

Package ‘broom.mixed’

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Type Package

Title Tidying Methods for Mixed Models

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Description Convert fitted objects from various R mixed-model packages into tidy data frames along the lines of the 'broom' package.

The package provides three

S3 generics for each model: tidy(), which summarizes a model's statistical findings such as coefficients of a regression; augment(), which adds columns to the original data such as predictions, residuals and cluster assignments; and glance(), which provides a one-row summary of model-level statistics.

Imports broom, coda, dplyr, forcats, methods, nlme, purrr, stringr, tibble, tidyr, furrr

Suggests brms, dotwhisker, knitr, testthat, gamlss, gamlss.data, ggplot2, GLMMadaptive, glmmADMB, glmmTMB, lmerTest, lme4, Matrix, MCMCglmm, mediation, mgcv, ordinal, pander, pbkrtest, posterior, rstan, rstanarm, rstantools, R2jags, TMB, rmarkdown

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BugReports <https://github.com/bbolker/broom.mixed/issues>

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augment.clmm	<i>name ordinal_tidiers</i>
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Description

the tidy method for clmm objects (from the ordinal package) lives in the broom package.

Usage

```
## S3 method for class 'clmm'
augment(x, data = model.frame(x), newdata, ...)
```

Arguments

x	An object of class merMod, such as those from lmer, glmer, or nlmer
data	original data this was fitted on; if not given this will attempt to be reconstructed
newdata	new data to be used for prediction; optional
...	Additional arguments (passed to confint.merMod for tidy; augment_columns for augment; ignored for glance)

augment.ranef.mer	<i>Augmentation for random effects (for caterpillar plots etc.)</i>
-------------------	---

Description

Augmentation for random effects (for caterpillar plots etc.)

Usage

```
## S3 method for class 'ranef.mer'
augment(x, ci.level = 0.9, reorder = TRUE, order.var = 1, ...)
```

Arguments

x	ranef (conditional mode) information from an lme4 fit, using ranef(. , condVar=TRUE)
ci.level	level for confidence intervals
reorder	reorder levels by conditional mode values?
order.var	numeric or character: which variable to use for ordering levels?
...	additional arguments (unused: for generic consistency)

Examples

```
if (require("lme4")) {
  load(system.file("extdata", "lme4_example.rda", package="broom.mixed"))
  rr <- ranef(lmm1, condVar=TRUE)
  aa <- broom::augment(rr)
  ## Q-Q plot:
  if (require(ggplot2) && require(dplyr)) {
    g0 <- ggplot(aa, aes(estimate, qq, xmin=lb, xmax=ub))+
      geom_errorbarh(height=0)+
      geom_point()+facet_wrap(~variable, scale="free_x")
    ## regular caterpillar plot:
    g1 <- ggplot(aa, aes(estimate, level, xmin=lb, xmax=ub))+
      geom_errorbarh(height=0)+
      geom_vline(xintercept=0, lty=2)+
      geom_point()+facet_wrap(~variable, scale="free_x")
    ## emphasize extreme values
    aa2 <- group_by(aa, grp, level)
    aa3 <- mutate(aa2, keep=any(estimate/std.error>2))
    ## Update caterpillar plot with extreme levels highlighted
    ## (highlight all groups with *either* extreme intercept *or*
    ##   extreme slope)
    ggplot(aa3, aes(estimate, level, xmin=lb, xmax=ub, colour=factor(keep)))+
      geom_errorbarh(height=0)+
      geom_vline(xintercept=0, lty=2)+
      geom_point()+facet_wrap(~variable, scale="free_x")+
      scale_colour_manual(values=c("black", "red"), guide=FALSE)
  }
}
```

brms_tidiers

Tidying methods for a brms model

Description

These methods tidy the estimates from [brmsfit-objects](#) (fitted model objects from the **brms** package) into a summary.

Usage

```
## S3 method for class 'brmsfit'
tidy(
  x,
  parameters = NA,
  effects = c("fixed", "ran_pars"),
  robust = FALSE,
  conf.int = TRUE,
  conf.level = 0.95,
  conf.method = c("quantile", "HPDinterval"),
  rhat = FALSE,
```

```

    ess = FALSE,
    fix.intercept = TRUE,
    exponentiate = FALSE,
    ...
)

## S3 method for class 'brmsfit'
glance(x, looic = FALSE, ...)

## S3 method for class 'brmsfit'
augment(x, data = stats::model.frame(x), newdata = NULL, se.fit = TRUE, ...)

```

Arguments

<code>x</code>	Fitted model object from the brms package. See brmsfit-class .
<code>parameters</code>	Names of parameters for which a summary should be returned, as given by a character vector or regular expressions. If NA (the default) summarized parameters are specified by the <code>effects</code> argument.
<code>effects</code>	A character vector including one or more of "fixed", "ran_vals", or "ran_pars". See the Value section for details.
<code>robust</code>	Whether to use median and median absolute deviation of the posterior distribution, rather than mean and standard deviation, to derive point estimates and uncertainty
<code>conf.int</code>	If TRUE columns for the lower (<code>conf.low</code>) and upper bounds (<code>conf.high</code>) of posterior uncertainty intervals are included.
<code>conf.level</code>	Defines the range of the posterior uncertainty <code>conf.int</code> , such that $100 * \text{conf.level}\%$ of the parameter's posterior distribution lies within the corresponding interval. Only used if <code>conf.int = TRUE</code> .
<code>conf.method</code>	method for computing confidence intervals ("quantile" or "HPDinterval")
<code>rhat</code>	whether to calculate the *Rhat* convergence metric (FALSE by default)
<code>ess</code>	whether to calculate the *effective sample size* (ESS) convergence metric (FALSE by default)
<code>fix.intercept</code>	rename "Intercept" parameter to "(Intercept)", to match behaviour of other model types?
<code>exponentiate</code>	whether to exponentiate the fixed-effect coefficient estimates and confidence intervals (common for logistic regression); if TRUE, also scales the standard errors by the exponentiated coefficient, transforming them to the new scale
<code>...</code>	Extra arguments, not used
<code>looic</code>	Should the LOO Information Criterion (and related info) be included? See loo.stanfit for details. (This can be slow for models fit to large datasets.)
<code>data</code>	data frame
<code>newdata</code>	new data frame
<code>se.fit</code>	return standard errors of fit?

Value

All tidying methods return a `data.frame` without rownames. The structure depends on the method chosen.

When `parameters = NA`, the `effects` argument is used to determine which parameters to summarize.

Generally, `tidy.brmsfit` returns one row for each coefficient, with at least three columns:

<code>term</code>	The name of the model parameter.
<code>estimate</code>	A point estimate of the coefficient (mean or median).
<code>std.error</code>	A standard error for the point estimate (sd or mad).

When `effects = "fixed"`, only population-level effects are returned.

When `effects = "ran_vals"`, only group-level effects are returned. In this case, two additional columns are added:

<code>group</code>	The name of the grouping factor.
<code>level</code>	The name of the level of the grouping factor.

Specifying `effects = "ran_pars"` selects the standard deviations and correlations of the group-level parameters.

If `conf.int = TRUE`, columns for the lower and upper bounds of the posterior `conf.int` computed.

Note

The names `'fixed'`, `'ran_pars'`, and `'ran_vals'` (corresponding to "non-varying", "hierarchical", and "varying" respectively in previous versions of the package), while technically inappropriate in a Bayesian setting where "fixed" and "random" effects are not well-defined, are used for compatibility with other (frequentist) mixed model types.

At present, the components of parameter estimates are separated by parsing the column names of `as_draws` (e.g. `r_patient[1, Intercept]` for the random effect on the intercept for patient 1, or `b_Tr1` for the fixed effect `Trt1`). We try to detect underscores in parameter names and warn, but detection may be imperfect.

See Also

[brms](#), [brmsfit-class](#)

Examples

```
## original model
## Not run:
brms_crossedRE <- brm(mpg ~ wt + (1|cyl) + (1+wt|gear), data = mtcars,
  iter = 500, chains = 2)

## End(Not run)

## too slow for CRAN (>5 seconds)
## load stored object
```

```

if (require("rstan") && require("brms")) {
  load(system.file("extdata", "brms_example.rda", package="broom.mixed"))

  fit <- brms_crossedRE
  tidy(fit)
  tidy(fit, parameters = "^sd_", conf.int = FALSE)
  tidy(fit, effects = "fixed", conf.method="HPDinterval")
  tidy(fit, effects = "ran_vals")
  tidy(fit, effects = "ran_pars", robust = TRUE)
  if (require("posterior")) {
    tidy(fit, effects = "ran_pars", rhat = TRUE, ess = TRUE)
  }
}
if (require("rstan") && require("brms")) {
  # glance method
  glance(fit)
  ## this example will give a warning that it should be run with
  ## reloo=TRUE; however, doing this will fail
  ## because the \code{fit} object has been stripped down to save space
  suppressWarnings(glance(fit, looic = TRUE, cores = 1))
  head(augment(fit))
}

```

compact

Remove NULL items in a vector or list

Description

Remove NULL items in a vector or list

Usage

```
compact(x)
```

Arguments

x a vector or list

fixef.MCMCglmm

Extract fixed effects from an MCMCglmm object

Description

Function designed to extract the fixed effects from an MCMCglmm model object. Can either extract all samples from the fixed effects posteriors or return the posterior means.

Usage

```
## S3 method for class 'MCMCglmm'
fixef(object, use = c("all", "mean"), ...)
```

Arguments

object	An MCMCglmm model object to extract the effects from
use	A character string indicating whether to extract all posterior samples or the mean of the posteriors. Defaults to "all".
...	Arguments passed on to the worker function.

Value

A matrix of the fixed effects

See Also

[ranef.MCMCglmm](#)

Examples

```
## Not run:
# a simple MCMCglmm model
data(PlodiaPO)
m <- MCMCglmm(PO ~ 1, random= ~ FSfamily, data=PlodiaPO, verbose=FALSE)

# only extract average fixed effects
fixef(m, use = "mean")

# histogram of posterior samples of fixed effects
hist(fixef(m))
# matches the mean
rowMeans(fixef(m))

## End(Not run)
```

Description

Tidying methods for "gamlss" objects from the gamlss package.

Usage

```
## S3 method for class 'gamlss'
tidy(x, quick = FALSE, conf.int = FALSE, conf.level = 0.95, ...)
```

Arguments

x	A "gamlss" object
quick	Whether to perform a fast version, and return only the coefficients
conf.int	whether to return confidence intervals
conf.level	confidence level for CI
...	arguments passed to <code>confint.gamlss</code>

Value

All tidying methods return a data.frame without rownames, whose structure depends on the method chosen.

A tibble with one row for each coefficient, containing columns:

parameter	type of coefficient being estimated: mu, sigma, nu, or tau
term	term in the model being estimated and tested
estimate	estimated coefficient
std.error	standard error
statistic	t-statistic
p.value	two-sided p-value

Examples

```
if (requireNamespace("gamlss", quietly = TRUE) &&
    requireNamespace("gamlss.data", quietly = TRUE)) {
  data(abdom, package="gamlss.data")
  ## Not run:
  mod <- gamlss(y~pb(x), sigma.fo=~pb(x), family=BCT,
               data=abdom, method=mixed(1,20))

  ## End(Not run)
  ## load stored object
  mod <- readRDS(system.file("extdata", "gamlss_example.rds",
```

```

      tidy(mod)
    }
    package="broom.mixed"))

```

get_methods	<i>Retrieve all method/class combinations currently provided by the broom.mixed package</i>
-------------	---

Description

Retrieve all method/class combinations currently provided by the broom.mixed package

Usage

```
get_methods()
```

Examples

```
print(get_methods(), n = Inf)
```

glmmadmb_tidiers	<i>Tidying methods for glmmADMB models</i>
------------------	--

Description

These methods tidy the coefficients of glmmADMB models

Usage

```

## S3 method for class 'glmmadmb'
tidy(
  x,
  effects = c("fixed", "ran_pars"),
  component = "cond",
  scales = NULL,
  ran_prefix = NULL,
  conf.int = FALSE,
  conf.level = 0.95,
  conf.method = "Wald",
  ...
)

## S3 method for class 'glmmadmb'
augment(x, data = stats::model.frame(x), newdata, ...)

## S3 method for class 'glmmadmb'
glance(x, ...)

```

Arguments

<code>x</code>	An object of class <code>glmmadmb</code> <code>glmer</code> , or <code>nlmer</code>
<code>effects</code>	A character vector including one or more of "fixed" (fixed-effect parameters), "ran_pars" (variances and covariances or standard deviations and correlations of random effect terms) or "ran_vals" (conditional modes/BLUPs/latent variable estimates)
<code>component</code>	Which component(s) to report for (e.g., conditional, zero-inflation, dispersion: at present only works for "cond")
<code>scales</code>	scales on which to report the variables: for random effects, the choices are "sd-cor" (standard deviations and correlations: the default if <code>scales</code> is <code>NULL</code>) or "varcov" (variances and covariances). <code>NA</code> means no transformation, appropriate e.g. for fixed effects; inverse-link transformations (exponentiation or logistic) are not yet implemented, but may be in the future.
<code>ran_prefix</code>	a length-2 character vector specifying the strings to use as prefixes for self- (variance/standard deviation) and cross- (covariance/correlation) random effects terms
<code>conf.int</code>	whether to include a confidence interval
<code>conf.level</code>	confidence level for CI
<code>conf.method</code>	method for computing confidence intervals (see confint.merMod)
<code>...</code>	extra arguments (not used)
<code>data</code>	original data this was fitted on; if not given this will attempt to be reconstructed
<code>newdata</code>	new data to be used for prediction; optional

Details

When the modeling was performed with `na.action = "na.omit"` (as is the typical default), rows with `NA` in the initial data are omitted entirely from the augmented data frame. When the modeling was performed with `na.action = "na.exclude"`, one should provide the original data as a second argument, at which point the augmented data will contain those rows (typically with `NA`s in place of the new columns). If the original data is not provided to augment and `na.action = "na.exclude"`, a warning is raised and the incomplete rows are dropped.

Value

All tidying methods return a `tbl_df` without rownames. The structure depends on the method chosen.

`tidy` returns one row for each estimated effect, either with groups depending on the `effects` parameter. It contains the columns

<code>group</code>	the group within which the random effect is being estimated: <code>NA</code> for fixed effects
<code>level</code>	level within group (<code>NA</code> except for modes)
<code>term</code>	term being estimated
<code>estimate</code>	estimated coefficient
<code>std.error</code>	standard error

statistic	t- or Z-statistic (NA for modes)
p.value	P-value computed from t-statistic (may be missing/NA)

augment returns one row for each original observation, with columns (each prepended by a .) added. Included are the columns

.fitted	predicted values
.resid	residuals
.fixed	predicted values with no random effects

Also added for "merMod" objects, but not for "mer" objects, are values from the response object within the model (of type lmResp, glmResp, nlsResp, etc). These include ".mu", ".offset", ".sqrtXwt", ".sqrtwt", ".eta".

glance returns one row with the columns

sigma	the square root of the estimated residual variance
logLik	the data's log-likelihood under the model
AIC	the Akaike Information Criterion
BIC	the Bayesian Information Criterion
deviance	deviance

See Also

[na.action](#)

Examples

```
if (require("glmmADMB") && require("lme4")) {
  ## original model
  ## Not run:
  data("sleepstudy", package="lme4")
  lmm1 <- glmmadmb(Reaction ~ Days + (Days | Subject), sleepstudy,
    family="gaussian")

  ## End(Not run)
  ## load stored object
  load(system.file("extdata", "glmmADMB_example.rda", package="broom.mixed"))
  tidy(lmm1, effects = "fixed")
  tidy(lmm1, effects = "fixed", conf.int=TRUE)
  ## tidy(lmm1, effects = "fixed", conf.int=TRUE, conf.method="profile")
  ## tidy(lmm1, effects = "ran_vals", conf.int=TRUE)
  head(augment(lmm1, sleepstudy))
  glance(lmm1)

  glmm1 <- glmmadmb(cbind(incidence, size - incidence) ~ period + (1 | herd),
    data = cbpp, family = "binomial")
  tidy(glmm1)
  tidy(glmm1, effects = "fixed")
  head(augment(glmm1, cbpp))
  glance(glmm1)
```

```
}
```

glmmTMB_tidiers

Tidying methods for glmmTMB models

Description

These methods tidy the coefficients of mixed effects models, particularly responses of the `merMod` class

Usage

```
## S3 method for class 'glmmTMB'
tidy(
  x,
  effects = c("ran_pars", "fixed"),
  component = c("cond", "zi"),
  scales = NULL,
  ran_prefix = NULL,
  conf.int = FALSE,
  conf.level = 0.95,
  conf.method = "Wald",
  exponentiate = FALSE,
  ...
)

## S3 method for class 'glmmTMB'
augment(x, data = stats::model.frame(x), newdata = NULL, ...)

## S3 method for class 'glmmTMB'
glance(x, ...)
```

Arguments

<code>x</code>	An object of class <code>merMod</code> , such as those from <code>lmer</code> , <code>glmer</code> , or <code>nlmer</code>
<code>effects</code>	A character vector including one or more of "fixed" (fixed-effect parameters), "ran_pars" (variances and covariances or standard deviations and correlations of random effect terms) or "ran_vals" (conditional modes/BLUPs/latent variable estimates)
<code>component</code>	which component to extract (e.g. <code>cond</code> for conditional effects (i.e., traditional fixed effects); <code>zi</code> for zero-inflation model; <code>disp</code> for dispersion model)
<code>scales</code>	scales on which to report the variables: for random effects, the choices are "sd-cor" (standard deviations and correlations: the default if <code>scales</code> is <code>NULL</code>) or "varcov" (variances and covariances). <code>NA</code> means no transformation, appropriate e.g. for fixed effects; inverse-link transformations (exponentiation or logistic) are not yet implemented, but may be in the future.

<code>ran_prefix</code>	a length-2 character vector specifying the strings to use as prefixes for self- (variance/standard deviation) and cross- (covariance/correlation) random effects terms
<code>conf.int</code>	whether to include a confidence interval
<code>conf.level</code>	confidence level for CI
<code>conf.method</code>	method for computing confidence intervals (see confint.merMod)
<code>exponentiate</code>	whether to exponentiate the fixed-effect coefficient estimates and confidence intervals (common for logistic regression); if TRUE, also scales the standard errors by the exponentiated coefficient, transforming them to the new scale
<code>...</code>	extra arguments (not used)
<code>data</code>	original data this was fitted on; if not given this will attempt to be reconstructed
<code>newdata</code>	new data to be used for prediction; optional

Details

When the modeling was performed with `na.action = "na.omit"` (as is the typical default), rows with NA in the initial data are omitted entirely from the augmented data frame. When the modeling was performed with `na.action = "na.exclude"`, one should provide the original data as a second argument, at which point the augmented data will contain those rows (typically with NAs in place of the new columns). If the original data is not provided to augment and `na.action = "na.exclude"`, a warning is raised and the incomplete rows are dropped.

Value

All tidying methods return a tibble. The structure depends on the method chosen.

`tidy` returns one row for each estimated effect, either with groups depending on the effects parameter. It contains the columns

<code>group</code>	the group within which the random effect is being estimated: NA for fixed effects
<code>level</code>	level within group (NA except for modes)
<code>term</code>	term being estimated
<code>estimate</code>	estimated coefficient
<code>std.error</code>	standard error
<code>statistic</code>	t- or Z-statistic (NA for modes)
<code>p.value</code>	P-value computed from t-statistic (may be missing/NA)

`augment` returns one row for each original observation, with columns (each prepended by a `.`) added. Included are the columns

<code>.fitted</code>	predicted values
<code>.resid</code>	residuals
<code>.fixed</code>	predicted values with no random effects

`glance` returns one row with the columns

<code>sigma</code>	the square root of the estimated residual variance
--------------------	--

logLik	the data's log-likelihood under the model
AIC	the Akaike Information Criterion
BIC	the Bayesian Information Criterion
deviance	deviance

Note

zero-inflation parameters (including the intercept) are reported on the logit scale

See Also

[na.action](#)

Examples

```

if (require("glmmTMB") && require("lme4"))
  ## &&
  ## make sure package versions are OK
  ## checkDepPackageVersion(dep_pkg = "TMB",
  ##                          this_pkg = "glmmTMB",
  ##                          warn = FALSE) &&
  ## checkDepPackageVersion(dep_pkg = "Matrix",
  ##                          this_pkg = "TMB",
  ##                          warn = FALSE)
)
{
  data("sleepstudy", package="lme4")
  ## original model:
  ## Not run:
  lmm1 <- glmmTMB(Reaction ~ Days + (Days | Subject), sleepstudy)

## End(Not run)
## load stored object
L <- load(system.file("extdata", "glmmTMB_example.rda", package="broom.mixed"))
for (obj in L) {
  assign(obj, glmmTMB::up2date(get(obj)))
}
tidy(lmm1)
tidy(lmm1, effects = "fixed")
tidy(lmm1, effects = "fixed", conf.int=TRUE)
tidy(lmm1, effects = "fixed", conf.int=TRUE, conf.method="uniroot")
## FIX: tidy(lmm1, effects = "ran_vals", conf.int=TRUE)
head(augment(lmm1, sleepstudy))
glance(lmm1)

## original model:
## glmm1 <- glmmTMB(incidence/size ~ period + (1 | herd),
##                  data = cbpp, family = binomial, weights=size)
tidy(glmm1)
tidy(glmm1, effects = "fixed")
tidy(glmm1, effects = "fixed", exponentiate=TRUE)

```

```

  tidy(glm1, effects = "fixed", conf.int=TRUE, exponentiate=TRUE)
  head(augment(glm1, cbpp))
  head(augment(glm1, cbpp, type.residuals="pearson"))
  glance(glm1)
## Not run:
  ## profile CIs - a little bit slower but more accurate
  tidy(glm1, effects = "fixed", conf.int=TRUE, exponentiate=TRUE, conf.method="profile")

## End(Not run)
}

```

insert_NAs	<i>insert a row of NAs into a data frame wherever another data frame has NAs</i>
------------	--

Description

insert a row of NAs into a data frame wherever another data frame has NAs

Usage

```
insert_NAs(x, original)
```

Arguments

x	data frame that has one row for each non-NA row in original
original	data frame with NAs

lme4_tidiers	<i>Tidying methods for mixed effects models</i>
--------------	---

Description

These methods tidy the coefficients of `lme4::lmer` and `lme4::glmer` models (i.e., `merMod` objects). Methods are also provided for `allFit` objects.

Usage

```

## S3 method for class 'merMod'
tidy(
  x,
  effects = c("ran_pars", "fixed"),
  scales = NULL,
  exponentiate = FALSE,
  exponentiate_ran_coefs = FALSE,
  ran_prefix = NULL,

```

```

    conf.int = FALSE,
    conf.level = 0.95,
    conf.method = "Wald",
    ddf.method = NULL,
    profile = NULL,
    debug = FALSE,
    ...
)

## S3 method for class 'rmlmerMod'
tidy(
  x,
  effects = c("ran_pars", "fixed"),
  scales = NULL,
  exponentiate = FALSE,
  exponentiate_ran_coefs = FALSE,
  ran_prefix = NULL,
  conf.int = FALSE,
  conf.level = 0.95,
  conf.method = "Wald",
  ddf.method = NULL,
  profile = NULL,
  debug = FALSE,
  ...
)

## S3 method for class 'merMod'
augment(x, data = stats::model.frame(x), newdata, ...)

## S3 method for class 'merMod'
glance(x, ...)

```

Arguments

<code>x</code>	An object of class <code>merMod</code> , such as those from <code>lmer</code> , <code>glmer</code> , or <code>nlmer</code>
<code>effects</code>	A character vector including one or more of "fixed" (fixed-effect parameters); "ran_pars" (variances and covariances or standard deviations and correlations of random effect terms); "ran_vals" (conditional modes/BLUPs/latent variable estimates); or "ran_coefs" (predicted parameter values for each group, as returned by <code>coef.merMod</code>)
<code>scales</code>	scales on which to report the variables: for random effects, the choices are "sd-cor" (standard deviations and correlations: the default if <code>scales</code> is <code>NULL</code>) or "vcov" (variances and covariances). <code>NA</code> means no transformation, appropriate e.g. for fixed effects.
<code>exponentiate</code>	whether to exponentiate the fixed-effect coefficient estimates and confidence intervals (common for logistic regression); if <code>TRUE</code> , also scales the standard errors by the exponentiated coefficient, transforming them to the new scale

<code>exponentiate_ran_coefs</code>	whether to exponentiate the predicted parameter values for each group
<code>ran_prefix</code>	a length-2 character vector specifying the strings to use as prefixes for self- (variance/standard deviation) and cross- (covariance/correlation) random effects terms
<code>conf.int</code>	whether to include a confidence interval
<code>conf.level</code>	confidence level for CI
<code>conf.method</code>	method for computing confidence intervals (see <code>lme4::confint.merMod</code>)
<code>ddf.method</code>	the method for computing the degrees of freedom and t-statistics (only applicable when using the lmerTest package: see summary.lmerModLmerTest)
<code>profile</code>	pre-computed profile object, for speed when using <code>conf.method="profile"</code>
<code>debug</code>	print debugging output?
<code>...</code>	Additional arguments (passed to <code>confint.merMod</code> for <code>tidy</code> ; <code>augment_columns</code> for <code>augment</code> ; ignored for <code>glance</code>)
<code>data</code>	original data this was fitted on; if not given this will attempt to be reconstructed
<code>newdata</code>	new data to be used for prediction; optional

Details

When the modeling was performed with `na.action = "na.omit"` (as is the typical default), rows with NA in the initial data are omitted entirely from the augmented data frame. When the modeling was performed with `na.action = "na.exclude"`, one should provide the original data as a second argument, at which point the augmented data will contain those rows (typically with NAs in place of the new columns). If the original data is not provided to `augment` and `na.action = "na.exclude"`, a warning is raised and the incomplete rows are dropped.

Value

All tidying methods return a `data.frame` without rownames. The structure depends on the method chosen.

`tidy` returns one row for each estimated effect, either with groups depending on the effects parameter. It contains the columns

<code>group</code>	the group within which the random effect is being estimated: "fixed" for fixed effects
<code>level</code>	level within group (NA except for modes)
<code>term</code>	term being estimated
<code>estimate</code>	estimated coefficient
<code>std.error</code>	standard error
<code>statistic</code>	t- or Z-statistic (NA for modes)
<code>p.value</code>	P-value computed from t-statistic (may be missing/NA)

`augment` returns one row for each original observation, with columns (each prepended by a `.`) added. Included are the columns

.fitted	predicted values
.resid	residuals
.fixed	predicted values with no random effects

Also added for "merMod" objects, but not for "mer" objects, are values from the response object within the model (of type `lmResp`, `glmResp`, `nlsResp`, etc). These include `".mu"`, `".offset"`, `".sqrtXwt"`, `".sqrtwt"`, `".eta"`.

`glance` returns one row with the columns

nobs	the number of observations
sigma	the square root of the estimated residual variance
logLik	the data's log-likelihood under the model
AIC	the Akaike Information Criterion
BIC	the Bayesian Information Criterion
deviance	deviance

See Also

[na.action](#)

Examples

```
if (require("lme4")) {
  ## original model
  ## Not run:
  lmm1 <- lmer(Reaction ~ Days + (Days | Subject), sleepstudy)

  ## End(Not run)
  ## load stored object
  load(system.file("extdata", "lme4_example.rda", package="broom.mixed"))
  (tt <- tidy(lmm1))
  tidy(lmm1, effects = "fixed")
  tidy(lmm1, effects = "fixed", conf.int=TRUE)
  tidy(lmm1, effects = "fixed", conf.int=TRUE, conf.method="profile")
  ## lmm1_prof <- profile(lmm1) # generated by extdata/runexamples
  tidy(lmm1, conf.int=TRUE, conf.method="profile", profile=lmm1_prof)
  ## conditional modes (group-level deviations from population-level estimate)
  tidy(lmm1, effects = "ran_vals", conf.int=TRUE)
  ## coefficients (group-level estimates)
  (rcoef1 <- tidy(lmm1, effects = "ran_coefs"))
  if (require(tidyr) && require(dplyr)) {
    ## reconstitute standard coefficient-by-level table
    spread(rcoef1, key=term, value=estimate)
    ## split ran_pars into type + term; sort fixed/sd/cor
    (tt %>% separate(term, c("type", "term"), sep="___", fill="left")
     %>% arrange(!is.na(type), desc(type)))
  }
  head(augment(lmm1, sleepstudy))
  glance(lmm1)
```

```

glmm1 <- glmer(cbind(incidence, size - incidence) ~ period + (1 | herd),
              data = cbpp, family = binomial)
tidy(glmm1)
tidy(glmm1, exponentiate=TRUE)
tidy(glmm1, effects = "fixed")
## suppress warning about influence.merMod
head(suppressWarnings(augment(glmm1, cbpp)))
glance(glmm1)

startvec <- c(Asym = 200, xmid = 725, scal = 350)
nm1 <- nlmer(circumference ~ SSlogis(age, Asym, xmid, scal) ~ Asym|Tree,
            Orange, start = startvec)
## suppress warnings about var-cov matrix ...
op <- options(warn=-1)
tidy(nm1)
tidy(nm1, effects = "fixed")
options(op)
head(augment(nm1, Orange))
glance(nm1)
detach("package:lme4")
}
if (require("lmerTest")) {
  lmm1 <- lmer(Reaction ~ Days + (Days | Subject), sleepstudy)
  tidy(lmm1)
  glance(lmm1)
  detach("package:lmerTest") # clean up
}

```

mediation_tidiers

Tidying methods for mediation analyses involving mixed effects models

Description

These methods tidy the coefficients of `mediation::mediate` output (i.e., `mediate.mer` objects) when `lme4::lmer` and `lme4::glmer` models (i.e., `merMod` objects) provide the input.

Usage

```

## S3 method for class 'mediate.mer'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

```

Arguments

<code>x</code>	an object of class <code>mediate.mer</code> , as from <code>mediate</code> using <code>lmer</code> , <code>glmer</code> , or <code>nlmer</code> models
<code>conf.int</code>	whether to include a confidence interval
<code>conf.level</code>	confidence level for CI
<code>...</code>	additional arguments (unused: for generic consistency)

Value

All tidying methods return a `data.frame` without rownames. The structure depends on the method chosen.

`tidy` returns one row for each estimated effect: first the mediated effect in the control and treatment groups, respectively, then the direct effect in each group. It contains the columns

<code>term</code>	term being estimated
<code>estimate</code>	estimated coefficient
<code>std.error</code>	standard error
<code>p.value</code>	P-value computed from t-statistic (may be missing/NA)

See Also

[mediate](#), [tidy.mediate](#)

Examples

```
if (require("lme4") && require("mediation")) {
  ## Borrowed from \code{help(mediation::mediate)}:
  ## Not run:
  ## Varying intercept for mediator
  mod_m <- glmer(job_dich ~ treat + econ_hard + (1 | educ),
                 family = binomial(link = "probit"), data = jobs)
  ## Varying intercept and slope for outcome
  mod_y <- glmer(work1 ~ treat + job_dich + econ_hard + (1 + treat | occp),
                 family = binomial(link = "probit"), data = jobs)
  ## Output based on mediator group ("educ")
  mod_med <- mediate(mod_m, mod_y, treat = "treat",
                    mediator = "job_dich", sims=50, group.out="educ")

  ## End(Not run)
  ## Load stored objects
  load(system.file("extdata", "mediation_example.rda", package="broom.mixed"))
  ## Tidy outputs
  tidy(mod_m)
  tidy(mod_y)
  tidy(mod_med)
}
```

Description

These methods tidy the coefficients of mixed effects models of the `lme` class from functions of the `nlme` package.

Usage

```
## S3 method for class 'lme'
tidy(
  x,
  effects = c("var_model", "ran_pars", "fixed"),
  scales = NULL,
  conf.int = FALSE,
  conf.level = 0.95,
  ...
)

## S3 method for class 'lme'
augment(x, data = x$data, newdata, ...)

## S3 method for class 'lme'
glance(x, ...)

## S3 method for class 'gls'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

## S3 method for class 'gls'
augment(x, data = nlme::getData(x), newdata, ...)
```

Arguments

<code>x</code>	An object of class <code>lme</code> , such as those from <code>lme</code> or <code>nlme</code>
<code>effects</code>	One or more of "var_model", "ran_pars", "fixed", "ran_vals", and/or "ran_coefs".
<code>scales</code>	scales on which to report the variables: for random effects, the choices are "sd-cor" (standard deviations and correlations: the default if <code>scales</code> is <code>NULL</code>) or "vcov" (variances and covariances). <code>NA</code> means no transformation, appropriate e.g. for fixed effects.
<code>conf.int</code>	whether to include a confidence interval
<code>conf.level</code>	confidence level for CI
<code>...</code>	extra arguments (not used)
<code>data</code>	original data this was fitted on; if not given this will attempt to be reconstructed
<code>newdata</code>	new data to be used for prediction; optional

Details

When the modeling was performed with `na.action = "na.omit"` (as is the typical default), rows with `NA` in the initial data are omitted entirely from the augmented data frame. When the modeling was performed with `na.action = "na.exclude"`, one should provide the original data as a second argument, at which point the augmented data will contain those rows (typically with `NA`s in place of the new columns). If the original data is not provided to augment and `na.action = "na.exclude"`, a warning is raised and the incomplete rows are dropped.

Value

All tidying methods return a `data.frame` without rownames. The structure depends on the method chosen.

`tidy` returns one row for each estimated effect, either random or fixed depending on the `effects` parameter. If `effects = "ran_vals"` (or `"ran_pars"`), it contains the columns

<code>group</code>	the group within which the random effect is being estimated
<code>level</code>	level within group
<code>term</code>	term being estimated
<code>estimate</code>	estimated coefficient
<code>estimated</code>	This column is only included if some parameters are fixed. TRUE if the residual error is estimated and FALSE if the residual error is fixed.

If `effects="fixed"`, `tidy` returns the columns

<code>term</code>	fixed term being estimated
<code>estimate</code>	estimate of fixed effect
<code>std.error</code>	standard error
<code>statistic</code>	t-statistic
<code>p.value</code>	P-value computed from t-statistic

If `effects="var_model"` (the `weights` argument to the model), `tidy` returns the columns defined in the help for `tidy.varFunc`.

`augment` returns one row for each original observation, with columns (each prepended by a `.`) added. Included are the columns

<code>.fitted</code>	predicted values
<code>.resid</code>	residuals
<code>.fixed</code>	predicted values with no random effects

`glance` returns one row with the columns

<code>sigma</code>	the square root of the estimated residual variance
<code>logLik</code>	the data's log-likelihood under the model
<code>AIC</code>	the Akaike Information Criterion
<code>BIC</code>	the Bayesian Information Criterion
<code>deviance</code>	returned as NA. To quote Brian Ripley on R-help https://stat.ethz.ch/pipermail/r-help/2006-May/104744.html , "McCullagh & Nelder (1989) would be the authoritative [sic] reference, but the 1982 first edition manages to use 'deviance' in three separate senses on one page."

See Also

[na.action](#)

Examples

```

if (require("nlme") && require("lme4")) {
  data("sleepstudy", package="lme4")
  ## original model
  ## Not run:
  lmm1 <- lme(Reaction ~ Days, random=~ Days|Subject, sleepstudy)

  ## End(Not run)
  ## load stored object
  load(system.file("extdata", "nlme_example.rda", package="broom.mixed"))
  tidy(lmm1)
  tidy(lmm1, effects = "fixed")
  tidy(lmm1, conf.int = TRUE)
  tidy(lmm1, conf.int = TRUE, conf.level = 0.8)
  tidy(lmm1, effects = "ran_pars")
  tidy(lmm1, effects = "ran_vals")
  tidy(lmm1, effects = "ran_coefs")
  head(augment(lmm1, sleepstudy))
  glance(lmm1)

  startvec <- c(Asym = 200, xmid = 725, scal = 350)
  nm1 <- nlme(circumference ~ SSlogis(age, Asym, xmid, scal),
    data = Orange,
    fixed = Asym + xmid + scal ~1,
    random = Asym ~1,
    start = startvec)
  tidy(nm1)
  tidy(nm1, effects = "fixed")
  head(augment(nm1, Orange))
  glance(nm1)

  gls1 <- gls(follicles ~ sin(2*pi*Time) + cos(2*pi*Time), Ovary,
    correlation = corAR1(form = ~ 1 | Mare))
  tidy(gls1)
  glance(gls1)
  head(augment(gls1))
}

```

ranef.MCMCglmm

Extract random effects from an MCMCglmm object

Description

Function designed to extract the random effects from an MCMCglmm model object. Can either extract all samples from the random effects posteriors or return the posterior means.

Usage

```

## S3 method for class 'MCMCglmm'
ranef(object, use = c("all", "mean"), ...)

```

Arguments

object	An MCMCglmm model object to extract the effects from
use	A character string indicating whether to extract all posterior samples or the mean of the posteriors. Defaults to "all".
...	Arguments passed on to the worker function.

Value

A matrix of the fixed effects

See Also

[fixef.MCMCglmm](#)

Examples

```
## Not run:
# a simple MCMCglmm model
data(PlodiaPO)
m <- MCMCglmm(PO ~ 1, random= ~ FSfamily, data=PlodiaPO, pr=TRUE, verbose=FALSE)

# only extract average fixed effects
head(ranef(m, use = "mean"))

# histogram of posterior samples of fixed effects
hist(ranef(m)[1, ])
# matches the mean
rowMeans(ranef(m)[1:6, ])

## End(Not run)
```

ranefLevels	<i>Extract the levels of factors used for random effects in MCMCglmm objects</i>
-------------	--

Description

Extract the levels of factors used for random effects in MCMCglmm objects

Usage

```
ranefLevels(object, data, ...)
```

Arguments

object	An MCMCglmm model object
data	The dataset used for the model
...	Not currently used

See Also

[paramNamesMCMCglmm](#), [ranef.MCMCglmm](#)

Examples

```
## Not run:
# a simple MCMCglmm model
data(PlodiaPO)
m <- MCMCglmm(PO ~ 1, random = ~ FSfamily, data = PlodiaPO, verbose=FALSE)

# extract the random effects levels
ranefLevels(m, PlodiaPO)

## End(Not run)
```

rstanarm_tidiers

Tidying methods for an rstanarm model

Description

These methods tidy the estimates from rstanarm fits (stan_glm, stan_glmer, etc.) into a summary.

Usage

```
## S3 method for class 'stanreg'
tidy(
  x,
  effects = c("fixed", "ran_pars"),
  conf.int = FALSE,
  conf.level = 0.9,
  conf.method = c("quantile", "HPDinterval"),
  exponentiate = FALSE,
  ...
)

## S3 method for class 'stanreg'
glance(x, looic = FALSE, ...)
```

Arguments

x	Fitted model object from the rstanarm package. See stanreg-objects .
effects	A character vector including one or more of "fixed", "ran_vals", or "ran_pars". See the Value section for details.
conf.int	If TRUE columns for the lower (conf.low) and upper (conf.high) bounds of the 100*prob% posterior uncertainty intervals are included. See posterior_interval for details.

<code>conf.level</code>	See posterior_interval .
<code>conf.method</code>	method for computing confidence intervals ("quantile" or "HPDinterval")
<code>exponentiate</code>	whether to exponentiate the fixed-effect coefficient estimates and confidence intervals (common for logistic regression); if TRUE, also scales the standard errors by the exponentiated coefficient, transforming them to the new scale
<code>...</code>	For glance, if looic=TRUE, optional arguments to loo.stanfit .
<code>looic</code>	Should the LOO Information Criterion (and related info) be included? See loo.stanfit for details. (This can be slow for models fit to large datasets.)

Value

All tidying methods return a `data.frame` without rownames. The structure depends on the method chosen.

When `effects="fixed"` (the default), `tidy.stanreg` returns one row for each coefficient, with three columns:

<code>term</code>	The name of the corresponding term in the model.
<code>estimate</code>	A point estimate of the coefficient (posterior median).
<code>std.error</code>	A standard error for the point estimate based on mad . See the <i>Uncertainty estimates</i> section in print.stanreg for more details.

For models with group-specific parameters (e.g., models fit with [stan_glmer](#)), setting `effects="ran_vals"` selects the group-level parameters instead of the non-varying regression coefficients. Additional columns are added indicating the level and group. Specifying `effects="ran_pars"` selects the standard deviations and (for certain models) correlations of the group-level parameters.

Setting `effects="auxiliary"` will select parameters other than those included by the other options. The particular parameters depend on which **rstanarm** modeling function was used to fit the model. For example, for models fit using [stan_glm](#) the overdispersion parameter is included if `effects="aux"`, for [stan_lm](#) the auxiliary parameters include the residual SD, R^2 , and $\log(\text{fit_ratio})$, etc.

`glance` returns one row with the columns

<code>algorithm</code>	The algorithm used to fit the model.
<code>pss</code>	The posterior sample size (except for models fit using optimization).
<code>nobs</code>	The number of observations used to fit the model.
<code>sigma</code>	The square root of the estimated residual variance, if applicable. If not applicable (e.g., for binomial GLMs), <code>sigma</code> will be given the value 1 in the returned object.

If `looic=TRUE`, then the following additional columns are also included:

<code>looic</code>	The LOO Information Criterion.
<code>elpd_loo</code>	The expected log predictive density ($\text{elpd_loo} = -2 * \text{looic}$).
<code>p_loo</code>	The effective number of parameters.

See Also

[summary, stanfit-method](#)

Examples

```
if (require("rstanarm")) {
## Not run:
#'    ## original models
fit <- stan_glmer(mpg ~ wt + (1|cyl) + (1+wt|gear), data = mtcars,
                 iter = 500, chains = 2)
fit2 <- stan_glmer((mpg>20) ~ wt + (1 | cyl) + (1 + wt | gear),
                  data = mtcars,
                  family = binomial,
                  iter = 500, chains = 2)

## End(Not run)
## load example data
load(system.file("extdata", "rstanarm_example.rda", package="broom.mixed"))

# non-varying ("population") parameters
tidy(fit, conf.int = TRUE, conf.level = 0.5)
tidy(fit, conf.int = TRUE, conf.method = "HPDinterval", conf.level = 0.5)

# exponentiating (in this case, from log-odds to odds ratios)
(tidy(fit2, conf.int = TRUE, conf.level = 0.5)
 |> dplyr::filter(term != "(Intercept)"))
)
(tidy(fit2, conf.int = TRUE, conf.level = 0.5, exponentiate = TRUE)
 |> dplyr::filter(term != "(Intercept)"))
)

# hierarchical sd & correlation parameters
tidy(fit, effects = "ran_pars")

# group-specific deviations from "population" parameters
tidy(fit, effects = "ran_vals")

# glance method
glance(fit)
## Not run:
glance(fit, looic = TRUE, cores = 1)

## End(Not run)
} ## if require("rstanarm")
```

Description

Function designed to extract the standard deviation of the random effects from an MCMCglmm model object. Note that this is not the same as the posterior distribution of (co)variance matrices. It is based on the posterior distribution of the random effects. This also means it requires `pr=TRUE` to be set in the model for the information to be saved. Can optionally return standard deviation of random effects after back transforming to the response metric. Currently probabilities, but only for ordinal family models (`family="ordinal"`).

Usage

```
stdranef(object, which, type = c("lp", "response"), ...)
```

Arguments

<code>object</code>	An MCMCglmm model object to extract the effects from
<code>which</code>	A list of random effects to extract or their numeric positions. If there are two numbers in a list, effects are simultaneous.
<code>type</code>	A character string indicating whether to calculate the standard deviation on the linear predictor metric, 'lp' or response, 'response'.
<code>...</code>	Not currently used.

Value

A list of class `postMCMCglmmRE` with means (M) and individual estimates (Data)

Examples

```
## Not run:
# a simple MCMCglmm model
data(PlodiaP0)
PlodiaP0 <- within(PlodiaP0, {
  P02 <- cut(P0, quantile(P0, c(0, .33, .66, 1)))
  plate <- factor(plate)
})

m <- MCMCglmm(P02 ~ 1, random = ~ FSfamily + plate,
  family = "ordinal", data = PlodiaP0,
  prior = list(
    R = list(V = 1, fix = 1),
    G = list(
      G1 = list(V = 1, nu = .002),
      G2 = list(V = 1, nu = .002)
    )
  ), verbose=FALSE, thin=1, pr=TRUE)

# summary of the model
summary(m)

# examples of extracting standard deviations of
# different random effects on the linear predictor metric
```

```
# or after transformation to probabilities (only for ordinal)
stdranef(m, which = list(1), type = "lp")
stdranef(m, which = list(2), type = "lp")
stdranef(m, which = list(1, 2, c(1, 2)), type = "lp")
stdranef(m, type = "lp")

## error because no 3rd random effect
#stdranef(m, which = list(1, 2, 3), type = "lp")

stdranef(m, which = list("FSfamily", "plate"), type = "lp")

# mean standard deviations on the probability metric
# also the full distributions, if desired in the Data slot.
res <- stdranef(m, type = "response")
res$M # means
hist(res$Data$FSfamily[, 1]) # histogram

## End(Not run)
```

tidy.lqmm

Tidying methods for lqmm models (EXPERIMENTAL)

Description

These methods, suggested by David Luke Thiessen on [Stack Exchange](#), provide support for linear quantile mixed models. They have not been carefully tested - please check output carefully and report problems!

Usage

```
## S3 method for class 'lqmm'
tidy(x, conf.int = FALSE, conf.level = 0.95, ...)

## S3 method for class 'lqmm'
glance(x, ...)
```

Arguments

x	An object of class merMod, such as those from lmer, glmer, or nlmer
conf.int	whether to include a confidence interval
conf.level	confidence level for CI
...	Additional arguments (passed to confint.merMod for tidy; augment_columns for augment; ignored for glance)

tidy.MCMCglmm

*Tidying methods for MCMC (Stan, JAGS, etc.) fits***Description**

Tidying methods for MCMC (Stan, JAGS, etc.) fits

Usage

```
## S3 method for class 'MCMCglmm'
tidy(x, effects = c("fixed", "ran_pars"), scales = NULL, ...)

tidyMCMC(
  x,
  pars,
  robust = FALSE,
  conf.int = FALSE,
  conf.level = 0.95,
  conf.method = c("quantile", "HPDinterval"),
  drop.pars = c("lp__", "deviance"),
  rhat = FALSE,
  ess = FALSE,
  index = FALSE,
  ...
)

## S3 method for class 'rjags'
tidy(
  x,
  robust = FALSE,
  conf.int = FALSE,
  conf.level = 0.95,
  conf.method = "quantile",
  ...
)

## S3 method for class 'stanfit'
tidy(
  x,
  pars,
  robust = FALSE,
  conf.int = FALSE,
  conf.level = 0.95,
  conf.method = c("quantile", "HPDinterval"),
  drop.pars = c("lp__", "deviance"),
  rhat = FALSE,
  ess = FALSE,
```

```

    index = FALSE,
    ...
  )

## S3 method for class 'mcmc'
tidy(
  x,
  pars,
  robust = FALSE,
  conf.int = FALSE,
  conf.level = 0.95,
  conf.method = c("quantile", "HPDinterval"),
  drop.pars = c("lp__", "deviance"),
  rhat = FALSE,
  ess = FALSE,
  index = FALSE,
  ...
)

## S3 method for class 'mcmc.list'
tidy(
  x,
  pars,
  robust = FALSE,
  conf.int = FALSE,
  conf.level = 0.95,
  conf.method = c("quantile", "HPDinterval"),
  drop.pars = c("lp__", "deviance"),
  rhat = FALSE,
  ess = FALSE,
  index = FALSE,
  ...
)

```

Arguments

<code>x</code>	a model fit to be converted to a data frame
<code>effects</code>	which effects (fixed, random, etc.) to return
<code>scales</code>	scales on which to report results
<code>...</code>	mostly unused; for <code>tidy.MCMCglmm</code> , these represent options passed through to <code>tidy.mcmc</code> (e.g. <code>robust</code> , <code>conf.int</code> , <code>conf.method</code> , ...)
<code>pars</code>	(character) specification of which parameters to include
<code>robust</code>	use mean and standard deviation (if <code>FALSE</code>) or median and mean absolute deviation (if <code>TRUE</code>) to compute point estimates and uncertainty?
<code>conf.int</code>	(logical) include confidence interval?
<code>conf.level</code>	probability level for CI

conf.method	method for computing confidence intervals ("quantile" or "HPDinterval")
drop.pars	Parameters not to include in the output (such as log-probability information)
rhat, ess	(logical) include Rhat and/or effective sample size estimates?
index	Add index column, remove index from term. For example, term a[13] becomes term a and index 13.

Examples

```
if (require("MCMCglmm")) {
  ## original model
  ## Not run:
  mm0 <- MCMCglmm(Reaction ~ Days,
    random = ~Subject, data = sleepstudy,
    nitt=4000,
    pr = TRUE
  )

  ## End(Not run)
  ## load stored object
  load(system.file("extdata", "MCMCglmm_example.rda",
    package="broom.mixed"))

  tidy(mm0)
  tidy(mm1)
  tidy(mm2)
  tail(tidy(mm0, effects="ran_vals"))
}

# Using example from "RStan Getting Started"
# https://github.com/stan-dev/rstan/wiki/RStan-Getting-Started

model_file <- system.file("extdata", "8schools.stan", package = "broom.mixed")
schools_dat <- list(J = 8,
  y = c(28, 8, -3, 7, -1, 1, 18, 12),
  sigma = c(15, 10, 16, 11, 9, 11, 10, 18))

## original model
## Not run:
set.seed(2015)
rstan_example <- rstan::stan(file = model_file, data = schools_dat,
  iter = 1000, chains = 2, save_dso = FALSE)

## End(Not run)
if (require(rstan)) {
  ## load stored object
  rstan_example <- readRDS(system.file("extdata", "rstan_example.rds", package = "broom.mixed"))
  tidy(rstan_example)
  tidy(rstan_example, conf.int = TRUE, pars = "theta")
  td_mean <- tidy(rstan_example, conf.int = TRUE)
  td_median <- tidy(rstan_example, conf.int = TRUE, robust = TRUE)

  if (require(dplyr) && require(ggplot2)) {
    tds <- (dplyr::bind_rows(list(mean=td_mean, median=td_median), .id="method")
      %>% mutate(type=ifelse(grepl("^theta", term), "theta",
```

```

      ifelse(grepl("^eta",term),"eta",
             "other"))))
    )

    ggplot(tds, aes(estimate, term)) +
      geom_errorbarh(aes(xmin = conf.low, xmax = conf.high),height=0) +
      geom_point(aes(color = method))+
      facet_wrap(~type,scale="free",ncol=1)
  } ## require(dplyr,ggplot2)
} ## require(rstan)
if (require(R2jags)) {
  ## see help("jags",package="R2jags")
  ## and  example("jags",package="R2jags")
  ## for details
  ## load stored object
  R2jags_example <- readRDS(system.file("extdata", "R2jags_example.rds", package = "broom.mixed"))
  tidy(R2jags_example)
  tidy(R2jags_example, conf.int=TRUE, conf.method="quantile")
}

```

tidy.TMB

*Tidying methods for TMB models***Description**

Tidying methods for TMB models

Usage

```

## S3 method for class 'TMB'
tidy(
  x,
  effects = c("fixed", "random"),
  conf.int = FALSE,
  conf.level = 0.95,
  conf.method = c("wald", "uniroot", "profile"),
  ...
)

```

Arguments

<code>x</code>	An object of class TMB (you may need to use <code>class(obj) <- "TMB"</code> on your results from TMB)
<code>effects</code>	which effects should be returned?
<code>conf.int</code>	whether to include a confidence interval
<code>conf.level</code>	confidence level for CI
<code>conf.method</code>	method for computing confidence intervals
<code>...</code>	additional arguments passed to <code>confint</code> function (<code>tmbroot</code> , <code>tmbprofile</code>)

Examples

```
if (require("TMB")) {

  ## Not run:
  runExample("simple", thisR=TRUE)
  class(obj) <- "TMB"
  tidy(obj, conf.int=TRUE, conf.method="wald")

  ## End(Not run)
  ## Not run: tidy(obj, conf.int=TRUE, conf.method="uniroot")
  ## Not run: tidy(obj, conf.int=TRUE, conf.method="profile")
}
```

tidy.varFunc

Tidy variance structure for the nlme package.

Description

Returns a tibble with the following columns:

group type of varFunc, along with the right hand side of the formula in parentheses e.g. "varExp(age | Sex)".

term terms included in the formula of the variance model, specifically the names of the coefficients. If the value is fixed, it will be appended with " ; fixed".

estimate estimated coefficient

estimated This column is only included if some parameters are fixed. TRUE if the parameter is estimated and FALSE if the parameter is fixed.

Usage

```
## S3 method for class 'varFunc'
tidy(x, ...)
```

```
## S3 method for class 'varComb'
tidy(x, ...)
```

Arguments

x	An object of class varFunc, such as those used as the weights argument from the nlme package
...	Ignored

Value

If the varFunc is uninitialized or has no parameters, the function will return an empty tibble. Otherwise, it will return a tibble with names described in the details section.

Examples

```
## Not run:
if (require("nlme")) {
  ChickWeight_arbitrary_group <- datasets::ChickWeight
  ChickWeight_arbitrary_group$group_arb_n <-
    1 + (
      as.integer(ChickWeight_arbitrary_group$Chick) >
      median(as.integer(ChickWeight_arbitrary_group$Chick))
    )
  ChickWeight_arbitrary_group$group_arb <- c("low", "high")[ChickWeight_arbitrary_group$group_arb_n]

  fit_with_fixed <-
    lme(
      weight ~ Diet * Time,
      random = ~Time | Chick,
      data =ChickWeight_arbitrary_group,
      weights=varIdent(fixed=c("low"=5), form=~1|group_arb)
    )
  # Show all parameters
  tidy(fit_with_fixed)
  # Exclude fixed parameters
  tidy(fit_with_fixed) %>%
    filter(across(any_of("estimated"), ~.x))
}

## End(Not run)
```

unrowname

strip rownames from an object

Description

strip rownames from an object

Usage

```
unrowname(x)
```

Arguments

x a data frame

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