Package ‘glmmTMB’

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Title Generalized Linear Mixed Models using Template Model Builder
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Description Fit linear and generalized linear mixed models with various extensions, including zero-inflation. The models are fitted using maximum likelihood estimation via 'TMB' (Template Model Builder). Random effects are assumed to be Gaussian on the scale of the linear predictor and are integrated out using the Laplace approximation. Gradients are calculated using automatic differentiation.
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Description

Methods have been written that allow glmmTMB objects to be used with several downstream packages that enable different forms of inference. In particular,

- `car::Anova` constructs type-II and type-III Anova tables for the fixed effect parameters of the conditional model (this might work with the fixed effects of the zero-inflation or dispersion models, but has not been tested)
- the `effects` package computes graphical tabular effect displays (again, for the fixed effects of the conditional component)
- the `emmeans` package computes estimated marginal means (aka least-squares means) for the fixed effects of the conditional component

Usage

```r
Anova.glmmTMB(mod, type = c("II", "III", 2, 3),
    test.statistic = c("Chisq", "F"), component = "cond",
    vcov = vcov(mod)[[component]], singular.ok, ...)
```

```r
Effect.glmmTMB(focal.predictors, mod, ...)
recover_data.glmmTMB(object, ...)
emm_basis.glmmTMB(object, trms, xlev, grid, component = "cond", ...)
```

Arguments

- `mod` a glmmTMB model
- `type` type of test, "II", "III", 2, or 3. Roman numerals are equivalent to the corresponding Arabic numerals. See `Anova` for details.
- `test.statistic` unused: only valid choice is "Chisq" (i.e., Wald chi-squared test)
- `component` which component of the model to compute emmeans for (conditional ("cond"), zero-inflation ("zi"), or dispersion ("disp"))
- `vcov.` variance-covariance matrix (usually extracted automatically)
- `singular.ok` OK to do ANOVA with singular models (unused)?
- `...` Additional parameters that may be supported by the method.
- `focal.predictors` a character vector of one or more predictors in the model in any order.
- `object` a glmmTMB model
- `trms` The terms component of object (typically with the response deleted, e.g. via `delete.response`
**confint.glmmTMB**

- **xlev**
  - Named list of factor levels (*excluding* ones coerced to factors in the model formula)

- **grid**
  - A `data.frame` (provided by `ref_grid`) containing the predictor settings needed in the reference grid

### Examples

```r
warp.lm <- glmmTMB(breaks ~ wool * tension, data = warpbreaks)
if (require(emmeans)) {
  emmeans (warp.lm, poly ~ tension | wool)
}
if (require(car)) {
  Anova(warp.lm, type="III")
}
if (require/effects()) {
  plot(allEffects(warp.lm))
}
```

### Description

Calculate confidence intervals

### Usage

```r
confint(object, parm = NULL, level = 0.95,
  method = c("wald", "Wald", "profile", "uniroot"),
  component = c("all", "cond", "zi", "other"), estimate = TRUE,
  parallel = c("no", "multicore", "snow"),
  ncpus = getOption("profile.ncpus", 1L), cl = NULL, ...)
```

### Arguments

- **object**
  - `glmmTMB` fitted object.
- **parm**
  - Specification of a parameter subset *after* component subset has been applied.
- **level**
  - Confidence level.
- **method**
  - One of: `"wald"`, `"Wald"`, `"profile"`, or `"uniroot"`: see `Details` function.
- **component**
  - Which of the three components ‘cond’, ‘zi’ or ‘other’ to select. Default is to select ‘all’.
- **estimate**
  - (logical) add a third column with estimate?
- **parallel**
  - Method (if any) for parallel computation
- **ncpus**
  - Number of CPUs/cores to use for parallel computation
- **cl**
  - Cluster to use for parallel computation
- **...**
  - Arguments may be passed to `profile.merMod` or `tmbroot`
Details

Available methods are

- **wald**: These intervals are based on the standard errors calculated for parameters on the scale of their internal parameterization depending on the family. Derived quantities such as standard deviation parameters and dispersion parameters are backtransformed. It follows that confidence intervals for these derived quantities are asymmetric.

- **profile**: This method computes a likelihood profile for the specified parameter(s) using `profile.glmmTMB`; fits a spline function to each half of the profile; and inverts the function to find the specified confidence interval.

- **uniroot**: This method uses the `uniroot` function to find critical values of one-dimensional profile functions for each specified parameter.

Examples

data(sleepstudy, package="lme4")
model <- glmmTMB(Reaction ~ Days + (1|Subject), sleepstudy)
model2 <- glmmTMB(Reaction ~ Days + (1|Subject), sleepstudy,
dispformula = ~I(Days>8))
confint(model)  ## Wald/delta-method CIs
confint(model,parm="theta_")  ## Wald/delta-method CIs
confint(model,parm=1,method="profile")

epil2

Seizure Counts for Epileptics - Extended

Description

Extended version of the `epil` dataset of the `MASS` package. The three transformed variables `visit`, `base`, and `age` used by Booth et al. (2003) have been added to `epil`.

Usage

epil2

Format

A data frame with 236 observations on the following 12 variables:

- y  an integer vector.
- trt a factor with levels "placebo" and "progabide".
- base  an integer vector.
- age  an integer vector.
- V4  an integer vector.
- subject  an integer vector.
- period  an integer vector.
findReTrmClasses

Description

list of specials – taken from enum.R

Usage

findReTrmClasses()
**fixef**

*Extract fixed-effects estimates*

**Description**

Extract the fixed-effects estimates

**Usage**

```r
default method for class 'glmmTMB'
fixef(object, ...)
```

**Arguments**

- `object`: any fitted model object from which fixed effects estimates can be extracted.
- `...`: optional additional arguments. Currently none are used in any methods.

**Details**

Extract the estimates of the fixed-effects parameters from a fitted model.

**Value**

a named, numeric vector of fixed-effects estimates.

**Examples**

```r
data(sleepstudy, package = "lme4")
fixef(glmmTMB(Reaction ~ Days + (1|Subject) + (0+Days|Subject), sleepstudy))
```
Arguments

varcor: a VarCorr (-like) matrix with attributes.
digits: the number of significant digits.
comp: character vector of length one or two indicating which columns out of "Variance" and "Std.Dev." should be shown in the formatted output.
formatter: the function to be used for formatting the standard deviations and or variances (but not the correlations which (currently) are always formatted as "0.nnn")
useScale: whether to report a scale parameter (e.g. residual standard deviation)
... optional arguments for formatter(*) in addition to the first (numeric vector) and digits.

Value

a character matrix of formatted VarCorr entries from varc.

formula.glmmTMB  

Extract the formula of a glmmTMB object

Description

Extract the formula of a glmmTMB object

Usage

## S3 method for class 'glmmTMB'
formula(x, fixed.only = FALSE, component = c("cond", "zi", "disp"), ...)

Arguments

x: a glmmTMB object
fixed.only: (logical) drop random effects, returning only the fixed-effect component of the formula?
component: formula for which component of the model to return (conditional, zero-inflation, or dispersion)
... unused, for generic consistency
getCapabilities

List model options that glmmTMB knows about

Description

List model options that glmmTMB knows about

Usage

getCapabilities(what = "all", check = FALSE)

Arguments

what (character) which type of model structure to report on ("all","family","link","covstruct")
check (logical) do brute-force checking to test whether families are really implemented (only available for what="family")

Value

if check==FALSE, returns a vector of the names (or a list of name vectors) of allowable entries; if check==TRUE, returns a logical vector of working families

Note

these are all the options that are defined internally; they have not necessarily all been implemented (FIXME!)

getME.glmmTMB

Extract or Get Generalize Components from a Fitted Mixed Effects Model

Description

Extract or Get Generalize Components from a Fitted Mixed Effects Model

Usage

## S3 method for class 'glmmTMB'
getME(object, name = c("X", "Xzi", "Z", "Zzi", "Xd", "theta", "beta"), ...)

Arguments

object a fitted glmmTMB object
name of the component to be retrieved
... ignored, for method compatibility
getReStruc

See Also
getME

getReStruc

Calculate random effect structure Calculates number of random effects, number of parameters, blocksize and number of blocks. Mostly for internal use.

Description
Calculate random effect structure Calculates number of random effects, number of parameters, blocksize and number of blocks. Mostly for internal use.

Usage
getReStruc(reTrms, ss = NULL)

Arguments
reTrms random-effects terms list
ss a character string indicating a valid covariance structure. Must be one of names(glmmTMB::valid_covstruct); default is to use an unstructured variance-covariance matrix ("us") for all blocks.

Value
a list

blockNumTheta number of variance covariance parameters per term
blockSize size (dimension) of one block
blockReps number of times the blocks are repeated (levels)
covCode structure code

Examples
data(sleepstudy, package="lme4")
rt <- lme4::lFormula(Reaction~Days+(1|Subject)+(0+Days|Subject),
sleepstudy)$reTrms
rt2 <- lme4::lFormula(Reaction~Days+(Days|Subject),
sleepstudy)$reTrms
getReStruc(rt)
**getXReTrms**

Create X and random effect terms from formula

**Description**

Create X and random effect terms from formula

**Usage**

`getXReTrms(formula, mf, fr, ranOK = TRUE, type = "", contrasts)`

**Arguments**

- `formula`: current formula, containing both fixed & random effects
- `mf`: matched call
- `fr`: full model frame
- `ranOK`: random effects allowed here?
- `type`: label for model type
- `contrasts`: a list of contrasts (see `?glmmTMB`)

**Value**

a list composed of

- `X`: design matrix for fixed effects
- `Z`: design matrix for random effects
- `reTrms`: output from `mkReTrms` from `lme4`
- `offset`: offset vector, or vector of zeros if offset not specified

**get_cor**

Translate vector of correlation parameters to correlation values, following the definition at [http://kaskr.github.io/adcomp/classUNSTRUCTURED_CORR_t.html](http://kaskr.github.io/adcomp/classUNSTRUCTURED_CORR_t.html): if `L` is the lower-triangular matrix with 1 on the diagonal and the correlation parameters in the lower triangle, then the correlation matrix is defined as \( \Sigma = D^{-1/2}LL^\top D^{-1/2} \), where \( D = \text{diag}(LL^\top) \). For a single correlation parameter \( \theta_0 \), this works out to \( \rho = \theta_0 / \sqrt{1 + \theta_0^2} \).

**Description**

Translate vector of correlation parameters to correlation values, following the definition at [http://kaskr.github.io/adcomp/classUNSTRUCTURED_CORR_t.html](http://kaskr.github.io/adcomp/classUNSTRUCTURED_CORR_t.html): if `L` is the lower-triangular matrix with 1 on the diagonal and the correlation parameters in the lower triangle, then the correlation matrix is defined as \( \Sigma = D^{-1/2}LL^\top D^{-1/2} \), where \( D = \text{diag}(LL^\top) \). For a single correlation parameter \( \theta_0 \), this works out to \( \rho = \theta_0 / \sqrt{1 + \theta_0^2} \).
Usage

glmmTMB

Description

Fit models with TMB

Usage

glmmTMB(formula, data = NULL, family = gaussian(), ziformula = ~0,
dispformula = ~1, weights = NULL, offset = NULL,
contrasts = NULL, na.action = na.fail, se = TRUE,
verbose = FALSE, doFit = TRUE, control = glmmTMBControl(),
REML = FALSE)

Arguments

formula combined fixed and random effects formula, following lme4 syntax
data data frame
family a family function, a character string naming a family function, or the result of a
call to a family function family (variance/link function) information; see family
for generic discussion of families or family_glmmTMB for details of glmmTMB-
specific families.
ziformula a one-sided (i.e., no response variable) formula for zero-inflation combining
fixed and random effects: the default ~0 specifies no zero-inflation. Specifying
~. sets the zero-inflation formula identical to the right-hand side of formula
(i.e., the conditional effects formula); terms can also be added or subtracted.
When using ~. as the zero-inflation formula in models where the conditional
effects formula contains an offset term, the offset term will automatically be dropped. The zero-inflation model uses a logit link.
dispformula  a one-sided formula for dispersion containing only fixed effects: the default ~1
specifies the standard dispersion given any family. The argument is ignored for families that do not have a dispersion parameter. For an explanation of the dispersion parameter for each family, see (sigma). The dispersion model uses a log link. In Gaussian mixed models, dispformula=~0 fixes the parameter to be 0, forcing variance into the random effects.

weights  weights, as in glm. Not automatically scaled to have sum 1.
offset  offset for conditional model (only)
contrasts  an optional list, e.g. list(fac1="contr.sum"). See the contrasts.arg of model.matrix.default.
n.action  how to handle missing values (see na.action and model.frame); from lm, “The
default is set by the na.action setting of options, and is na.fail if that is unset. The ‘factory-fresh’ default is na.omit.”
se  whether to return standard errors
verbose  logical indicating if some progress indication should be printed to the console.
dofit  whether to fit the full model, or (if FALSE) return the preprocessed data and parameter objects, without fitting the model
control  control parameters; see glmmTMBControl.
REML  Logical; Use REML estimation rather than maximum likelihood.

Details

• binomial models with more than one trial (i.e., not binary/Bernoulli) can either be specified in the form prob ~ ..., weights = N or in the more typical two-column matrix (cbind(successes,failures)~...) form.
• Behavior of REML=TRUE for Gaussian responses matches lme4::lmer. It may also be useful in some cases with non-Gaussian responses (Millar 2011). Simulations should be done first to verify.
• Because the df.residual method for glmmTMB currently counts the dispersion parameter, one would need to multiply by sqrt(nobs(fit)/(1+df.residual(fit))) when comparing with lm ...
• by default, vector-valued random effects are fitted with unstructured (general positive definite) variance-covariance matrices. Structured variance-covariance matrices can be specified in the form struc(terms|group), where struc is one of
  – diag (diagonal, heterogeneous variance)
  – ar1 (autoregressive order-1, homogeneous variance)
  – cs (compound symmetric, heterogeneous variance)
  – ou (* Ornstein-Uhlenbeck, homogeneous variance)
  – exp (* exponential autocorrelation)
  – gau (* Gaussian autocorrelation)
  – mat (* Matérn process correlation)
  – toep (* Toeplitz)

(note structures marked with * are experimental/untested)
• For backward compatibility, the family argument can also be specified as a list comprising the name of the distribution and the link function (e.g. `list(family="binomial", link="logit")`). However, **this alternative is now deprecated** (it produces a warning and will be removed at some point in the future). Furthermore, certain capabilities such as Pearson residuals or predictions on the data scale will only be possible if components such as variance and linkfun are present (see `family`).

**References**


**Examples**

```r
(m1 <- glmmTMB(count~ mined + (1|site),
    zi=mined,
    family=poisson, data=Salamanders))
summary(m1)

## Zero-inflated negative binomial model
(m2 <- glmmTMB(count=spp + mined + (1|site),
    zi=spp + mined,
    family=nbinom2, Salamanders))

## Hurdle Poisson model
(m3 <- glmmTMB(count=spp + mined + (1|site),
    zi=spp + mined,
    family=truncated_poisson, Salamanders))

## Binomial model
data(cbpp, package="lme4")
(tmbm1 <- glmmTMB(cbind(incidence, size-incidence) ~ period + (1 | herd),
    family=binomial, data=cbpp))

## Dispersion model
sim1=function(nfac=40, nt=100, facsd=.1, tsd=.15, mu=0, residsd=1)
{
    dat=expand.grid(fac=factor(letters[1:nfac]), t= 1:nt)
    n=nrow(dat)
    dat$REfac=rnorm(nfac, sd= facsd)[dat$fac]
    dat$RET=rnorm(nt, sd= tsd)[dat$t]
    dat$x=rnorm(n, mean=mu, sd=residsd) + dat$REfac + dat$RET
    return(dat)
}
set.seed(101)
d1 = sim1(mu=100, residsd =10)
d2 = sim1(mu=200, residsd =5)
d1$sd="ten"
d2$sd="five"
set = rbind(d1, d2)
m0 = glmmTMB(x~sd+(1|t), dispformula=~sd, dat)
fixef(m0)$disp
```
glmmTMBControl

Usage

```r
glmmtmbcontrol(optCtrl = list(iter.max = 300, eval.max = 400),
profile = FALSE, collect = FALSE)
```

Arguments

- `optCtrl`: Passed as argument control to `nlminb`.
- `profile`: Logical; Experimental option to improve speed and robustness when a model has many fixed effects.
- `collect`: Logical; Experimental option to improve speed by recognizing duplicated observations.

Details

The general non-linear optimizer `nlminb` is used by `glmmTMB` for parameter estimation. It may sometimes be necessary to tweak some tolerances in order to make a model converge. For instance, the warning ‘iteration limit reached without convergence’ may be fixed by increasing the number of iterations using something like:

```r
glmmtmbcontrol(optCtrl=list(iter.max=1e3,eval.max=1e3)).
```

The argument `profile` allows `glmmTMB` to use some special properties of the optimization problem in order to speed up estimation in cases with many fixed effects. Enable this option using:

```r
glmmtmbcontrol(profile=TRUE).
```

Control parameters may depend on the model specification, because each control component is evaluated inside `TMBStruc`, the output of `mktMBStruc`. To specify that `profile` should be enabled for more than 5 fixed effects one can use:

```r
glmmtmbcontrol(profile=quote(length(parameters$beta)>=5)).
```
Description

see `refit` and `isLMM` for details

Usage

```r
## S3 method for class 'glmmTMB'
isLMM(object)

## S3 method for class 'glmmTMB'
refit(object, newresp, ...)
```

Arguments

- `object`: a fitted glmmTMB object
- `newresp`: a new response vector
- `...`: additional arguments (for generic consistency; ignored)

Details

These methods are still somewhat experimental (check your results carefully!), but they should allow parametric bootstrapping. They work by copying and replacing the original response column in the data frame passed to `glmmtmb`, so they will only work properly if (1) the data frame is still available in the environment and (2) the response variable is specified as a single symbol (e.g. proportion or a two-column matrix constructed on the fly with `cbind()`). Untested with binomial models where the response is specified as a factor.

Examples

```r
if (requireNamespace("lme4")) {
  ## Not run:
  fm1 <- glmmTMB(count~mined+(1|spp),
                 ziformula=~mined,
                 data=Salamanders,
                 family=nbinom1)
  b1 <- lme4::bootMer(fm1, FUN=function(x) fixef(x)$zi, nsim=20, .progress="txt")
  if (requireNamespace("boot")) {
    boot.ci(b1, type="perc")
  }

  ## End(Not run)
}
```
Description
Family functions for glmmTMB

Usage
nbinom2(link = "log")
nbinom1(link = "log")
compois(link = "log")
truncated_compois(link = "log")
genpois(link = "log")
truncated_genpois(link = "log")
truncated_poisson(link = "log")
truncated_nbinom2(link = "log")
truncated_nbinom1(link = "log")
beta_family(link = "logit")
betabinomial(link = "logit")
tweedie(link = "log")

Arguments

link (character) link function for the conditional mean ("log", "logit", "probit", "inverse", "cloglog", or "identity")

Details
If specified, the dispersion model uses a log link. Denoting the dispersion parameter as \(\phi = \exp(\eta)\) (where \(\eta\) is the linear predictor from the dispersion model) and the predicted mean as \(\mu\):

**gaussian** (from base R): constant variance=\(\phi\)

**Gamma** (from base R) \(\phi\) is the shape parameter, i.e variance=\(\mu\)*\(\phi\)

**nbinom2** variance increases quadratically with the mean (Hardin & Hilbe 2007), i.e. variance=\(\mu\)*(1+\(\mu/\phi\))

**nbinom1** variance increases linearly with the mean (Hardin & Hilbe 2007), i.e. variance=\(\mu\)*(1+\(\phi\))
compois is the Conway-Maxwell Poisson parameterized with the exact mean which differs from the COMPoissonReg package (Sellers & Lotze 2015)

genpois is the generalized Poisson distribution

beta follows the parameterization of Ferrari and Cribari-Neto (2004) and the betareg package, i.e. variance=mu*(1-mu)

Value
returns a list with (at least) components

family length-1 character vector giving the family name
link length-1 character vector specifying the link function
variance a function of either 1 (mean) or 2 (mean and dispersion parameter) arguments giving a value proportional to the predicted variance (scaled by sigma(.))

References
Details

Some \texttt{glmmTMB} covariance structures require extra information, such as temporal or spatial coordinates. \texttt{numFactor} allows to associate such extra information as part of a factor via the factor levels. The original numeric coordinates are recoverable without loss of precision using the function \texttt{parsenumLevels}. Factor levels are sorted coordinate wise from left to right: first coordinate is fastest running.

Value

Factor with specialized coding of levels.

Examples

```r
## 1D example
numFactor(sample(1:5, 20, TRUE))
## 2D example
coords <- cbind(sample(1:5, 20, TRUE), sample(1:5, 20, TRUE))
(f <- numFactor(coords))
parsenumLevels(levels(f)) ## Sorted
## Used as part of a model.matrix
model.matrix(~ f)
## parsenumLevels( colnames(model.matrix(~ f)) )
## Error: 'Failed to parse numeric levels: (Intercept)'
parsenumLevels( colnames(model.matrix(~ f - 1)) )
```

---

**Owls**

*Begging by Owl Nestlings*

**Description**

Begging by owl nestlings

**Usage**

data(Owls)

**Format**

The \texttt{Owls} data set is a data frame with 599 observations on the following variables:

- **Nest** a factor describing individual nest locations
- **FoodTreatment** (factor) food treatment: Deprived or Satiated
- **SexParent** (factor) sex of provisioning parent: Female or Male
- **ArrivalTime** a numeric vector
- **SiblingNegotiation** a numeric vector
- **BroodSize** brood size
- **NegPerChick** number of negotiations per chick
Note

Access to data kindly provided by Alain Zuur

Source


References


Examples

data(Owls, package = "glmmTMB")
require("lattice")
bwplot(reorder(Nest,NegPerChick) ~ NegPerChick | FoodTreatment:SexParent,
data=Owls)
dotplot(reorder(Nest,NegPerChick) ~ NegPerChick| FoodTreatment:SexParent,
data=Owls)

## Not run:
## Fit negative binomial model with "constant" Zero Inflation:
owls_nb1 <- glmmTMB(SiblingNegotiation ~ FoodTreatment*SexParent +
          (1|Nest)*offset(log(BroodSize)),
          family = nbinom1(), zi = ~1, data=Owls)
owls_nb1_bs <- update(owls_nb1,.
          ~ . - offset(log(BroodSize)) + log(BroodSize))
fixef(owls_nb1_bs)

## End(Not run)

---

predict.glmmTMB prediction

Description

prediction

Usage

## S3 method for class 'glmmTMB'
predict(object, newdata = NULL, se.fit = FALSE, re.form, allow.new.levels = FALSE, type = c("link", "response", "conditional", "zprob", "zlink"), ztype = NULL, na.action = na.pass, debug = FALSE, ...)
predict.glmmTMB

Arguments

- **object**: a `glmmTMB` object
- **newdata**: new data for prediction
- **se.fit**: return the standard errors of the predicted values?
- **re.form**: (not yet implemented) specify which random effects to condition on when predicting. To compute population-level predictions for a given grouping variable (i.e., setting _all_ random effects for that grouping variable to zero), set the group value to `NA`.
- **allow.new.levels**: allow previously unobserved levels in random-effects variables? see details.
- **type**: Denoting \( \mu \) as the mean of the conditional distribution and \( p \) as the zero-inflation probability, the possible choices are:
  - "link": conditional mean on the scale of the link function, or equivalently the linear predictor of the conditional model
  - "response": expected value; this is \( \mu \times (1 - p) \) for zero-inflated models and \( \mu \) otherwise
  - "conditional": mean of the conditional response; \( \mu \) for all models (i.e., synonymous with "response" in the absence of zero-inflation
  - "zprob": the probability of a structural zero (gives an error for non-zero-inflated models)
  - "zlink": predicted zero-inflation probability on the scale of the logit link function
- **ztype**: deprecated: formerly used to specify type of zero-inflation probability. Now synonymous with type
- **na.action**: how to handle missing values in `newdata` (see `na.action`); the default (`na.pass`) is to predict `NA`
- **debug**: (logical) return the `TMBS` object that will be used internally for debugging?
- **...**: unused - for method compatibility

Details

- Prediction of new random effect levels is possible as long as the model specification (fixed effects and parameters) is kept constant. However, to ensure intentional usage, a warning is triggered if `allow.new.levels=FALSE` (the default).
- Prediction using "data-dependent bases" (variables whose scaling or transformation depends on the original data, e.g. `poly`, `ns`, or `poly`) should work properly; however, users are advised to check results extra-carefully when using such variables. Models with different versions of the same data-dependent basis type in different components (e.g. `formula= y ~ poly(x,3)`, `dispformula= ~poly(x)` will probably not produce correct predictions.

Examples

```r
data(sleepstudy, package="lme4")
g0 <- glmmTMB(Reaction~Days+(Days|Subject), sleepstudy)
predict(g0, sleepstudy)
```
## profile.glmmTMB

Compute likelihood profiles for a fitted model

### Description

Compute likelihood profiles for a fitted model

### Usage

```r
## S3 method for class 'profile.glmmTMB'
confint(object, parm = NULL, level = 0.95, ...
```

### Arguments

- `object`: An object of class `profile.glmmTMB`
- `parm`: A character vector of parameters to be extracted.
- `level`: The level of confidence for the intervals.
- `...`: Further parameters passed to the `confint` method.

## print.VarCorr.glmmTMB

Printing The Variance and Correlation Parameters of a glmmTMB

### Description

Printing The Variance and Correlation Parameters of a glmmTMB

### Usage

```r
## S3 method for class 'VarCorr.glmmTMB'
print(x, digits = max(3,getOption("digits") -
  2), comp = "Std.Dev.", formatter = format, ...)
```

### Arguments

- `x`: A result of `VarCorr(<glmmTMB>)`
- `digits`: Number of significant digits to use.
- `comp`: A string specifying the component to format and print.
- `formatter`: A function.
- `...`: Optional further arguments, passed the next `print` method.

## Examples

```r
# Predict new Subject
nd <- sleepstudy[1,]
nd$Subject <- "new"
predict(g0, newdata=nd, allow.new.levels=TRUE)
# population-level prediction
nd_pop <- data.frame(Days=unique(sleepstudy$Days),
Subject=NA)
predict(g0, newdata=nd_pop)
```
profile.glmmTMB

Arguments

- **fitted**: a fitted glmmTMB object
- **parm**: which parameters to profile, specified
  - by index (position)
  - by name (matching the row/column names of vcov(object, full=TRUE))
  - as "theta_" (random-effects variance-covariance parameters) or "beta_" (conditional and zero-inflation parameters)
- **level_max**: maximum confidence interval target for profile
- **npts**: target number of points in (each half of) the profile (approximate)
- **stepfac**: initial step factor (fraction of estimated standard deviation)
- **stderr**: standard errors to use as a scaling factor when picking step sizes to compute the profile; by default (if stderr is NULL, or NA for a particular element), uses the estimated (Wald) standard errors of the parameters
- **trace**: print tracing information? If trace=FALSE or 0, no tracing; if trace=1, print names of parameters currently being profiled; if trace>1, turn on tracing for the underlying `tmbprofile` function
- **parallel**: method (if any) for parallel computation
- **ncpus**: number of CPUs/cores to use for parallel computation
- **cl**: cluster to use for parallel computation
- **...**: additional arguments passed to `tmbprofile`
- **object**: a fitted profile (`profile.glmmTMB`) object
- **level**: confidence level

Details

Fits natural splines separately to the points from each half of the profile for each specified parameter (i.e., values above and below the MLE), then finds the inverse functions to estimate the endpoints of the confidence interval

Value

An object of class `profile.glmmTMB`, which is also a data frame, with columns .par (parameter being profiled), .focal (value of focal parameter), value (negative log-likelihood).

Examples

```r
## Not run:
m1 <- glmmTMB(count ~ mined + (1|site),
         zi=~mined, family=poisson, data=Salamanders)
salamander_prof1 <- profile(m1, parallel="multicore",
              ncpus=2, trace=1)
## testing
salamander_prof1 <- profile(m1, trace=1, parm=1)
salamander_prof1M <- profile(m1, trace=1, parm=1, npts = 4)
salamander_prof2 <- profile(m1, parm="theta_")
```
## ranef.glmmTMB

### Extract Random Effects

**Description**

Generic function to extract random effects from glmmTMB models, both for the conditional model and zero inflation.

**Usage**

```r
## S3 method for class 'glmmTMB'
ranef(object, condVar = TRUE, ...)
```

```r
## S3 method for class 'ranef.glmmTMB'
as.data.frame(x, ..., 
    stringsAsFactors = default.stringsAsFactors())
```

```r
## S3 method for class 'glmmTMB'
coef(object, condVar = FALSE, ...)
```

**Arguments**

- `object`: a glmmTMB model.
- `condVar`: include conditional variances in result?
- `...`: some methods for this generic function require additional arguments.
- `x`: a ranef.glmmTMB object (i.e., the result of running ranef on a fitted glmmTMB model).
- `stringsAsFactors`: see `data.frame`
ranef.glmmTMB

Value

- For `ranef`, an object of class `ranef.glmmTMB` with two components:
  - `cond` a list of data frames, containing random effects for the conditional model.
  - `zi` a list of data frames, containing random effects for the zero inflation.

If `condVar=TRUE`, the individual list elements within the `cond` and `zi` components (corresponding to individual random effects terms) will have associated `condVar` attributes giving the conditional variances of the random effects values. These are in the form of three-dimensional arrays: see `ranef.merMod` for details (the only difference between the packages is that the attributes are called `"postVar"` in `lme4`, vs. `"condVar"` in `glmmTMB`.

- For `coef.glmmTMB`: a similar list, but containing the overall coefficient value for each level (i.e., the sum of the fixed effect estimate and the random effect value for that level). Conditional variances are not yet available as an option for `coef.glmmTMB`.

- For `as.data.frame`: a data frame with components
  - `component` part of the model to which the random effects apply (conditional or zero-inflation)
  - `grpvar` grouping variable
  - `term` random-effects term (e.g., intercept or slope
  - `grp` group, or level of the grouping variable
  - `condval` value of the conditional mode
  - `condsd` conditional standard deviation

Note

When a model has no zero inflation, the the `ranef` and `coef` print methods simplify the structure shown, by default. To show the full list structure, use `print(ranef(model), simplify=FALSE)` (or the analogous code for `coef`). In all cases, the full list structure is used to access the data frames (see example).

See Also

`fixef.glmmTMB`.

Examples

```r
if (requireNamespace("lme4")) {
  data(sleepstudy, package="lme4")
  model <- glmmTMB(Reaction ~ Days + (1|Subject), sleepstudy)
  rr <- ranef(model)
  print(rr, simplify=FALSE)
  ## extract Subject conditional modes for conditional model
  rr$cond$Subject
  as.data.frame(rr)
}
```
residuals.glmmTMB  
*Compute residuals for a glmmTMB object*

---

**Description**

Compute residuals for a glmmTMB object

**Usage**

```r
## S3 method for class 'glmmTMB'
residuals(object, type = c("response", "pearson"), ...)
```

**Arguments**

- `object` a “glmmTMB” object
- `type` (character) residual type
- `...` ignored, for method compatibility

---

**Salamanders**  
*Repeated counts of salamanders in streams*

---

**Description**

A data set containing counts of salamanders with site covariates and sampling covariates. Each of 23 sites was sampled 4 times. When using this data set, please cite Price et al. (2016) as well as the Dryad data package (Price et al. 2015).

**Usage**

```r
data(Salamanders)
```

**Format**

A data frame with 644 observations on the following 10 variables:

- `site` name of a location where repeated samples were taken
- `mined` factor indicating whether the site was affected by mountain top removal coal mining
- `cover` amount of cover objects in the stream (scaled)
- `sample` repeated sample
- `DOP` Days since precipitation (scaled)
- `Wtemp` water temperature (scaled)
- `DOY` day of year (scaled)
- `spp` abbreviated species name, possibly also life stage
- `count` number of salamanders observed
sigma.glmmTMB

References


Examples

```
require("glmmTMB")
data(Salamanders)

zpm3 = glmmTMB(count~spp * mined + (1|site), zi=spp * mined, Salamanders, family="poisson")
```

sigma.glmmTMB

Extract residual standard deviation or dispersion parameter

Description

For Gaussian models, sigma returns the value of the residual standard deviation; for other families, it returns the dispersion parameter, however it is defined for that particular family. See details for each family below.

Usage

```
## S3 method for class 'glmmTMB'
sigma(object, ...)
```

Arguments

- `object`: a “glmmTMB” fitted object
- `...`: (ignored; for method compatibility)

Details

The value returned varies by family:

- **gaussian**: returns the maximum likelihood estimate of the standard deviation (i.e., smaller than the results of `sigma(lm(...))` by a factor of (n-1)/n)
- **nbinom1**: returns an overdispersion parameter (usually denoted $\alpha$ as in Hardin and Hilbe (2007)): such that the variance equals $\mu(1 + \alpha)$.
- **nbinom2**: returns an overdispersion parameter (usually denoted $\theta$ or $k$); in contrast to most other families, larger $\theta$ corresponds to a lower variance which is $\mu(1 + \mu/\theta)$. 
**Gamma** Internally, glmmTMB fits Gamma responses by fitting a mean and a shape parameter; sigma is estimated as \((1/\text{sqrt}(\text{shape}))\), which will typically be close (but not identical to) that estimated by stats::sigma.default, which uses \(\text{sqrt}(\text{deviance}/\text{df.residual})\).

**beta** returns the value of \(\phi\), where the conditional variance is \(\mu(1 - \mu)/(1 + \phi)\) (i.e., increasing \(\phi\) decreases the variance.) This parameterization follows Ferrari and Cribari-Neto (2004) (and the betareg package):

**betabinomial** This family uses the same parameterization (governing the Beta distribution that underlies the binomial probabilities) as beta.

**genpois** returns the value of \(\phi\), where the variance is \(\mu\phi\)

**compois** returns the value of \(1/\nu\). When \(\nu = 1\), compois is equivalent to the Poisson distribution. There is no closed form equation for the variance, but it is approximately undersidpersed when \(1/\nu < 1\) and approximately oversidpersed when \(1/\nu > 1\). In this implementation, \(\mu\) is exactly the mean, which differs from the COMPoissonReg package (Sellers & Lotze 2015).

The most commonly used GLM families (binomial, poisson) have fixed dispersion parameters which are internally ignored.

**References**


---

**simulate.glmmTMB**

*Simulate from a glmmTMB fitted model*

**Description**

Simulate from a glmmTMB fitted model

**Usage**

```r
## S3 method for class 'glmmTMB'
simulate(object, nsim = 1, seed = NULL, ...)
```

**Arguments**

- `object` glmmTMB fitted model
- `nsim` number of response lists to simulate. Defaults to 1.
- `seed` random number seed
- `...` extra arguments
Details

Random effects are also simulated from their estimated distribution. Currently, it is not possible to condition on estimated random effects.

Value

returns a list of vectors. The list has length nsim. Each simulated vector of observations is the same size as the vector of response variables in the original data set. In the binomial family case each simulation is a two-column matrix with success/failure.

---

**tmbroot**

*Compute likelihood profile confidence intervals of a TMB object by root-finding (generalized from TMB::tmbprofile)*

---

Description

Compute likelihood profile confidence intervals of a TMB object by root-finding (generalized from TMB::tmbprofile)

Usage

`tmbroot(obj, name, target = 0.5 * qchisq(0.95, df = 1), lincomb, parm.range = c(NA, NA), sd.range = 7, trace = FALSE, continuation = FALSE)`

Arguments

- `obj`: a fitted glmmTMB object
- `name`: parameter index/name
- `target`: desired deviation from minimum log-likelihood. Default is set to retrieve the 95 if the objective function is a negative log-likelihood function
- `lincomb`: linear combination of parameters
- `parm.range`: lower and upper limits; if NA, a value will be guessed based on the parameter value and sd.range
- `sd.range`: in the absence of explicit parm.range values, the range chosen will be the parameter value plus or minus sd.range. May be specified as a two-element vector for different ranges below and above the parameter value.
- `trace`: report information?
- `continuation`: use continuation method, i.e. set starting parameters for non-focal parameters to solutions from previous fits?

Value

a two-element numeric vector containing the lower and upper limits (or NA if the target is not achieved in the range), with an attribute giving the total number of function iterations used
Calculate Variance-Covariance Matrix for a Fitted glmmTMB model

Calculate Variance-Covariance Matrix for a Fitted glmmTMB model

Usage

```r
## S3 method for class 'glmmTMB'
vcov(object, full = FALSE, ...)
```

Arguments

- `object`: a “glmmTMB” fit
- `full`: return a full variance-covariance matrix?
- `...`: ignored, for method compatibility

Value

By default (full==FALSE), a list of separate variance-covariance matrices for each model component (conditional, zero-inflation, dispersion). If full==TRUE, a single square variance-covariance matrix for all top-level model parameters (conditional, dispersion, and variance-covariance parameters)
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