- mixed_model(fixed = y ~ year, random = ~ 1 | id, data = DF,</pre>
                   family = binomial())
# likelihood ratio test between the two models
anova(fm2, fm1)
# the same hypothesis but with a Wald test
anova(fm1, L = rbind(c(0, 0, 1, 0), c(0, 0, 0, 1)))
################
# An effects plot for the mean subject (i.e., with random effects equal to 0)
nDF <- with(DF, expand.grid(year = seq(min(year), max(year), length.out = 15),</pre>
    group = levels(group)))
plot_data <- effectPlotData(fm2, nDF)</pre>
require("lattice")
xyplot(pred + low + upp ~ year | group, data = plot_data,
       type = "l", lty = c(1, 2, 2), col = c(2, 1, 1), lwd = 2,
       xlab = "Follow-up time", ylab = "log odds")
expit <- function (x) exp(x) / (1 + exp(x))
xyplot(expit(pred) + expit(low) + expit(upp) ~ year | group, data = plot_data,
       type = "1", lty = c(1, 2, 2), col = c(2, 1, 1), lwd = 2,
       xlab = "Follow-up time", ylab = "Probabilities")
# An effects plots for the marginal probabilities
plot_data_m <- effectPlotData(fm2, nDF, marginal = TRUE, cores = 1L)</pre>
expit <- function (x) exp(x) / (1 + exp(x))
xyplot(expit(pred) + expit(low) + expit(upp) ~ year | group, data = plot_data_m,
       type = "l", lty = c(1, 2, 2), col = c(2, 1, 1), lwd = 2,
       xlab = "Follow-up time", ylab = "Probabilities")
# include random slopes
fm1_slp <- update(fm1, random = ~ year | id)</pre>
# increase the number of quadrature points to 15
fm1_slp_q15 <- update(fm1_slp, nAGQ = 15)</pre>
```

a diagonal covariance matrix for the random effects

```
fm1_slp_diag <- update(fm1, random = ~ year || id)
anova(fm1_slp_diag, fm1_slp)</pre>
```

MixMod Methods Various Methods for Standard Generics

Description

Methods for object of class "MixMod" for standard generic functions.

Usage

```
coef(object, ...)
## S3 method for class 'MixMod'
coef(object, sub_model = c("main", "zero_part"),
    ...)
fixef(object, ...)
## S3 method for class 'MixMod'
fixef(object, sub_model = c("main", "zero_part"), ...)
ranef(object, ...)
## S3 method for class 'MixMod'
ranef(object, post_vars = FALSE, ...)
confint(object, parm, level = 0.95, ...)
## S3 method for class 'MixMod'
confint(object,
  parm = c("fixed-effects", "var-cov","extra", "zero_part"),
  level = 0.95, sandwich = FALSE, ...)
anova(object, ...)
## S3 method for class 'MixMod'
anova(object, object2, test = TRUE,
 L = NULL, sandwich = FALSE, ...)
fitted(object, ...)
## S3 method for class 'MixMod'
fitted(object,
```

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```
type = c("mean_subject", "subject_specific", "marginal"),
  link_fun = NULL, ...)
residuals(object, ...)
## S3 method for class 'MixMod'
residuals(object,
  type = c("mean_subject", "subject_specific", "marginal"),
 link_fun = NULL, tasnf_y = function (x) x, ...)
predict(object, ...)
## S3 method for class 'MixMod'
predict(object, newdata, newdata2 = NULL,
    type_pred = c("response", "link"),
    type = c("mean_subject", "subject_specific", "marginal", "zero_part"),
    se.fit = FALSE, M = 300, df = 10, scale = 0.3, level = 0.95,
    seed = 1, return_newdata = FALSE, sandwich = FALSE, ...)
simulate(object, nsim = 1, seed = NULL, ...)
## S3 method for class 'MixMod'
simulate(object, nsim = 1, seed = NULL,
    type = c("subject_specific", "mean_subject"), new_RE = FALSE,
    acount_MLEs_var = FALSE, sim_fun = NULL,
    sandwich = FALSE, ...)
terms(x, ...)
## S3 method for class 'MixMod'
terms(x, type = c("fixed", "random", "zi_fixed", "zi_random"), ...)
formula(x, ...)
## S3 method for class 'MixMod'
formula(x, type = c("fixed", "random", "zi_fixed", "zi_random"), ...)
model.frame(formula, ...)
## S3 method for class 'MixMod'
model.frame(formula, type = c("fixed", "random", "zi_fixed", "zi_random"), ...)
model.matrix(object, ...)
## S3 method for class 'MixMod'
model.matrix(object, type = c("fixed", "random", "zi_fixed", "zi_random"), ...)
```

```
nobs(object, ...)
## S3 method for class 'MixMod'
nobs(object, level = 1, ...)
VIF(object, ...)
## S3 method for class 'MixMod'
VIF(object, type = c("fixed", "zi_fixed"), ...)
cooks.distance(model, ...)
## S3 method for class 'MixMod'
cooks.distance(model, cores = max(parallel::detectCores() - 1, 1), ...)
```

Arguments

object, object2, x, formula, model

	objects inheriting from class "MixMod". When object2 is also provided, then the model behind object must be nested within the model behind object2.
sub_model	character string indicating for which sub-model to extract the estimated coefficients; it is only relevant for zero-inflated models.
post_vars	logical; if TRUE the posterior variances of the random effects are returned as an extra attribute of the numeric matrix produced by $ranef()$.
parm	character string; for which type of parameters to calculate confidence intervals. Option "var-cov" corresponds to the variance-covariance matrix of the random effects. Option extra corresponds to extra (shape/dispersion) parameters in the distribution of the outcome (e.g., the θ parameter in the negative binomial family). Option zero_inflated corresponds to the coefficients of the zero-inflated sub-model.
level	numeric scalar between 0 and 1 denoting the level of the confidence interval. In the nobs() method it denotes the level at which the number of observations is counted. The value 0 corresponds to the number of independent sample units determined by the number of levels of the grouping variable. If set to a value greater than zero, it returns the total number of observations.
test	logical; should a p-value be calculated.
L	a numeric matrix representing a contrasts matrix. This is only used when in anova() only object is provided, and it can only be specified for the fixed effects. When L is used, a Wald test is performed.
sandwich	logical; if TRUE the sandwich estimator is used in the calculation of standard errors.
type	character string indicating the type of fitted values / residuals / predictions / vari- ance inflation factors to calculate. Option "mean_subject" corresponds to only using the fixed-effects part; option "subject_specific" corresponds to using

both the fixed- and random-effects parts; option "marginal" is based in multiplying the fixed effects design matrix with the marginal coefficients obtained by marginal_coefs.

link_fun the link_fun of marginal_coefs.

tasnf_y a function to transform the grouped / repeated measurements outcome before calculating the residuals; for example, relevant in two-part models for semicontinuous data, in which it is assumed that the log outcome follows a normal distribution.

newdata, newdata2

a data frame based on which predictions are to be calculated. newdata2 is only relevant when level = "subject_specific"; see **Details** for more information.

type_pred character string indicating at which scale to calculate predictions. Options are "link" indicating to calculate predictions at the link function / linear predictor scale, and "response" indicating to calculate predictions at the scale of the response variable.

se.fit logical, if TRUE standard errors of predictions are returned.

- M numeric scalar denoting the number of Monte Carlo samples; see **Details** for more information.
- df numeric scalar denoting the degrees of freedom for the Student's t proposal distribution; see **Details** for more information.
- scale numeric scalar or vector denoting the scaling applied to the subject-specific covariance matrices of the random effects; see **Details** for more information.
- seed numerical scalar giving the seed to be used in the Monte Carlo scheme.

return_newdata logical; if TRUE the predict() method returns a copy of the newdata and of newdata2 if the corresponding argument was not NULL, with extra columns the predictions, and the lower and upper limits of the cofidence intervals when type = "subject_specific".

- nsim numeric scalar giving the number of times to simulate the response variable.
- new_RE logical; if TRUE, new random effects will be simulated, and new outcome data will be simulated by simulate() using these new random effect. Otherwise, the empirical Bayes estimates of the random effects from the fitted model will be used.

acount_MLEs_var

logical; if TRUE it accounts for the variability of the maximum likelihood estimates (MLEs) by simulating a new value for the parameters from a multivariate normal distribution with mean the MLEs and covariance matrix the covariance matrix of the MLEs.

sim_fun a function based on which to simulate the response variable. This is relevant for non-standard models. The simulate() function also tries to extract this function from the family component of object. The function should have the following four arguments: n a numeric scalar denoting the number of observations to simulate, mu a numeric vector of means, phis a numeric vector of extra dispersion/scale parameters, and eta_zi a numeric vector for the zero-part of the model, if this is relevant. coresthe number of cores to use in the computation....further arguments; currently none is used.

Details

In generic terms, we assume that the mean of the outcome y_i (i = 1, ..., n denotes the subjects) conditional on the random effects is given by the equation:

$$gE(y_i|b_i) = \eta_i = X_i\beta + Z_ib_i,$$

where g(.) denotes the link function, b_i the vector of random effects, β the vector of fixed effects, and X_i and Z_i the design matrices for the fixed and random effects, respectively.

Argument type_pred of predict() specifies whether predictions will be calculated in the link / linear predictor scale, i.e., η_i or in the response scale, i.e., $gE(y_i|b_i)$.

When type = "mean_subject", predictions are calculated using only the fixed effects, i.e., the $X_i\beta$ part, where X_i is evaluated in newdata. This corresponds to predictions for the 'mean' subjects, i.e., subjects who have random effects value equal to 0. Note, that in the case of nonlinear link functions this does not correspond to the averaged over the population predictions (i.e., marginal predictions).

When type = "marginal", predictions are calculated using only the fixed effects, i.e., the $X_i\beta$ part, where X_i is evaluated in newdata, but with β coefficients the marginalized coefficients obtain from marginal_coefs. These predictions will be marginal / population averaged predictions.

When type = "zero_part", predictions are calculated for the logistic regression of the extra zeropart of the model (i.e., applicable for zero-inflated and hurdle models).

When type = "subject_specific", predictions are calculated using both the fixed- and randomeffects parts, i.e., $X_i\beta + Z_ib_i$, where X_i and Z_i are evaluated in newdata. Estimates for the random effects of each subject are obtained as modes from the posterior distribution $[b_i|y_i;\theta]$ evaluated in newdata and with *theta* (denoting the parameters of the model, fixed effects and variance components) replaced by their maximum likelihood estimates.

Notes: (i) When se.fit = TRUE and type_pred = "response", the standard errors returned are on the linear predictor scale, not the response scale. (ii) When se.fit = TRUE and the model contains an extra zero-part, no standard errors are computed when type = "mean_subject". (iii) When the model contains an extra zero-part, type = "marginal" predictions are not yet implemented.

When se.fit = TRUE and type = "subject_specific", standard errors and confidence intervals for the subject-specific predictions are obtained by a Monte Carlo scheme entailing three steps repeated M times, namely

- **Step I** Account for the variability of maximum likelihood estimates (MLES) by simulating a new value θ^* for the parameters θ from a multivariate normal distribution with mean the MLEs and covariance matrix the covariance matrix of the MLEs.
- **Step II** Account for the variability in the random effects estimates by simulating a new value b_i^* for the random effects b_i from the posterior distribution $[b_i|y_i; \theta^*]$. Because the posterior distribution does not have a closed-form, a Metropolis-Hastings algorithm is used to sample the new value b_i^* using as proposal distribution a multivariate Student's-t distribution with degrees of freedom df, centered at the mode of the posterior distribution $[b_i|y_i; \theta]$ with θ the MLEs, and scale matrix the inverse Hessian matrix of the log density of $[b_i|y_i; \theta]$ evaluated at the modes, but multiplied by scale. The scale and df parameters can be used to adjust the acceptance rate.

Step III The predictions are calculated using $X_i\beta^* + Z_ib_i^*$.

Argument newdata2 can be used to calculate dynamic subject-specific predictions. I.e., using the observed responses y_i in newdata, estimates of the random effects of each subject are obtained. For the same subjects we want to obtain predictions in new covariates settings for which no response data are yet available. For example, in a longitudinal study, for a subject we have responses up to a follow-up t (newdata) and we want the prediction at $t + \Delta t$ (newdata2).

Value

The estimated fixed and random effects, coefficients (this is similar as in package **nlme**), confidence intervals fitted values (on the scale on the response) and residuals.

Author(s)

Dimitris Rizopoulos <d.rizopoulos@erasmusmc.nl>

See Also

mixed_model, marginal_coefs

Examples

```
# simulate some data
set.seed(123L)
n <- 500
K <- 15
t.max <- 25
betas <- c(-2.13, -0.25, 0.24, -0.05)
D <- matrix(0, 2, 2)
D[1:2, 1:2] <- c(0.48, -0.08, -0.08, 0.18)
times <- c(replicate(n, c(0, sort(runif(K-1, 0, t.max)))))</pre>
group <- sample(rep(0:1, each = n/2))
DF <- data.frame(year = times, group = factor(rep(group, each = K)))</pre>
X <- model.matrix(~ group * year, data = DF)</pre>
Z <- model.matrix(~ year, data = DF)</pre>
b <- cbind(rnorm(n, sd = sqrt(D[1, 1])), rnorm(n, sd = sqrt(D[2, 2])))</pre>
id <- rep(1:n, each = K)
eta.y <- as.vector(X %*% betas + rowSums(Z * b[id, ]))</pre>
DF$y <- rbinom(n * K, 1, plogis(eta.y))</pre>
DF$id <- factor(id)</pre>
****
fm1 <- mixed_model(fixed = y \sim year + group, random = \sim year | id, data = DF,
                   family = binomial())
```

head(coef(fm1))
fixef(fm1)

negative.binomial Family function for Negative Binomial Mixed Models

Description

Specifies the information required to fit a Negative Binomial generalized linear mixed model, using mixed_model().

Usage

negative.binomial()

Note

Currently only the log-link is implemented.

Author(s)

Dimitris Rizopoulos <d.rizopoulos@erasmusmc.nl>

Examples

```
summary(gm1)
```

```
# We do a likelihood ratio test with the Poisson mixed model
gm0 <-mixed_model(fixed = y ~ f1 * f2, random = ~ 1 | g, data = dd,
                     family = poisson())
anova(gm0, gm1)
# Define a cutom-made family function to be used with mixed_model()
# the required components are 'family', 'link', 'linkfun', 'linkinv' and 'log_dens';
# the extra functions 'score_eta_fun' and 'score_phis_fun' can be skipped and will
# internally approximated using numeric derivatives (though it is better that you provide
# them).
my_negBinom <- function (link = "log") {</pre>
    stats <- make.link(link)</pre>
    log_dens <- function (y, eta, mu_fun, phis, eta_zi) {</pre>
        # the log density function
        phis <- exp(phis)</pre>
        mu <- mu_fun(eta)</pre>
        log_mu_phis <- log(mu + phis)</pre>
        comp1 <- lgamma(y + phis) - lgamma(phis) - lgamma(y + 1)</pre>
        comp2 <- phis * log(phis) - phis * log_mu_phis</pre>
        comp3 <- y * log(mu) - y * log_mu_phis</pre>
        out <- comp1 + comp2 + comp3</pre>
        attr(out, "mu_y") <- mu</pre>
        out
    }
    score_eta_fun <- function (y, mu, phis, eta_zi) {</pre>
        # the derivative of the log density w.r.t. mu
        phis <- exp(phis)</pre>
        mu_phis <- mu + phis</pre>
        comp2 <- - phis / mu_phis</pre>
        comp3 <- y / mu - y / mu_phis</pre>
        # the derivative of mu w.r.t. eta (this depends on the chosen link function)
        mu.eta <- mu
        (comp2 + comp3) * mu.eta
    }
    score_phis_fun <- function (y, mu, phis, eta_zi) {</pre>
        # the derivative of the log density w.r.t. phis
        phis <- exp(phis)</pre>
        mu_phis <- mu + phis</pre>
        comp1 <- digamma(y + phis) - digamma(phis)</pre>
        comp2 <- log(phis) + 1 - log(mu_phis) - phis / mu_phis</pre>
        comp3 <- - y / mu_phis</pre>
        comp1 + comp2 + comp3
    }
    structure(list(family = "user Neg Binom", link = stats$name, linkfun = stats$linkfun,
                    linkinv = stats$linkinv, log_dens = log_dens,
                    score_eta_fun = score_eta_fun,
                    score_phis_fun = score_phis_fun),
               class = "family")
}
```

summary(fm)

scoring_rules Pa

Proper Scoring Rules for Categorical Data

Description

Calculates the logarithmic, quadratic/Brier and spherical based on a fitted mixed model for categorical data.

Usage

```
scoring_rules(object, newdata, newdata2 = NULL, max_count = 2000,
    return_newdata = FALSE)
```

Arguments

object	an object inheriting from class "MixMod".
newdata	a data.frame based on which to estimate the random effect and calculate predic- tions. It should contain the response variable.
newdata2	a data.frame based on which to estimate the random effect and calculate predic- tions. It should contain the response variable.
max_count	numeric scalar denoting the maximum count up to which to calculate probabili- ties; this is relevant for count response data.
return_newdata	logical; if TRUE the values of the scoring rules are ruturned as extra columns of the newdata or newdata2 data.frame.

Value

A data.frame with (extra) columns the values of the logarithmic, quadratic and spherical scoring rules calculated based on the fitted model and the observed responses in newdata or newdata2.

Author(s)

Dimitris Rizopoulos <d.rizopoulos@erasmusmc.nl>

References

Carvalho, A. (2016). An overview of applications of proper scoring rules. *Decision Analysis* 13, 223–242. doi:10.1287/deca.2016.0337

scoring_rules

See Also

mixed_model, predict.MixMod

Examples

NA NA

NA

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