Package 'multilevelcoda'

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```
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Description Implement Bayesian multilevel modelling for compositional data.
     Compute multilevel compositional data and
     perform log-ratio transforms at between and within-person levels,
     fit Bayesian multilevel models for compositional predictors and outcomes,
     and run post-hoc analyses such as isotemporal substitution models.
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as.data.frame.complr Extract Compositional Data from complr object.

Description

Extract amounts and compositions in conventional formats as data.frames, matrices, or arrays.

Usage

```
## S3 method for class 'complr'
as.data.frame(x, row.names = NULL, optional = TRUE, ...)
## S3 method for class 'complr'
as.matrix(x, ...)
```

Arguments

```
    x An object of class complr.
    row.names, optional
    Unused and only added for consistency with the as.data.frame generic.
    generic argument, not in use.
```

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bayes_factor.brmcoda Bayes Factors from Marginal Likelihoods

Description

Compute Bayes factors from marginal likelihoods

Usage

```
## S3 method for class 'brmcoda'
bayes_factor(x1, x2, ...)
```

Arguments

х1 A brmcoda object.

х2 Another brmcoda object based on the same responses.

Other arguments passed to bayes_factor.brmsfit.

See Also

```
bayes_factor.brmsfit
```

brmcoda

Fit Bayesian generalised (non-)linear multilevel compositional model via full Bayesian inference

Description

Fit a brm model with multilevel ILR coordinates

Usage

```
brmcoda(complr, formula, ...)
```

Arguments

complr	A complr object containing data of composition, ILR coordinates, and other
	variables used in the model.
formula	A object of class formula, brmsformula: A symbolic description of the model

A object of class formula, brmsformula: A symbolic description of the model

to be fitted. Details of the model specification can be found in brmsformula.

Further arguments passed to brm.

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Value

A brmcoda with two elements

complr An object of class complr used in the brm model.

model An object of class brmsfit, which contains the posterior draws along with many

other useful information about the model.

Examples

```
if(requireNamespace("cmdstanr")){
 cilr <- complr(data = mcompd, sbp = sbp,</pre>
                 parts = c("TST", "WAKE", "MVPA", "LPA", "SB"), idvar = "ID")
 # inspects ILRs before passing to brmcoda
 names(cilr$between_logratio)
 names(cilr$within_logratio)
 names(cilr$logratio)
 # model with compositional predictor at between and within-person levels
 m1 <- brmcoda(complr = cilr,</pre>
                formula = Stress ~ bilr1 + bilr2 + bilr3 + bilr4 +
                                    wilr1 + wilr2 + wilr3 + wilr4 + (1 | ID),
                chain = 1, iter = 500,
                backend = "cmdstanr")
 # model with compositional outcome
 m2 <- brmcoda(complr = cilr,</pre>
                formula = mvbind(ilr1, ilr2, ilr3, ilr4) ~ Stress + Female + (1 | ID),
                chain = 1, iter = 500,
                backend = "cmdstanr")
 }
```

bsub

Between-person Simple Substitution

Description

This function is an alias of substitution to estimates the the difference in an outcome when compositional parts are substituted for specific unit(s) at *between* level using a single reference composition (e.g., compositional mean at sample level). It is recommended that users run substitution model using the substitution function.

Usage

```
bsub(
  object,
  delta,
  basesub,
```

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```
summary = TRUE,
ref = "grandmean",
level = "between",
weight = "equal",
aorg = TRUE,
scale = c("response", "linear"),
comparison = "one-to-one",
cores = NULL,
...
)
```

Arguments

object A fitted brmcoda object.

delta A integer, numeric value or vector indicating the amount of substituted change

between compositional parts.

basesub A base substitution. Can be a data.frame or data.table of the base possi-

ble substitution of compositional parts, which can be computed using function build.basesub. If "one-to-all", all possible one-to-remaining reallocations

are estimated. If NULL, all possible one-to-one reallocations are estimated.

summary A logical value to obtain summary statistics instead of the raw values. Default is

TRUE. Currently only support outputing raw values for model using grandmean

as reference composition.

ref Either a character value or vector or a dataset. Can be "grandmean" and/or

"clustermean", or a data.frame or data.table of user's specified reference grid consisting of combinations of covariates over which predictions are made. User's specified reference grid is only possible for simple substitution. Single

level models are default to "grandmean".

level A character string or vector. Should the estimate of multilevel models focus on

the "between" and/or "within" or "aggregate" variance? Single-level models

are default to "aggregate".

weight A character value specifying the weight to use in calculation of the reference

composition. If "equal", give equal weight to units (e.g., individuals). If "proportional", weights in proportion to the frequencies of units being averaged (e.g., observations across individuals). Default to "equal" for ref =

"grandmean" and "proportional" for ref = "clustermean".

aorg A logical value to obtain (a) verage prediction (o) ver the (r) eference (g) rid. Should

the estimate at each level of the reference grid (FALSE) or their average (TRUE) be returned? Default is TRUE. Only applicable for model with covariates in addition

to the isometric log-ratio coordinates (i.e., adjusted model).

scale Either "response" or "linear". If "response", results are returned on the

scale of the response variable. If "linear", results are returned on the scale of the linear predictor term, that is without applying the inverse link function or

other transformations.

comparison internally used only.

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Number of cores to use when executing the chains in parallel, we recommend

setting the mc.cores option to be as many processors as the hardware and RAM allow (up to the number of compositional parts). For non-Windows OS in non-

interactive R sessions, forking is used instead of PSOCK clusters.

... currently ignored.

Value

A list containing the results of multilevel compositional substitution model. The first six lists contain the results of the substitution estimation for a compositional part.

Mean Posterior means.

CI_low and CI_high

95% credible intervals.

Delta Amount substituted across compositional parts.

From Compositional part that is substituted from.

To Compositional parts that is substituted to.

Level Where changes in composition takes place.

Reference Either grandmean, clustermean, or users.

See Also

substitution

Examples

bsubmargins

Between-person Average Substitution

Description

This function is an alias of substitution to estimates the difference in an outcome when compositional parts are substituted for specific unit(s) at *between* level using cluster mean (e.g., compositional mean at individual level) as reference composition. It is recommended that users run substitution model using the substitution function.

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Usage

```
bsubmargins(
  object,
  delta,
  basesub,
  summary = TRUE,
  ref = "clustermean",
  level = "between",
  weight = "proportional",
  scale = c("response", "linear"),
  comparison = "one-to-one",
  cores = NULL,
  ...
)
```

Arguments

object A fitted brmcoda object.

delta A integer, numeric value or vector indicating the amount of substituted change

between compositional parts.

basesub A base substitution. Can be a data.frame or data.table of the base possi-

ble substitution of compositional parts, which can be computed using function build.basesub. If "one-to-all", all possible one-to-remaining reallocations

are estimated. If NULL, all possible one-to-one reallocations are estimated.

summary A logical value to obtain summary statistics instead of the raw values. Default is

TRUE. Currently only support outputing raw values for model using grandmean

as reference composition.

ref Either a character value or vector or a dataset. Can be "grandmean" and/or

"clustermean", or a data.frame or data.table of user's specified reference grid consisting of combinations of covariates over which predictions are made. User's specified reference grid is only possible for simple substitution. Single

level models are default to "grandmean".

level A character string or vector. Should the estimate of multilevel models focus on

the "between" and/or "within" or "aggregate" variance? Single-level models

are default to "aggregate".

weight A character value specifying the weight to use in calculation of the reference

composition. If "equal", give equal weight to units (e.g., individuals). If "proportional", weights in proportion to the frequencies of units being averaged (e.g., observations across individuals). Default to "equal" for ref =

"grandmean" and "proportional" for ref = "clustermean".

scale Either "response" or "linear". If "response", results are returned on the

scale of the response variable. If "linear", results are returned on the scale of the linear predictor term, that is without applying the inverse link function or

other transformations.

comparison internally used only.

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cores Number of cores to use when executing the chains in parallel, we recommend

setting the mc.cores option to be as many processors as the hardware and RAM allow (up to the number of compositional parts). For non-Windows OS in non-

interactive R sessions, forking is used instead of PSOCK clusters.

... currently ignored.

Value

A list containing the results of multilevel compositional substitution model. The first six lists contain the results of the substitution estimation for a compositional part.

Mean Posterior means.

CI_low and CI_high

95% credible intervals.

Delta Amount substituted across compositional parts.

From Compositional part that is substituted from.

To Compositional parts that is substituted to.

Level Where changes in composition takes place.

Reference Either grandmean, clustermean, or users.

See Also

substitution

Examples

build.basesub

Build Base Pairwise Substitution

Description

Make a data set of all possible pairwise substitution of a composition which can be used as the base for substitution models.

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Usage

```
build.basesub(parts, comparison = NULL)
```

Arguments

parts A character vector specifying the names of compositional variables to be used. comparison Either "one-to-one" or "one-to-all". Default is "one-to-one".

Value

A data table of all possible pairwise substitution.

Examples

```
ps1 <- build.basesub(parts = c("TST", "WAKE", "MVPA", "LPA", "SB"))
print(ps1)

ps2 <- build.basesub(c("WAKE", "MVPA", "LPA", "SB"), comparison = "one-to-all")
print(ps2)</pre>
```

build.rg

Reference Grid for substitution model.

Description

Build a dataset for fitted.brmcoda used in substitution model

Usage

```
build.rg(object, ref, level, weight, fill = FALSE)
```

Arguments

object	A fitted b	rmcoda	object.

ref Either a character value or vector or a dataset. Can be "grandmean" and/or

"clustermean", or a data.frame or data.table of user's specified reference grid consisting of combinations of covariates over which predictions are made. User's specified reference grid is only possible for simple substitution. Single

level models are default to "grandmean".

level A character string or vector. Should the estimate of multilevel models focus on

the "between" and/or "within" or "aggregate" variance? Single-level models

are default to "aggregate".

weight A character value specifying the weight to use in calculation of the reference

composition. If "equal", give equal weight to units (e.g., individuals). If "proportional", weights in proportion to the frequencies of units being averaged (e.g., observations across individuals). Default to "equal" for ref =

"grandmean" and "proportional" for ref = "clustermean".

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fill

Logical value only relevant when ref is an user's specified reference grid in which information about some, but not all covariates is provided (e.g., models including age and sex as covariate but only age was provided in the reference grid). If TRUE, the unspecified covariates are filled with the default reference grid. If FALSE, users will be asked to provide a full reference grid. Currently only support the default to FALSE.

Value

A reference grid consisting of a combination of covariates in brmcoda

build.sbp

Build Sequential Binary Partition

Description

Build a default sequential binary partition for complr object. The default sequential binary partition is a pivot balance that allows the effect of this first balance coordinate to be interpreted as the change in the prediction for the dependent variable when that given part increases while all remaining parts decrease by a common proportion.

Usage

```
build.sbp(parts)
```

Arguments

parts

A character vector specifying the names of compositional variables to be used.

Value

A matrix sequential binary partition.

```
sbp1 <- build.sbp(parts = c("TST", "WAKE", "MVPA", "LPA", "SB"))
print(sbp1)

sbp2 <- build.sbp(c("WAKE", "MVPA", "LPA", "SB"))
print(sbp2)</pre>
```

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coef.brmcoda

Model Coefficients

Description

Extract model coefficients, which are the sum of population-level effects and corresponding group-level effects of the brmsfit object in a brmcoda object.

Usage

```
## S3 method for class 'brmcoda'
coef(object, ...)
```

Arguments

object An object of class brmcoda.
... Further arguments passed to coef.brmsfit.

Value

A list of 3D arrays (one per grouping factor). If summary is TRUE, the 1st dimension contains the factor levels, the 2nd dimension contains the summary statistics (see posterior_summary), and the 3rd dimension contains the group-level effects. If summary is FALSE, the 1st dimension contains the posterior draws, the 2nd dimension contains the factor levels, and the 3rd dimension contains the group-level effects.

See Also

```
coef.brmsfit
```

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compilr

Indices from a (dataset of) Multilevel Composition(s) (deprecated.)

Description

Indices from a (dataset of) Multilevel Composition(s) (deprecated.)

Usage

```
compilr(...)
```

Arguments

... arguments passed to complr.

Value

A complr object with at least the following elements.

comp A vector of class acomp representing one closed composition or a matrix of class

acomp representing multiple closed compositions each in one row.

between_comp A vector of class acomp representing one closed between-person composition or

a matrix of class acomp representing multiple closed between-person composi-

tions each in one row.

within_comp A vector of class acomp representing one closed within-person composition or a

matrix of class acomp representing multiple closed within-person compositions

each in one row.

logratio Log ratio transform of composition.

between_logratio

Log ratio transform of between-person composition.

within_logratio

Log ratio transform of within-person composition.

data The user's dataset or imputed dataset if the input data contains zeros.

transform Type of transform applied on compositional data.

parts Names of compositional variables.

idvar Name of the variable containing IDs.

total Total amount to which the compositions is closed.

See Also

complr

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complr	Indices from a (dataset of) Multilevel Composition(s)	

Description

Compute sets of compositions and log ratio transformation for multilevel compositional data

Usage

```
complr(data, parts, sbp = NULL, total = 1, idvar = NULL, transform = "ilr")
```

Arguments

data	A data.frame or data.table containing data of all variables used in the analysis. It must include a composition and a ID variable. Required.
parts	A character vector specifying the names of compositional variables to be used.
sbp	A signary matrix indicating sequential binary partition.
total	A numeric value of the total amount to which the compositions should be closed. Default is 1.
idvar	Only for multilevel data, a character string specifying the name of the variable containing IDs.
transform	A character value naming a log ratio transformation to be applied on compositional data. Can be either "ilr" (isometric logratio), "alr" (additive logratio), or "clr" (centered logratio). Default is "ilr".

Details

The *ilr*-transform maps the D-part compositional data from the simplex into non-overlapping subgroups in the (D-1)-dimension Euclidean space isometrically by using an orthonormal basis, thereby preserving the compositional properties and yielding a full-rank covariance matrix. *ilr* transformation should be preferred. However, the *alr* and *clr* are alternatives. The *alr*-transform maps a D-part composition in the Aitchison-simplex non-isometrically to a (D-1)-dimension Euclidian vectors, commonly treating the last part as the common denominator of the others. *alr* transformation does not rely on distance which breaks the constraint of compositional data. *clr*-transform maps a D-part composition in the Aitchison-simplex isometrically to a D-dimensional Euclidian vector subspace. *clr* transformation is not injetive, resulting in singular covariance matrices.

Value

A complr object with at least the following elements.

comp	A vector of class acomp representing one closed composition or a matrix of class
	acomp representing multiple closed compositions each in one row.
between_comp	A vector of class acomp representing one closed between-person composition or
	a matrix of class acomp representing multiple closed between-person composi-
	tions each in one row

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 $\label{eq:within_comp} \textbf{A vector of class acomp representing one closed within-person composition or a}$

matrix of class acomp representing multiple closed within-person compositions

each in one row.

logratio Log ratio transform of composition.

between_logratio

Log ratio transform of between-person composition.

within_logratio

Log ratio transform of within-person composition.

data The user's dataset or imputed dataset if the input data contains zeros.

transform Type of transform applied on compositional data.

parts Names of compositional variables.

idvar Name of the variable containing IDs.

total Total amount to which the compositions is closed.

Examples

```
cilr <- complr(data = mcompd,</pre>
                 parts = c("TST", "WAKE", "MVPA", "LPA", "SB"),
                 idvar = "ID", total = 1440)
str(cilr)
calr <- complr(data = mcompd, sbp = sbp,</pre>
                 parts = c("TST", "WAKE", "MVPA", "LPA", "SB"),
                 idvar = "ID",
                 transform = "alr")
str(calr)
cclr <- complr(data = mcompd, sbp = sbp,</pre>
                 parts = c("TST", "WAKE", "MVPA", "LPA", "SB"),
                 idvar = "ID",
                  transform = "clr")
str(cclr)
cilr_wide <- complr(data = mcompd[!duplicated(ID)], sbp = sbp,</pre>
                 parts = c("TST", "WAKE", "MVPA", "LPA", "SB"))
str(cilr_wide)
```

create_substitution Constructor function for substitution class.

Description

Constructor function for substitution class.

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Usage

```
create_substitution(
  between_simple_sub,
  within_simple_sub,
  simple_sub,
  between_avg_sub,
  within_avg_sub,
  avg_sub,
  delta,
  ref,
  level,
  weight,
  parts,
  aorg
)
```

Arguments

between_simple_sub

A list of results from bsub or NULL

within_simple_sub

A list of results from wsub or NULL

simple_sub A list of results from sub or NULL

between_avg_sub

A list of results from bsubmargins or NULL

within_avg_sub A list of results from wsubmargins or NULL

avg_sub A list of results from submargins or NULL

delta A numeric vector of the amount of substitution ref A character value specifying the reference grid

level A character value specifying the level of substitution

weight The weight to use in calculation of the reference composition

parts The parts of the composition

aorg A logical value specifying whether to summarize the results (average over the

reference grid)

Value

An object of class substitution

See Also

substitution

```
diagnostic-quantities-brmcoda
```

Extract Diagnostic Quantities from brmsfit Models in brmcoda

Description

Extract Diagnostic Quantities from brmsfit Models in brmcoda

Usage

```
## S3 method for class 'brmcoda'
log_posterior(object, ...)
## S3 method for class 'brmcoda'
nuts_params(object, ...)
## S3 method for class 'brmcoda'
rhat(x, ...)
## S3 method for class 'brmcoda'
neff_ratio(object, ...)
```

Arguments

... Arguments passed to individual methods (if applicable).

x, object A brmcoda object or another R object for which the methods are defined.

Value

The exact form of the output depends on the method.

See Also

```
log_posterior.brmsfit
nuts_params.brmsfit
rhat.brmsfit
neff_ratio.brmsfit
```

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draws-index-brmcoda

Index brmcoda objects

Description

Index brmcoda objects

Usage

```
## S3 method for class 'brmcoda'
variables(x, ...)
## S3 method for class 'brmcoda'
nvariables(x, ...)
## S3 method for class 'brmcoda'
niterations(x)
## S3 method for class 'brmcoda'
nchains(x)
## S3 method for class 'brmcoda'
ndraws(x)
```

Arguments

x An object of class brmcoda.

... Arguments passed to individual methods.

See Also

```
variables.brmsfit
nvariables.brmsfit
niterations.brmsfit
nchains.brmsfit
ndraws.brmsfit
```

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fitted.brmcoda

Expected Values of the Posterior Predictive Distribution

Description

Compute posterior draws of the expected value of the posterior predictive distribution of a brmsfit model in the brmcoda object. Can be performed for the data used to fit the model (posterior predictive checks) or for new data. By definition, these predictions have smaller variance than the posterior predictions performed by the predict.brmcoda method. This is because only the uncertainty in the expected value of the posterior predictive distribution is incorporated in the draws computed by fitted while the residual error is ignored there. However, the estimated means of both methods averaged across draws should be very similar.

Usage

```
## S3 method for class 'brmcoda'
fitted(object, scale = c("linear", "response"), summary = TRUE, ...)
```

Arguments

object An object of class brmcoda. scale

Specifically for models with compositional responses, either "response" or "linear". If "linear", results are returned on the log-ratio scale. If "response",

results are returned on the compositional scale of the response variable.

Should summary statistics be returned instead of the raw values? Default is summary

TRUE.

Further arguments passed to fitted.brmsfit that control additional aspects of

prediction.

Value

An array of predicted *mean* response values. If summary = FALSE the output resembles those of posterior_epred.brmsfit.

If summary = TRUE the output depends on the family: For categorical and ordinal families, the output is an N x E x C array, where N is the number of observations, E is the number of summary statistics, and C is the number of categories. For all other families, the output is an N x E matrix. The number of summary statistics E is equal to 2 + length(probs): The Estimate column contains point estimates (either mean or median depending on argument robust), while the Est. Error column contains uncertainty estimates (either standard deviation or median absolute deviation depending on argument robust). The remaining columns starting with Q contain quantile estimates as specified via argument probs.

In multivariate models, an additional dimension is added to the output which indexes along the different response variables.

See Also

fitted.brmsfit

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Examples

```
## fit a model
if(requireNamespace("cmdstanr")){
 ## compute composition and ilr coordinates
 cilr <- complr(data = mcompd, sbp = sbp,</pre>
                 parts = c("TST", "WAKE", "MVPA", "LPA", "SB"),
                 idvar = "ID", total = 1440)
 ## fit a model
 m1 <- brmcoda(complr = cilr,</pre>
                formula = Stress ~ bilr1 + bilr2 + bilr3 + bilr4 +
                  wilr1 + wilr2 + wilr3 + wilr4 + (1 | ID),
                chain = 1, iter = 500,
                backend = "cmdstanr")
 ## compute expected predictions
 epred <- fitted(m1)</pre>
 head(epred)
 ## fit a model with compositional outcome
 m2 <- brmcoda(complr = cilr,</pre>
                formula = mvbind(ilr1, ilr2, ilr3, ilr4) ~ Stress + Female + (1 | ID),
                chain = 1, iter = 500,
                backend = "cmdstanr")
 ## expected predictions on compositional scale
 epredcomp <- fitted(m2, scale = "response")</pre>
 head(epredcomp)
}
```

fixef.brmcoda

Population-Level Estimates

Description

Extract the population-level ('fixed') effects from the brmsfit object in a brmcoda object.

Usage

```
## S3 method for class 'brmcoda'
fixef(object, ...)
```

Arguments

```
object An object of class brmcoda.
... Further arguments passed to fixef.brmsfit.
```

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Value

If summary is TRUE, a matrix returned by posterior_summary for the population-level effects. If summary is FALSE, a matrix with one row per posterior draw and one column per population-level effect.

See Also

```
fixef.brmsfit
```

Examples

get-substitution

Helper functions used only internally to estimate substitution model

Description

Helper functions used only internally to estimate substitution model

is.brmcoda

Checks if argument is a brmcoda object

Description

Checks if argument is a brmcoda object

Usage

```
is.brmcoda(x)
```

Arguments

An object of class brmcoda.

is.complr

Checks if argument is a complr object

Description

Checks if argument is a complr object

Usage

```
is.complr(x)
```

Arguments

Χ

An object of class complr.

is.substitution

Checks if argument is a substitution object

Description

Checks if argument is a substitution object

Usage

```
is.substitution(x)
```

Arguments

Х

An object of class substitution.

launch_shinystan.brmcoda

Interface to shinystan

Description

Provide an interface to shinystan for models fitted with brms

Usage

```
## S3 method for class 'brmcoda'
launch_shinystan(object, ...)
```

loo.brmcoda 23

Arguments

object A fitted model object of class brmcoda.

... Optional arguments to pass to launch_shinystan or runApp.

Value

An S4 shinystan object

See Also

launch_shinystan

loo.brmcoda

Efficient approximate leave-one-out cross-validation (LOO)

Description

Perform approximate leave-one-out cross-validation based on the posterior likelihood using the **loo** package. For more details see **loo**.

Usage

```
## S3 method for class 'brmcoda' loo(x, ...)
```

Arguments

x A brmcoda object.

... More brmsfit objects or further arguments passed to the underlying post-processing functions. In particular, see prepare_predictions for further supported arguments.

Value

If just one object is provided, an object of class loo. If multiple objects are provided, an object of class loolist.

See Also

loo.brmsfit

24 mcmc_plot.brmcoda

mcmc_plot.brmcoda

MCMC Plots Implemented in bayesplot

Description

Call MCMC plotting functions implemented in the bayesplot package.

Usage

```
## S3 method for class 'brmcoda'
mcmc_plot(object, ...)
```

Arguments

object A brmcoda class object.
... Further arguments passed to mcmc_plot.brmsfit.

Value

A ggplot object that can be further customized using the ggplot2 package.

See Also

```
mcmc_plot.brmsfit
```

mcompd 25

mcompd

Multilevel Compositional Data

Description

A simulated dataset containing multiple days of compositional data.

Usage

mcompd

Format

A data table containing 10 variables.

ID A unique identifier for each individual

Time Recurrence time of repeated measures by individual

Stress Self report stress measures on a 0 to 10 scale — repeated measure

TST Total Sleep Time (minutes) — repeated measure

WAKE Wake time while in bed, trying to sleep (minutes) — repeated measure

MVPA Moderate to Vigorous Physical Activity (minutes) — repeated measure

LPA Light Physical Activity (minutes) — repeated measure

SB Sedentary Behavior (minutes) — repeated measure

Age Age in years — baseline measure only

Female Binary: whether participants identified as female (1) or not (0) — baseline measure only

mean.complr

Mean amounts and mean compositions presented in a complr object.

Description

Mean amounts and mean compositions presented in a complr object.

Usage

```
## S3 method for class 'complr'
mean(x, weight = c("equal", "proportional"), ...)
```

26 multilevelcoda_sim

Arguments

An object of class complr.

A character value specifying the weight to use in calculation of the reference composition. If "equal", give equal weight to units (e.g., individuals). If "proportional", weights in proportion to the frequencies of units being averaged (e.g., observations across individuals) Default is equal.

... generic argument, not in use.

Examples

 ${\tt model.frame.brmcoda}$

Extracting the Model Frame from a Formula or Fit from brmcoda object

Description

Extracting the Model Frame from a Formula or Fit from brmcoda object

Usage

```
## S3 method for class 'brmcoda'
model.frame(formula, ...)
```

Arguments

formula A brmcoda object.

Further arguments to be passed to methods.

multilevelcoda_sim

multilevelcoda Simulation Study Results

Description

Provide the full results for a simulation study testing the performance of multilevelcoda

Usage

```
multilevelcoda_sim()
```

Value

An S4 shiny object

nobs.brmcoda 27

nobs.brmcoda

Extract Number of Observations from brmcoda object

Description

Extract Number of Observations from brmcoda object

Usage

```
## S3 method for class 'brmcoda'
nobs(object, ...)
```

Arguments

object A brmcoda object.

... Further arguments to be passed to methods.

pairs.brmcoda

Create a matrix of output plots from a brmcoda's brmsfit object

Description

A pairs method that is customized for MCMC output.

Usage

```
## S3 method for class 'brmcoda'
pairs(x, ...)
```

Arguments

x A brmcoda class object.

Further arguments passed to pairs.brmsfit.

See Also

```
pairs.brmsfit
```

28 pivot_coord

Examples

pivot_coord

Estimate pivot balance coordinates

Description

Estimate pivot balance coordinates

Usage

```
pivot_coord(object, summary = TRUE, method = c("rotate", "refit"), ...)
```

Arguments

object An object of class brmcoda.

Should summary statistics be returned instead of the raw values? Default is TRUE.

method A character string. Should the pivot balance coordinates be estimated by "rotate" the sequential binary partition using the same brmcoda object or "refit" the brmcoda object? Default is "rotate".

... currently ignored.

Value

A list of brmcoda for each pivot balance coordinate.

pivot_coord_refit 29

pivot_coord_refit

Estimate pivot balance coordinates by refitting model.

Description

Estimate pivot balance coordinates by refitting model.

Usage

```
pivot_coord_refit(object, ...)
```

Arguments

object An object of class brmcoda.
... Further arguments passed to brm.

Value

A list of brmcoda for each pivot balance coordinate.

30 pivot_coord_rotate

Description

Estimate pivot balance coordinates by rotating sequential binary partition.

Usage

```
pivot_coord_rotate(object, summary = TRUE, ...)
```

Arguments

object An object of class brmcoda.

summary Should summary statistics be returned instead of the raw values? Default is TRUE.

... currently ignored.

Value

A list of brmcoda for each pivot balance coordinate.

plot.brmcoda 31

plot.brmcoda

Trace and Density Plots for MCMC Draws plot

Description

Make a plot of brmcoda model results.

Usage

```
## S3 method for class 'brmcoda' plot(x, ...)
```

Arguments

x A brmcoda class object.

... Further arguments passed to plot.brmsfit.

Value

An invisible list of gtable objects.

See Also

```
plot.brmsfit
```

32 pp_check.brmcoda

plot.substitution Suit

Substitution Plot

Description

Make a plot of substitution model results.

Usage

```
## S3 method for class 'substitution'
plot(x, to, ref, level, ...)
```

Arguments

X	A substitution class object.
to	A character value or vector specifying the names of the compositional parts that were reallocated to in the model.
ref	A character value of (("grandmean" or "clustermean" or "users"),
level	A character value of ("between", "within"), or "aggregate").

... Further components to the plot, followed by a plus sign (+).

Value

A ggplot graph object showing the estimated difference in outcome when each pair of compositional variables are substituted for a specific time.

pp_check.brmcoda

Posterior Predictive Checks for brmcoda Objects

Description

Perform posterior predictive checks with the help of the **bayesplot** package.

Usage

```
## S3 method for class 'brmcoda'
pp_check(object, ...)
```

Arguments

object An object of class brmcoda.

... Further arguments passed to predict.brmsfit as well as to the PPC function

specified in type.

See Also

```
pp_check.brmsfit
```

predict.brmcoda 33

predict.brmcoda Draws from the Posterior Predictive Distribution	
--	--

Description

Compute posterior draws of the posterior predictive distribution of a brmsfit model in the brmcoda object. Can be performed for the data used to fit the model (posterior predictive checks) or for new data. By definition, these draws have higher variance than draws of the expected value of the posterior predictive distribution computed by fitted.brmcoda. This is because the residual error is incorporated in posterior_predict. However, the estimated means of both methods averaged across draws should be very similar.

Usage

```
## S3 method for class 'brmcoda'
predict(object, scale = c("linear", "response"), summary = TRUE, ...)
```

Arguments

object	An object of class brmcoda.
scale	Specifically for models with compositional responses, either "response" or "linear". If "linear", results are returned on the log-ratio scale. If "response", results are returned on the compositional scale of the response variable.
summary	Should summary statistics be returned instead of the raw values? Default is TRUE.
	Further arguments passed to predict.brmsfit that control additional aspects of prediction.

Value

An array of predicted response values. If summary = FALSE the output resembles those of posterior_predict.brmsfit.

If summary = TRUE the output depends on the family: For categorical and ordinal families, the output is an N x C matrix, where N is the number of observations, C is the number of categories, and the values are predicted category probabilities. For all other families, the output is a N x E matrix where E = 2 + length(probs) is the number of summary statistics: The Estimate column contains point estimates (either mean or median depending on argument robust), while the Est. Error column contains uncertainty estimates (either standard deviation or median absolute deviation depending on argument robust). The remaining columns starting with Q contain quantile estimates as specified via argument probs.

See Also

```
predict.brmsfit
```

34 print.brmcoda

Examples

```
if(requireNamespace("cmdstanr")){
 ## fit a model
 cilr <- complr(data = mcompd, sbp = sbp,</pre>
                 parts = c("TST", "WAKE", "MVPA", "LPA", "SB"),
                 idvar = "ID", total = 1440)
 m1 <- brmcoda(complr = cilr,</pre>
                formula = Stress ~ bilr1 + bilr2 + bilr3 + bilr4 +
                  wilr1 + wilr2 + wilr3 + wilr4 + (1 | ID),
                 chain = 1, iter = 500,
                backend = "cmdstanr")
 ## predicted responses
 pred <- predict(m1)</pre>
 head(pred)
 ## fit a model with compositional outcome
 m2 <- brmcoda(complr = cilr,</pre>
                 formula = mvbind(ilr1, ilr2, ilr3, ilr4) ~ Stress + Female + (1 | ID),
                chain = 1, iter = 500,
                backend = "cmdstanr")
 ## predicted responses on compositional scale
 predcomp <- predict(m2, scale = "linear")</pre>
 head(predcomp)
}
```

print.brmcoda

Print a Summary for a fitted brmsfit model in a brmcoda object

Description

Print a Summary for a fitted brmsfit model in a brmcoda object

Usage

```
## S3 method for class 'brmcoda'
print(x, ...)
```

Arguments

x An object of class brmcoda.

... Other arguments passed to summary.brmcoda.

See Also

summary.brmcoda

print.complr 35

Examples

print.complr

Print a Summary for a complr object

Description

Print a Summary for a complr object

Usage

```
## S3 method for class 'complr'
print(x, ...)
```

Arguments

- x An object of class complr.
- ... Other arguments passed to summary.complr.

See Also

```
summary.complr
```

print.substitution

Print a Summary for a substitution *object*

Description

Print a Summary for a substitution object

Usage

```
## S3 method for class 'substitution' print(x, ...)
```

Arguments

x A substitution object.

... Additional arguments to be passed to to method summary of substitution.

See Also

```
summary.substitution
```

Examples

print.summary.complr Summary for a complr object

Description

Summary for a complr object

Usage

```
## S3 method for class 'summary.complr'
print(x, ...)
```

prior_summary.brmcoda

Arguments

```
x An object of class summary.complr.... Other arguments passed to summary.complr.
```

See Also

```
summary.complr
```

Examples

prior_summary.brmcoda Extract Priors of a brmsfit from a brmcoda object

Description

Compute Bayes factors from marginal likelihoods

Usage

```
## S3 method for class 'brmcoda'
prior_summary(object, ...)
```

Arguments

object An object of class brmcoda.

. . . Further arguments passed to or from other methods.

See Also

```
prior_summary.brmsfit
```

38 ranef.brmcoda

psub

Possible Pairwise Substitutions

Description

A dataset containing possible pairwise substitutions.

Usage

psub

Format

A data table containing 5 variables.

TST first compositional variable

WAKE second compositional variable

MVPA third compositional variable

LPA fourth compositional variable

SB fifth compositional variable

ranef.brmcoda

Group-Level Estimates

Description

Extract the group-level ('random') effects of each level of the brmsfit object in a brmcoda object.

Usage

```
## S3 method for class 'brmcoda'
ranef(object, ...)
```

Arguments

object An object of class brmcoda.

... Further arguments passed to ranef.brmsfit.

Value

A list of 3D arrays (one per grouping factor). If summary is TRUE, the 1st dimension contains the factor levels, the 2nd dimension contains the summary statistics (see posterior_summary), and the 3rd dimension contains the group-level effects. If summary is FALSE, the 1st dimension contains the posterior draws, the 2nd dimension contains the factor levels, and the 3rd dimension contains the group-level effects.

residuals.brmcoda 39

See Also

```
ranef.brmsfit
```

Examples

residuals.brmcoda

Posterior Draws of Residuals/Predictive Errors

Description

Compute posterior draws of residuals/predictive errors

Usage

```
## S3 method for class 'brmcoda'
residuals(object, ...)
```

Arguments

object An object of class brmcoda.

... Further arguments passed to residuals.brmsfit.

Value

An array of predictive error/residual draws. If summary = FALSE the output resembles those of $predictive_error.brmsfit$. If summary = TRUE the output is an N x E matrix, where N is the number of observations and E denotes the summary statistics computed from the draws.

See Also

```
residuals.brmsfit
```

40 sbp

Examples

sbp

Sequential Binary Partition

Description

A matrix of sequential binary partition.

Usage

sbp

Format

A matrix with 5 columns and 4 rows.

TST first compositional variable

WAKE second compositional variable

MVPA third compositional variable

LPA fourth compositional variable

SB fifth compositional variable

sim 41

sim

multilevelcoda Simulation Study results

Description

A list of 4 components

Usage

sim

Format

A list with 5 columns and 4 rows.

brmcoda_tab Simulation results for brmcoda() for tablessub_tab Simulation results for substitution() for tablesbrmcoda_plot Simulation results for brmcoda() for graphssub_plot Simulation results for substitution() for graphs

sub

Simple Substitution

Description

This function is an alias of substitution to estimates the difference in an outcome when compositional parts are substituted for specific unit(s) using a aggregate reference composition (e.g., compositional mean at sample level, not separated by between- and within effects). It is recommended that users run substitution model using the substitution function.

```
sub(
  object,
  delta,
  basesub,
  ref = "grandmean",
  level = "aggregate",
  weight = "equal",
  aorg = TRUE,
  summary = TRUE,
  scale = c("response", "linear"),
  comparison = "one-to-one",
  cores = NULL,
  ...
)
```

42 sub

Arguments

object A fitted brmcoda object.

delta A integer, numeric value or vector indicating the amount of substituted change

between compositional parts.

basesub A base substitution. Can be a data frame or data table of the base possi-

> ble substitution of compositional parts, which can be computed using function build.basesub. If "one-to-all", all possible one-to-remaining reallocations

are estimated. If NULL, all possible one-to-one reallocations are estimated.

ref Either a character value or vector or a dataset. Can be "grandmean" and/or

> "clustermean", or a data.frame or data.table of user's specified reference grid consisting of combinations of covariates over which predictions are made. User's specified reference grid is only possible for simple substitution. Single

level models are default to "grandmean".

level A character string or vector. Should the estimate of multilevel models focus on

the "between" and/or "within" or "aggregate" variance? Single-level models

are default to "aggregate".

A character value specifying the weight to use in calculation of the reference weight

> composition. If "equal", give equal weight to units (e.g., individuals). If "proportional", weights in proportion to the frequencies of units being averaged (e.g., observations across individuals). Default to "equal" for ref =

"grandmean" and "proportional" for ref = "clustermean".

A logical value to obtain (a) verage prediction (o) ver the (r) eference (g) rid. Should aorg

> the estimate at each level of the reference grid (FALSE) or their average (TRUE) be returned? Default is TRUE. Only applicable for model with covariates in addition

to the isometric log-ratio coordinates (i.e., adjusted model).

A logical value to obtain summary statistics instead of the raw values. Default is summary

TRUE. Currently only support outputing raw values for model using grandmean

as reference composition.

Either "response" or "linear". If "response", results are returned on the scale

> scale of the response variable. If "linear", results are returned on the scale of the linear predictor term, that is without applying the inverse link function or

other transformations.

comparison internally used only.

Number of cores to use when executing the chains in parallel, we recommend cores

> setting the mc. cores option to be as many processors as the hardware and RAM allow (up to the number of compositional parts). For non-Windows OS in non-

interactive R sessions, forking is used instead of PSOCK clusters.

currently ignored.

Value

A list containing the results of multilevel compositional substitution model. The first six lists contain the results of the substitution estimation for a compositional part.

Posterior means. Mean

submargins 43

```
CI_low and CI_high
95% credible intervals.

Delta Amount substituted across compositional parts.

From Compositional part that is substituted from.

To Compositional parts that is substituted to.

Level Level where changes in composition takes place.

Reference Either grandmean, clustermean, or users.
```

See Also

substitution

Examples

submargins

Average Substitution

Description

This function is an alias of substitution to estimates the difference in an outcome when compositional parts are substituted for specific unit(s) using cluster mean (e.g., compositional mean at individual level) as reference composition. It is recommended that users run substitution model using the substitution function.

```
submargins(
  object,
  delta,
  basesub,
  summary = TRUE,
  ref = "clustermean",
  level = "aggregate",
```

44 submargins

```
weight = "proportional",
  scale = c("response", "linear"),
  comparison = "one-to-one",
  cores = NULL,
   ...
)
```

Arguments

object A fitted brmcoda object.

delta A integer, numeric value or vector indicating the amount of substituted change

between compositional parts.

basesub A base substitution. Can be a data.frame or data.table of the base possi-

ble substitution of compositional parts, which can be computed using function build.basesub. If "one-to-all", all possible one-to-remaining reallocations are estimated. If NULL, all possible one-to-one reallocations are estimated.

summary A logical value to obtain summary statistics instead of the raw values. Default is

TRUE. Currently only support outputing raw values for model using grandmean

as reference composition.

ref Either a character value or vector or a dataset. Can be "grandmean" and/or

"clustermean", or a data.frame or data.table of user's specified reference grid consisting of combinations of covariates over which predictions are made. User's specified reference grid is only possible for simple substitution. Single

level models are default to "grandmean".

level A character string or vector. Should the estimate of multilevel models focus on

the "between" and/or "within" or "aggregate" variance? Single-level models

are default to "aggregate".

weight A character value specifying the weight to use in calculation of the reference

composition. If "equal", give equal weight to units (e.g., individuals). If "proportional", weights in proportion to the frequencies of units being averaged (e.g., observations across individuals). Default to "equal" for ref =

"grandmean" and "proportional" for ref = "clustermean".

scale Either "response" or "linear". If "response", results are returned on the

scale of the response variable. If "linear", results are returned on the scale of the linear predictor term, that is without applying the inverse link function or

other transformations.

comparison internally used only.

cores Number of cores to use when executing the chains in parallel, we recommend

setting the mc.cores option to be as many processors as the hardware and RAM allow (up to the number of compositional parts). For non-Windows OS in non-

interactive R sessions, forking is used instead of PSOCK clusters.

... currently ignored.

Value

A list containing the results of multilevel compositional substitution model. The first six lists contain the results of the substitution estimation for a compositional part.

substitution 45

Mean CI_low and CI_hi	Posterior means.
	95% credible intervals.
Delta	Amount substituted across compositional parts.
From	Compositional part that is substituted from.
То	Compositional parts that is substituted to.
Level	Level where changes in composition takes place.
Reference	Either grandmean, clustermean, or users.

See Also

```
substitution
```

Examples

substitution

Multilevel Compositional Substitution Analysis

Description

Estimate the difference in an outcome when compositional parts are substituted for specific unit(s). The substitution output encapsulates the substitution results for all compositional parts present in the brmcoda object.

```
substitution(
  object,
  delta,
  ref = c("grandmean", "clustermean"),
  level = c("between", "within", "aggregate"),
  basesub,
  aorg = TRUE,
```

46 substitution

```
summary = TRUE,
weight = c("equal", "proportional"),
scale = c("response", "linear"),
comparison = NULL,
cores = NULL,
...
)
```

Arguments

aorg

object A fitted brmcoda object.

delta A integer, numeric value or vector indicating the amount of substituted change

between compositional parts.

ref Either a character value or vector or a dataset. Can be "grandmean" and/or

"clustermean", or a data.frame or data.table of user's specified reference grid consisting of combinations of covariates over which predictions are made. User's specified reference grid is only possible for simple substitution. Single

level models are default to "grandmean".

level A character string or vector. Should the estimate of multilevel models focus on

the "between" and/or "within" or "aggregate" variance? Single-level models

are default to "aggregate".

basesub A base substitution. Can be a data.frame or data.table of the base possi-

ble substitution of compositional parts, which can be computed using function build.basesub. If "one-to-all", all possible one-to-remaining reallocations

are estimated. If NULL, all possible one-to-one reallocations are estimated.

A logical value to obtain (a)verage prediction (o)ver the (r)eference (g)rid. Should the estimate at each level of the reference grid (FALSE) or their average (TRUE) be returned? Default is TRUE. Only applicable for model with covariates in addition

to the isometric log-ratio coordinates (i.e., adjusted model).

summary A logical value to obtain summary statistics instead of the raw values. Default is

TRUE. Currently only support outputing raw values for model using grandmean

as reference composition.

weight A character value specifying the weight to use in calculation of the reference

composition. If "equal", give equal weight to units (e.g., individuals). If "proportional", weights in proportion to the frequencies of units being averaged (e.g., observations across individuals). Default to "equal" for ref =

"grandmean" and "proportional" for ref = "clustermean".

scale Either "response" or "linear". If "response", results are returned on the

scale of the response variable. If "linear", results are returned on the scale of the linear predictor term, that is without applying the inverse link function or

other transformations.

comparison internally used only.

cores Number of cores to use when executing the chains in parallel, we recommend

setting the mc.cores option to be as many processors as the hardware and RAM allow (up to the number of compositional parts). For non-Windows OS in non-

interactive R sessions, forking is used instead of PSOCK clusters.

.. currently ignored.

substitution 47

Value

A list containing the results of multilevel compositional substitution model. The first six lists contain the results of the substitution estimation for a compositional part.

Mean Posterior means.

CI_low and CI_high
95% credible intervals.

Delta Amount substituted across compositional parts.

From Compositional part that is substituted from.

To Compositional parts that is substituted to.

Level Level where changes in composition takes place.

Reference Either grandmean, clustermean, or users.

Examples

```
if(requireNamespace("cmdstanr")){
 cilr <- complr(data = mcompd, sbp = sbp,</pre>
                 parts = c("TST", "WAKE", "MVPA", "LPA", "SB"),
                 idvar = "ID", total = 1440)
 # model with compositional predictor at between and within-person levels
 fit1 <- brmcoda(complr = cilr,</pre>
                  formula = Stress ~ bilr1 + bilr2 + bilr3 + bilr4 +
                                      wilr1 + wilr2 + wilr3 + wilr4 + (1 | ID),
                  chain = 1, iter = 500, backend = "cmdstanr")
 # one to one reallocation at between and within-person levels
 sub1 <- substitution(object = fit1, delta = 5, level = c("between"))</pre>
 summary(sub1)
 # one to all reallocation at between and within-person levels
 sub2 <- substitution(object = fit1, delta = 5, level = c("between", "within"),</pre>
                        basesub = "one-to-all")
 summary(sub2)
 # model with compositional predictor at aggregate level of variance
 fit2 <- brmcoda(complr = cilr,</pre>
                  formula = Stress ~ ilr1 + ilr2 + ilr3 + ilr4 + (1 | ID),
                  chain = 1, iter = 500, backend = "cmdstanr")
 sub3 <- substitution(object = fit2, delta = 5, level = c("aggregate"))</pre>
}
```

48 summary.complr

summary.brmcoda

Create a Summary of a fitted brmsfit model in a brmcoda object

Description

Create a Summary of a fitted brmsfit model in a brmcoda object

Usage

```
## S3 method for class 'brmcoda'
summary(object, ...)
```

Arguments

object An object of class brmcoda.... Other arguments passed to summary.brmsfit.

Examples

summary.complr

Create a Summary of a complr object

Description

Create a Summary of a complr object

```
## S3 method for class 'complr'
summary(object, weight = c("equal", "proportional"), ...)
```

49 summary.pivot_coord

Arguments

object An object of class complr.

A character value specifying the weight to use in calculation of the reference weight

> composition. If "equal", give equal weight to units (e.g., individuals). If "proportional", weights in proportion to the frequencies of units being av-

eraged (e.g., observations across individuals). Default is equal.

generic argument, not in use.

Examples

```
cilr <- complr(data = mcompd, sbp = sbp,</pre>
                parts = c("TST", "WAKE", "MVPA", "LPA", "SB"),
                idvar = "ID")
summary(cilr)
cilr <- complr(data = mcompd, sbp = sbp,</pre>
                parts = c("TST", "WAKE", "MVPA", "LPA", "SB"))
summary(cilr)
```

Create a Summary of a fitted brmsfit model from a pivot_coord summary.pivot_coord object

Description

Create a Summary of a fitted brmsfit model from a pivot_coord object

Usage

```
## S3 method for class 'pivot_coord'
summary(object, digits = 2, ...)
```

Arguments

object An object of class pivot_coord. A integer value used for number formatting. Default is 2. digits currently ignored. . . .

Value

A data table of results.

50 summary.substitution

Examples

summary.substitution $Create\ a\ Summary\ of\ a\ Substitution\ Model\ represented\ by\ a\ substitution\ object$

Description

Create a Summary of a Substitution Model represented by a substitution object

Usage

```
## S3 method for class 'substitution'
summary(object, delta, to, from, ref, level, digits = 2, ...)
```

Arguments

object	A substitution class object.
delta	A integer, numeric value or vector indicating the desired delta at which substitution results should be summarised. Default to all delta available in the substitution object.
to	A character value or vector specifying the names of the compositional parts that were reallocated to in the model.
from	A character value or vector specifying the names of the compositional parts that were reallocated from in the model.
ref	Either a character value or vector (("grandmean" and/or "clustermean" or "users"), Default to all ref available in the substitution object.
level	A character string or vector ("between" and/or "within"). Default to all level available in the substitution object.
digits	A integer value used for number formatting. Default is 2.
	generic argument, not in use.

update.brmcoda 51

Value

A summary of substitution object.

Mean Posterior means.

CI_low and CI_high

95% credible intervals.

Delta Amount substituted across compositional parts.

From Compositional part that is substituted from.

To Compositional parts that is substituted to.

Level Level where changes in composition takes place. Either between or within.

Reference Either grandmean, clustermean, or users.

Examples

update.brmcoda

Update brmcoda models

Description

This method allows for updating an existing brmcoda object.

Usage

```
## S3 method for class 'brmcoda'
update(object, formula. = NULL, newdata = NULL, newcomplr = NULL, ...)
```

Arguments

object A fitted brmcoda object to be updated.

formula. Changes to the formula; for details see update.formula and brmsformula.

newdata A data.frame or data.table containing data of all variables used in the anal-

ysis. It must include a composition and the same ID variable as the existing

complr object.

52 update.complr

newcomplr A complr object containing data of composition, ILR coordinates, and other

variables used in the updated model.

... Further arguments passed to brm.

Value

A brmcoda with two elements

complr An object of class complr used in the brm model.

model An object of class brmsfit, which contains the posterior draws along with many

other useful information about the model.

See Also

brmcoda

Examples

update.complr

Update complr

Description

This method allows for updating an existing complr object.

```
## S3 method for class 'complr'
update(object, newdata, ...)
```

update.complr 53

Arguments

object A complr class object to be updated.

newdata A data.frame or data.table containing data of all variables used in the anal-

ysis. It must include a composition and the same ID variable as the existing

complr object.

... generic argument, not in use.

Value

A complr object with at least the following elements.

comp A vector of class acomp representing one closed composition or a matrix of class

acomp representing multiple closed compositions each in one row.

between_comp A vector of class acomp representing one closed between-person composition or

a matrix of class acomp representing multiple closed between-person composi-

tions each in one row.

within_comp A vector of class acomp representing one closed within-person composition or a

matrix of class acomp representing multiple closed within-person compositions

each in one row.

logratio Log ratio transform of composition.

between_logratio

Log ratio transform of between-person composition.

within_logratio

Log ratio transform of within-person composition.

data The user's dataset or imputed dataset if the input data contains zeros.

transform Type of transform applied on compositional data.

parts Names of compositional variables.

idvar Name of the variable containing IDs.

total Total amount to which the compositions is closed.

See Also

complr

Examples

54 VarCorr.brmcoda

var.complr	Variance of compositions presented in a complr object.	
var.complr	Variance of compositions presented in a complr object.	

Description

Variance of compositions presented in a complr object.

Usage

```
## S3 method for class 'complr'
var(x, weight = c("equal", "proportional"), ...)
```

Arguments

X	An object of class complr.
weight	A character value specifying the weight to use in calculation of the reference composition. If "equal", give equal weight to units (e.g., individuals). If "proportional", weights in proportion to the frequencies of units being averaged (e.g., observations across individuals) Default is equal.

generic argument, not in use.

VarCorr.brmcoda Extract Variance and Correlation Components

Description

. . .

Calculates the estimated standard deviations, correlations and covariances of the group-level terms of the brmsfit object in a brmcoda object.

Usage

```
## S3 method for class 'brmcoda'
VarCorr(x, ...)
```

Arguments

An object of class brmcoda. Х Further arguments passed to VarCorr.brmsfit. . . .

Value

A list of lists (one per grouping factor), each with three elements: a matrix containing the standard deviations, an array containing the correlation matrix, and an array containing the covariance matrix with variances on the diagonal.

vcov.brmcoda 55

See Also

```
VarCorr.brmsfit
```

Examples

vcov.brmcoda

Covariance and Correlation Matrix of Population-Level Effects

Description

Get a point estimate of the covariance or correlation matrix of population-level parameters of the brmsfit object in a brmcoda object.

Usage

```
## S3 method for class 'brmcoda'
vcov(object, ...)
```

Arguments

object An object of class brmcoda.
... Further arguments passed to vcov.brmsfit.

Value

covariance or correlation matrix of population-level parameters

See Also

```
vcov.brmsfit
```

56 wsub

Examples

wsub

Within-person Simple Substitution

Description

This function is an alias of substitution to estimates the difference in an outcome when compositional parts are substituted for specific unit(s) at *within* level using a single reference composition (e.g., compositional mean at sample level). It is recommended that users run substitution model using the substitution function.

Usage

```
wsub(
  object,
  basesub,
  delta,
  ref = "grandmean",
  level = "within",
  weight = "equal",
  aorg = TRUE,
  summary = TRUE,
  scale = c("response", "linear"),
  comparison = "one-to-one",
  cores = NULL,
  ...
)
```

Arguments

object

A fitted brmcoda object.

basesub

A base substitution. Can be a data.frame or data.table of the base possible substitution of compositional parts, which can be computed using function build.basesub. If "one-to-all", all possible one-to-remaining reallocations are estimated. If NULL, all possible one-to-one reallocations are estimated.

wsub 57

delta A integer, numeric value or vector indicating the amount of substituted change

between compositional parts.

ref Either a character value or vector or a dataset. Can be "grandmean" and/or

"clustermean", or a data.frame or data.table of user's specified reference grid consisting of combinations of covariates over which predictions are made. User's specified reference grid is only possible for simple substitution. Single

level models are default to "grandmean".

level A character string or vector. Should the estimate of multilevel models focus on

the "between" and/or "within" or "aggregate" variance? Single-level models

are default to "aggregate".

weight A character value specifying the weight to use in calculation of the reference

composition. If "equal", give equal weight to units (e.g., individuals). If "proportional", weights in proportion to the frequencies of units being averaged (e.g., observations across individuals). Default to "equal" for ref =

"grandmean" and "proportional" for ref = "clustermean".

aorg A logical value to obtain (a) verage prediction (o) ver the (r) eference (g) rid. Should

the estimate at each level of the reference grid (FALSE) or their average (TRUE) be returned? Default is TRUE. Only applicable for model with covariates in addition

to the isometric log-ratio coordinates (i.e., adjusted model).

summary A logical value to obtain summary statistics instead of the raw values. Default is

TRUE. Currently only support outputing raw values for model using grandmean

as reference composition.

scale Either "response" or "linear". If "response", results are returned on the

scale of the response variable. If "linear", results are returned on the scale of the linear predictor term, that is without applying the inverse link function or

other transformations.

comparison internally used only.

cores Number of cores to use when executing the chains in parallel, we recommend

setting the mc. cores option to be as many processors as the hardware and RAM allow (up to the number of compositional parts). For non-Windows OS in non-

interactive R sessions, forking is used instead of PSOCK clusters.

... currently ignored.

Value

A list containing the results of multilevel compositional substitution model. The first six lists contain the results of the substitution estimation for a compositional part.

Mean Posterior means.

CI_low and CI_high

95% credible intervals.

Delta Amount substituted across compositional parts.

From Compositional part that is substituted from.

To Compositional parts that is substituted to.

Level Level where changes in composition takes place.

Reference Either grandmean, clustermean, or users.

58 wsubmargins

See Also

```
substitution
```

Examples

wsubmargins

Within-person Average Substitution

Description

This function is an alias of substitution to estimates the difference in an outcome when compositional parts are substituted for specific unit(s) at *within* level using cluster mean (e.g., compositional mean at individual level) as reference composition. It is recommended that users run substitution model using the substitution function.

```
wsubmargins(
  object,
  delta,
  basesub,
  summary = TRUE,
  ref = "clustermean",
  level = "within",
  weight = "proportional",
  scale = c("response", "linear"),
  comparison = "one-to-one",
  cores = NULL,
  ...
)
```

wsubmargins 59

Arguments

object A fitted brmcoda object.

delta A integer, numeric value or vector indicating the amount of substituted change

between compositional parts.

basesub A base substitution. Can be a data.frame or data.table of the base possi-

ble substitution of compositional parts, which can be computed using function build.basesub. If "one-to-all", all possible one-to-remaining reallocations are estimated. If NULL, all possible one-to-one reallocations are estimated.

summary A logical value to obtain summary statistics instead of the raw values. Default is

TRUE. Currently only support outputing raw values for model using grandmean

as reference composition.

ref Either a character value or vector or a dataset. Can be "grandmean" and/or

"clustermean", or a data.frame or data.table of user's specified reference grid consisting of combinations of covariates over which predictions are made. User's specified reference grid is only possible for simple substitution. Single

level models are default to "grandmean".

level A character string or vector. Should the estimate of multilevel models focus on

the "between" and/or "within" or "aggregate" variance? Single-level models

are default to "aggregate".

weight A character value specifying the weight to use in calculation of the reference

composition. If "equal", give equal weight to units (e.g., individuals). If "proportional", weights in proportion to the frequencies of units being averaged (e.g., observations across individuals). Default to "equal" for ref =

"grandmean" and "proportional" for ref = "clustermean".

scale Either "response" or "linear". If "response", results are returned on the

scale of the response variable. If "linear", results are returned on the scale of the linear predictor term, that is without applying the inverse link function or

other transformations.

comparison internally used only.

cores Number of cores to use when executing the chains in parallel, we recommend

setting the mc. cores option to be as many processors as the hardware and RAM allow (up to the number of compositional parts). For non-Windows OS in non-

interactive R sessions, forking is used instead of PSOCK clusters.

... currently ignored.

Value

A list containing the results of multilevel compositional substitution model. The first six lists contain the results of the substitution estimation for a compositional part.

Mean Posterior means.

CI_low and CI_high

95% credible intervals.

Delta Amount substituted across compositional parts.

From Compositional part that is substituted from.

60 wsubmargins

To Compositional parts that is substituted to.

Level Where changes in composition takes place.

Reference Either grandmean, clustermean, or users.

See Also

substitution

Examples

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