

Package: bmm (via r-universe)

November 19, 2024

Title Easy and Accessible Bayesian Measurement Models Using 'brms'

Version 1.0.0

Description Fit computational and measurement models using full Bayesian inference. The package provides a simple and accessible interface by translating complex domain-specific models into 'brms' syntax, a powerful and flexible framework for fitting Bayesian regression models using 'Stan'. The package is designed so that users can easily apply state-of-the-art models in various research fields, and so that researchers can use it as a new model development framework.

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URL <https://github.com/venpopov/bmm>, <https://venpopov.github.io/bmm/>

BugReports <https://github.com/venpopov/bmm/issues>

Depends R (>= 3.6.0)

Imports brms (>= 2.21.0), crayon, dplyr, fs, glue, magrittr, matrixStats, methods, parallel, stats, tidyr, withr

Suggests bookdown, cmdstanr (>= 0.7.0), cowplot, fansi, ggplot2, gghthemes, knitr, mixtur, remotes, rmarkdown, stringr, testthat (>= 3.0.0), tidybayes, usethis, waldo

Additional_repositories <https://mc-stan.org/r-packages/>

Config/testthat/edition 3

Encoding UTF-8

LazyData true

Roxygen list(markdown = TRUE)

RoxygenNote 7.3.1

Config/pak/sysreqs make libicu-dev

Repository <https://popov-lab.r-universe.dev>

RemoteUrl <https://github.com/venpopov/bmm>

RemoteRef v1.0.1

RemoteSha 8ada976031db6f46deea8e1f52e68046e8aae8a6

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 bmm-package

Easy and Accesible Bayesian Measurement Models Using 'brms'

Description

Fit computational and measurement models using full Bayesian inference. The package provides a simple and accessible interface by translating complex domain-specific models into 'brms' syntax, a powerful and flexible framework for fitting Bayesian regression models using 'Stan'. The package is designed so that users can easily apply state-of-the-art models in various research fields, and so that researchers can use it as a new model development framework.

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See Also

Useful links:

- <https://github.com/venpopov/bmm>
- <https://venpopov.github.io/bmm/>
- Report bugs at <https://github.com/venpopov/bmm/issues>

bmm

Fit Bayesian Measurement Models

Description

Fit a Bayesian measurement model using **brms** as a backend interface to Stan.

Usage

```
bmm(  
  formula,  
  data,  
  model,  
  prior = NULL,  
  sort_data = getOption("bmm.sort_data", "check"),  
  silent = getOption("bmm.silent", 1),  
  backend = getOption("brms.backend", NULL),  
  file = NULL,  
  file_compress = TRUE,  
  file_refit = getOption("bmm.file_refit", FALSE),  
  ...  
)  
  
fit_model(  
  formula,  
  data,  
  model,  
  prior = NULL,
```

```

  sort_data = getOption("bmm.sort_data", "check"),
  silent = getOption("bmm.silent", 1),
  backend = getOption("brms.backend", NULL),
  ...
)

```

Arguments

formula	An object of class <code>bmmformula</code> . A symbolic description of the model to be fitted.
data	An object of class <code>data.frame</code> , containing data of all variables used in the model. The names of the variables must match the variable names passed to the <code>bmmmodel</code> object for required arguments.
model	A description of the model to be fitted. This is a call to a <code>bmmmodel</code> such as <code>mixture3p()</code> function. Every model function has a number of required arguments which need to be specified within the function call. Call <code>supported_models()</code> to see the list of supported models and their required arguments
prior	One or more <code>brmsprior</code> objects created by <code>brms::set_prior()</code> or related functions and combined using the <code>c</code> method or the <code>+</code> operator. See also <code>default_prior()</code> for more help. Not necessary for the default model fitting, but you can provide prior constraints to model parameters
sort_data	Logical. If <code>TRUE</code> , the data will be sorted by the predictor variables for faster sampling. If <code>FALSE</code> , the data will not be sorted, but sampling will be slower. If "check" (the default), <code>bmm()</code> will check if the data is sorted, and ask you via a console prompt if it should be sorted. You can set the default value for this option using <code>global_options(bmm.sort_data = TRUE/FALSE/"check")</code> or <code>viabmm_options(sort_data)</code>
silent	Verbosity level between 0 and 2. If 1 (the default), most of the informational messages of compiler and sampler are suppressed. If 2, even more messages are suppressed. The actual sampling progress is still printed. Set <code>refresh = 0</code> to turn this off as well. If using <code>backend = "rstan"</code> you can also set <code>open_progress = FALSE</code> to prevent opening additional progress bars.
backend	Character. The backend to use for fitting the model. Can be "rstan" or "cmdstanr". If <code>NULL</code> (the default), "cmdstanr" will be used if the <code>cmdstanr</code> package is installed, otherwise "rstan" will be used. You can set the default backend using <code>global_options(brms.backend = "rstan"/"cmdstanr")</code>
file	Either <code>NULL</code> or a character string. If a string, the fitted model object is saved via <code>saveRDS</code> in a file named after the string. The <code>.rds</code> extension is added automatically. If the file object for later usage. If the directory of the file does not exist, it will be created.
file_compress	Logical or a character string, specifying one of the compression algorithms supported by <code>saveRDS</code> when saving the fitted model object.
file_refit	Logical. Modifies when the fit stored via the <code>file</code> argument is re-used. Can be set globally for the current R session via the "bmm.file_refit" option (see <code>options</code>). If <code>TRUE</code> (the default), the model is re-used if the file exists. If <code>FALSE</code> , the model is re-fitted. Note that unlike <code>brms</code> , there is no "on_change" option
...	Further arguments passed to <code>brms::brm()</code> or Stan. See the description of <code>brms::brm()</code> for more details

Value

An object of class `brmsfit` which contains the posterior draws along with many other useful information about the model. Use `methods(class = "brmsfit")` for an overview on available methods.

Supported Models

The following models are supported:

- `imm(resp_error, nt_features, nt_distances, set_size, regex, version)`
- `mixture2p(resp_error)`
- `mixture3p(resp_error, nt_features, set_size, regex)`
- `sdm(resp_error, version)`

Type `?modelName` to get information about a specific model, e.g. `?imm`

bmmformula syntax

see [this online article](#) for a detailed description of the syntax and how it differs from the syntax for **brmsformula**

Default priors, Stan code and Stan data

For more information about the default priors in **bmm** and about how to extract the Stan code and data generated by `bmm` and #' brms, see [the online article](#).

Miscellaneous

Type `help(package=bmm)` for a full list of available help topics.

`fit_model()` is a deprecated alias for `bmm()`.

References

Frischkorn, G. T., & Popov, V. (2023). A tutorial for estimating mixture models for visual working memory tasks in brms: Introducing the Bayesian Measurement Modeling (bmm) package for R. <https://doi.org/10.31234/osf.io/umt57>

See Also

[supported_models\(\)](#), [brms::brm\(\)](#), [default_prior\(\)](#), [bmmformula\(\)](#), [stancode\(\)](#), [standata\(\)](#)

Examples

```
# generate artificial data from the Signal Discrimination Model
dat <- data.frame(y = rsdm(2000))

# define formula
ff <- bmmformula(c ~ 1, kappa ~ 1)

# fit the model
fit <- bmm(formula = ff,
```

```

data = dat,
model = sdm(resp_error = "y"),
cores = 4,
backend = 'cmdstanr')

```

bmmformula

Create formula for predicting parameters of a bmmmodel

Description

This function is used to specify the formulas predicting the different parameters of a bmmmodel.

Usage

```
bmmformula(...)
```

```
bmf(...)
```

Arguments

... Formulas for predicting a bmmmodel parameter. Each formula for a parameter should be specified as a separate argument, separated by commas

Value

A list of formulas for each parameters being predicted

General formula structure

The formula argument accepts formulas of the following syntax:

```
parameter ~ fixed_effects + (random_effects | grouping_variable)
```

bmm formulas are built on brms formulas and function in nearly the same way, so you can use most of the brms formula syntax. The main differences is that in bmm formulas, the response variable is not specified in the formula. Instead, each parameter of the model is explicitly specified as the left-hand side of the formula. In brms, the response variable is always specified as the left-hand side of the first formula, which implicitly means that any predictors in the first formula are predictors of the mu parameter of the model. In general, measurement models do not all have a mu parameter, therefore it is more straightforward to explicitly predict each parameter of the model.

For example, in the following brms formula for the drift diffusion model, the first line corresponds to the drift rate parameter, but this is not explicitly stated.

```

brmsformula(rt | dec(response) ~ condition + (condition | id),
            bs ~ 1 + (1 | id),
            ndt ~ 1 + (1 | id),
            bias ~ 1 + (1 | id))

```

In `bmm`, the same formula would be written as:

```
bmmformula(drift ~ condition + (condition | id),
           bs ~ 1 + (1 | id),
           ndt ~ 1 + (1 | id),
           bias ~ 1 + (1 | id))
```

and the `rt` and response variables would be specified in the `model` argument of the `bmm()` function.

Aside from that, the `bmm` formula syntax is the same as the `brms` formula syntax. For more information on the `brms` formula syntax, see [brms::brmsformula\(\)](#).

You can also use the `bmf()` function as a shorthand for `bmmformula()`.

Examples

```
imm_formula <- bmmformula(
  c ~ 0 + set_size + (0 + set_size | id),
  a ~ 1,
  kappa ~ 0 + set_size + (0 + set_size | id)
)

# or use the shorter alias 'bmf'
imm_formula2 <- bmf(
  c ~ 0 + set_size + (0 + set_size | id),
  a ~ 1,
  kappa ~ 0 + set_size + (0 + set_size | id)
)
identical(imm_formula, imm_formula2)
```

bmm_options

View or change global bmm options

Description

View or change global bmm options

Usage

```
bmm_options(
  sort_data,
  parallel,
  default_priors,
  silent,
  color_summary,
  file_refit,
  reset_options = FALSE
)
```

Arguments

sort_data	logical. If TRUE, the data will be sorted by the predictors. If FALSE, the data will not be sorted, but sampling will be slower. If "check" (the default), bmm() will check if the data is sorted, and ask you via a console prompt if it should be sorted. Default: "check"
parallel	logical. If TRUE, chains will be run in parallel. If FALSE, chains will be run sequentially. You can also set these value for each model separately via the argument parallel in bmm(). Default: FALSE
default_priors	logical. If TRUE (default), the default bmm priors will be used. If FALSE, only the basic brms priors will be used. Default: TRUE
silent	numeric. Verbosity level between 0 and 2. If 1 (the default), most of the informational messages of compiler and sampler are suppressed. If 2, even more messages are suppressed. The actual sampling progress is still printed. Default: 1
color_summary	logical. If TRUE, the summary of the model will be printed in color. Default: TRUE
file_refit	logical. If TRUE, bmm() will refit the model even if the file argument is specified. Default: FALSE
reset_options	logical. If TRUE, the options will be reset to their default values Default: FALSE

Details

The `bmm_options` function is used to view or change the current bmm options. If no arguments are provided, the function will return the current options. If arguments are provided, the function will change the options and return the old options invisibly. If you provide only some of the arguments, the other options will not be changed. The options are stored in the global options list and will be used by `bmm()` and other functions in the `bmm` package. Each of these options can also be set manually using the built-in `options()` function, by setting the `bmm.sort_data`, `bmm.default_priors`, and `bmm.silent` options.

Value

A message with the current bmm options and their values, and invisibly returns the old options for use with `on.exit()` and friends.

Examples

```
# view the current options
bmm_options()

# change the options to always sort the data and to use parallel sampling
bmm_options(sort_data = TRUE, parallel = TRUE)

# restore the default options
bmm_options(reset_options = TRUE)

# you can change the options using the options() function as well
```



```
options(bmm.sort_data = TRUE, bmm.parallel = TRUE)
bmm_options()

# reset the options to their default values
bmm_options(reset_options = TRUE)

# bmm_options(sort_data = TRUE, parallel = TRUE) will also return the old options
# so you can use it with on.exit()
old_op <- bmm_options(sort_data = TRUE, parallel = TRUE)
on.exit(bmm_options(old_op))

bmm_options(reset_options = TRUE)
```

calc_error_relative_to_nontargets

Calculate response error relative to non-target values

Description

Given a vector of responses, and the values of non-targets, this function computes the error relative to each of the non-targets.

Usage

```
calc_error_relative_to_nontargets(data, response, nt_features)
```

Arguments

data	A data.frame object where each row is a single observation
response	Character. The name of the column in data which contains the response
nt_features	Character vector. The names of the columns in data which contain the values of the non-targets

Value

A data.frame with $n*m$ rows, where n is the number of rows of data and m is the number of non-target variables. It preserves all other columns of data, except for the non-target locations, and adds a column `y_nt`, which contains the transformed response error relative to the non-targets

Examples

```
data <- oberauer_lin_2017
data <- calc_error_relative_to_nontargets(data, "dev_rad", paste0("col_nt", 1:7))
hist(data$y_nt, breaks = 100)
```

circle_transform *Convert degrees to radians or radians to degrees.*

Description

The helper functions deg2rad and rad2deg should add convenience in transforming data from degrees to radians and from radians to degrees.

Usage

```
deg2rad(deg)
```

```
rad2deg(rad)
```

Arguments

deg A numeric vector of values in degrees.

rad A numeric vector of values in radians.

Value

A numeric vector of the same length as deg or rad.

Examples

```
degrees <- runif(100, min = 0, max = 360)
radians <- deg2rad(degrees)
degrees_again <- rad2deg(radians)
```

c_parametrizations *Convert between parametrizations of the c parameter of the SDM distribution*

Description

Convert between parametrizations of the c parameter of the SDM distribution

Usage

```
c_sqrtexp2bessel(c, kappa)
```

```
c_bessel2sqrtexp(c, kappa)
```

Arguments

c Vector of memory strength values

kappa Vector of precision values

Details

`c_bessel2sqrtexp` converts the memory strength parameter (`c`) from the `bessel` parametrization to the `sqrtexp` parametrization, `c_sqrtexp2bessel` converts from the `sqrtexp` parametrization to the `bessel` parametrization.

See [the online article](#) for details on the parameterization. The `sqrtexp` parametrization is the default in the `bmm` package.

Value

A numeric vector of the same length as `c` and `kappa`.

Examples

```
c_bessel <- c_sqrtexp2bessel(c = 4, kappa = 3)
c_sqrtexp <- c_bessel2sqrtexp(c = c_bessel, kappa = 3)
```

```
default_prior.bmmformula
```

Get Default priors for Measurement Models specified in BMM

Description

Obtain the default priors for a Bayesian multilevel measurement model, as well as information for which parameters priors can be specified. Given the `model`, the `data` and the `formula` for the model, this function will return the default priors that would be used to estimate the model. Additionally, it will return all model parameters that have no prior specified (flat priors). This can help to get an idea about which priors need to be specified and also know which priors were used if no user-specified priors were passed to the `bmm()` function.

The default priors in `bmm` tend to be more informative than the default priors in `brms`, as we use domain knowledge to specify the priors.

Usage

```
## S3 method for class 'bmmformula'
default_prior(object, data, model, formula = object, ...)
```

Arguments

<code>object</code>	A <code>bmmformula</code> object
<code>data</code>	An object of class <code>data.frame</code> , containing data of all variables used in the model. The names of the variables must match the variable names passed to the <code>bmmmodel</code> object for required arguments.
<code>model</code>	A description of the model to be fitted. This is a call to a <code>bmmmodel</code> such as <code>mixture3p()</code> function. Every model function has a number of required arguments which need to be specified within the function call. Call <code>supported_models()</code> to see the list of supported models and their required arguments

formula An object of class `bmmformula`. A symbolic description of the model to be fitted.
 ... Further arguments passed to `brms::default_prior()`

Value

A `data.frame` with columns specifying the prior, the class, the coef and group for each of the priors specified. Separate rows contain the information on the parameters (or parameter classes) for which priors can be specified.

See Also

`supported_models()`, `brms::default_prior()`

Examples

```
default_prior(bmf(c ~ 1, kappa ~ 1),
  data = oberauer_lin_2017,
  model = sdm(resp_error = "dev_rad")
)
```

fit_info

Extract information from a brmsfit object

Description

Extract information from a `brmsfit` object

Usage

```
fit_info(fit, what)
```

Arguments

`fit` A `brmsfit` object, or a list of `brmsfit` objects
`what` String. What to return:

- "time" for the sampling time per chain
- "time_mean" for the mean sampling time

Value

Depends on `what` and the class of `fit`. For `brmsfit` objects, information about the single fit is returned. For `brmsfit_list` objects, a list or `data.frame` with the information for each fit is returned.

- "time": A `data.frame` with the sampling time per chain
- "time_mean": A named numeric vector with the mean sampling time

Examples

```
fit <- bmm(
  formula = bmmformula(c ~ 1, kappa ~ 1),
  data = data.frame(y = rsdm(1000)),
  model = sdm(resp_error = "y")
)

fit_info(fit, "time")
```

 imm

Interference measurement model by Oberauer and Lin (2017).

Description

Three versions of the Interference measurement model by Oberauer and Lin (2017). - the full, bsc, and abc. `IMMfull()`, `IMMbsc()`, and `IMMabc()` are deprecated and will be removed in the future. Please use `imm(version = 'full')`, `imm(version = 'bsc')`, or `imm(version = 'abc')` instead.

Usage

```
imm(
  resp_error,
  nt_features,
  nt_distances,
  set_size,
  regex = FALSE,
  version = "full",
  ...
)
```

```
IMMfull(resp_error, nt_features, nt_distances, set_size, regex = FALSE, ...)
```

```
IMMbsc(resp_error, nt_features, nt_distances, set_size, regex = FALSE, ...)
```

```
IMMabc(resp_error, nt_features, set_size, regex = FALSE, ...)
```

Arguments

- | | |
|--------------------------|--|
| <code>resp_error</code> | The name of the variable in the provided dataset containing the response error. The response Error should code the response relative to the to-be-recalled target in radians. You can transform the response error in degrees to radian using the <code>deg2rad</code> function. |
| <code>nt_features</code> | A character vector with the names of the non-target variables. The <code>non_target</code> variables should be in radians and be centered relative to the target. Alternatively, if <code>regex=TRUE</code> , a regular expression can be used to match the non-target feature columns in the dataset. |

nt_distances	A vector of names of the columns containing the distances of non-target items to the target item. Alternatively, if regex=TRUE, a regular expression can be used to match the non-target distances columns in the dataset. Only necessary for the bsc and full versions.
set_size	Name of the column containing the set size variable (if set_size varies) or a numeric value for the set_size, if the set_size is fixed.
regex	Logical. If TRUE, the nt_features and nt_distances arguments are interpreted as a regular expression to match the non-target feature columns in the dataset.
version	Character. The version of the IMM model to use. Can be one of full, bsc, or abc. The default is full.
...	used internally for testing, ignore it

Details

- **Domain:** Visual working memory
- **Task:** Continuous reproduction
- **Name:** Interference measurement model by Oberauer and Lin (2017).
- **Citation:**
 - Oberauer, K., & Lin, H.Y. (2017). An interference model of visual working memory. *Psychological Review*, 124(1), 21-59

Version: full:

- **Requirements:**
 - The response variable should be in radians and represent the angular error relative to the target
- The non-target features should be in radians and be centered relative to the target
- **Parameters:**
 - mu1: Location parameter of the von Mises distribution for memory responses (in radians). Fixed internally to 0 by default.
 - kappa: Concentration parameter of the von Mises distribution
 - a: General activation of memory items
 - c: Context activation
 - s: Spatial similarity gradient
- **Fixed parameters:**
 - mu1 = 0
 - mu2 = 0
 - kappa2 = -100
- **Default parameter links:**
 - mu1 = tan_half; kappa = log; a = log; c = log; s = log
- **Default priors:**
 - mu1:
 - * main: student_t(1, 0, 1)

- kappa:
 - * main: normal(2, 1)
 - * effects: normal(0, 1)
- a:
 - * main: normal(0, 1)
 - * effects: normal(0, 1)
- c:
 - * main: normal(0, 1)
 - * effects: normal(0, 1)
- s:
 - * main: normal(0, 1)
 - * effects: normal(0, 1)

Version: bsc:

- **Requirements:**

- The response variable should be in radians and represent the angular error relative to the target
- The non-target features should be in radians and be centered relative to the target

- **Parameters:**

- mu1: Location parameter of the von Mises distribution for memory responses (in radians). Fixed internally to 0 by default.
- kappa: Concentration parameter of the von Mises distribution
- c: Context activation
- s: Spatial similarity gradient

- **Fixed parameters:**

- mu1 = 0
- mu2 = 0
- kappa2 = -100

- **Default parameter links:**

- mu1 = tan_half; kappa = log; c = log; s = log

- **Default priors:**

- mu1:
 - * main: student_t(1, 0, 1)
- kappa:
 - * main: normal(2, 1)
 - * effects: normal(0, 1)
- c:
 - * main: normal(0, 1)
 - * effects: normal(0, 1)
- s:
 - * main: normal(0, 1)
 - * effects: normal(0, 1)

Version: abc:

- **Requirements:**
 - The response variable should be in radians and represent the angular error relative to the target
- The non-target features should be in radians and be centered relative to the target
- **Parameters:**
 - μ_1 : Location parameter of the von Mises distribution for memory responses (in radians). Fixed internally to 0 by default.
 - κ : Concentration parameter of the von Mises distribution
 - a : General activation of memory items
 - c : Context activation
- **Fixed parameters:**
 - $\mu_1 = 0$
 - $\mu_2 = 0$
 - $\kappa_2 = -100$
- **Default parameter links:**
 - $\mu_1 = \tan_half$; $\kappa = \log$; $a = \log$; $c = \log$
- **Default priors:**
 - μ_1 :
 - * main: $student_t(1, 0, 1)$
 - κ :
 - * main: $normal(2, 1)$
 - * effects: $normal(0, 1)$
 - a :
 - * main: $normal(0, 1)$
 - * effects: $normal(0, 1)$
 - c :
 - * main: $normal(0, 1)$
 - * effects: $normal(0, 1)$

Additionally, all imm models have an internal parameter that is fixed to 0 to allow the model to be identifiable. This parameter is not estimated and is not included in the model formula. The parameter is:

- $b = \text{"Background activation (internally fixed to 0)"}$

Value

An object of class `bmmode1`

Examples

```
# load data
data <- oberauer_lin_2017

# define formula
```



```

ff <- bmmformula(
  kappa ~ 0 + set_size,
  c ~ 0 + set_size,
  a ~ 0 + set_size,
  s ~ 0 + set_size
)

# specify the full IMM model with explicit column names for non-target features and distances
# by default this fits the full version of the model
model1 <- imm(resp_error = "dev_rad",
  nt_features = paste0('col_nt', 1:7),
  nt_distances = paste0('dist_nt', 1:7),
  set_size = 'set_size')

# fit the model
fit <- bmm(formula = ff,
  data = data,
  model = model1,
  cores = 4,
  backend = 'cmdstanr')

# alternatively specify the IMM model with a regular expression to match non-target features
# this is equivalent to the previous call, but more concise
model2 <- imm(resp_error = "dev_rad",
  nt_features = 'col_nt',
  nt_distances = 'dist_nt',
  set_size = 'set_size',
  regex = TRUE)

# fit the model
fit <- bmm(formula = ff,
  data = data,
  model = model2,
  cores = 4,
  backend = 'cmdstanr')

# you can also specify the `bsc` or `abc` versions of the model to fit a reduced version
model3 <- imm(resp_error = "dev_rad",
  nt_features = 'col_nt',
  set_size = 'set_size',
  regex = TRUE,
  version = 'abc')
fit <- bmm(formula = ff,
  data = data,
  model = model3,
  cores = 4,
  backend = 'cmdstanr')

```

Description

Density, distribution, and random generation functions for the interference measurement model with the location of μ , strength of cue- dependent activation c , strength of cue-independent activation a , the generalization gradient s , and the precision of memory representations κ .

Usage

```
dimmm(  
  x,  
  mu = c(0, 2, -1.5),  
  dist = c(0, 0.5, 2),  
  c = 5,  
  a = 2,  
  b = 1,  
  s = 2,  
  kappa = 5,  
  log = FALSE  
)
```

```
piimm(  
  q,  
  mu = c(0, 2, -1.5),  
  dist = c(0, 0.5, 2),  
  c = 1,  
  a = 0.2,  
  b = 0,  
  s = 2,  
  kappa = 5  
)
```

```
qiimm(  
  p,  
  mu = c(0, 2, -1.5),  
  dist = c(0, 0.5, 2),  
  c = 1,  
  a = 0.2,  
  b = 0,  
  s = 2,  
  kappa = 5  
)
```

```
riimm(  
  n,  
  mu = c(0, 2, -1.5),  
  dist = c(0, 0.5, 2),  
  c = 1,  
  a = 0.2,  
  b = 1,
```

```

    s = 2,
    kappa = 5
  )

```

Arguments

x	Vector of observed responses
mu	Vector of locations
dist	Vector of distances of the item locations to the cued location
c	Vector of strengths for cue-dependent activation
a	Vector of strengths for cue-independent activation
b	Vector of baseline activation
s	Vector of generalization gradients
kappa	Vector of precision values
log	Logical; if TRUE, values are returned on the log scale.
q	Vector of quantiles
p	Vector of probability
n	Number of observations to generate data for

Value

`dimm` gives the density of the interference measurement model, `pimm` gives the cumulative distribution function of the interference measurement model, `qimm` gives the quantile function of the interference measurement model, and `rimm` gives the random generation function for the interference measurement model.

References

Oberauer, K., Stoneking, C., Wabersich, D., & Lin, H.-Y. (2017). Hierarchical Bayesian measurement models for continuous reproduction of visual features from working memory. *Journal of Vision*, 17(5), 11.

Examples

```

# generate random samples from the imm and overlay the density
r <- rimm(10000, mu = c(0, 2, -1.5), dist = c(0, 0.5, 2),
         c = 5, a = 2, s = 2, b = 1, kappa = 4)
x <- seq(-pi,pi,length.out=10000)
d <- dimm(x, mu = c(0, 2, -1.5), dist = c(0, 0.5, 2),
         c = 5, a = 2, s = 2, b = 1, kappa = 4)
hist(r, breaks=60, freq=FALSE)
lines(x,d,type="l", col="red")

```

k2sd	<i>Transform kappa of the von Mises distribution to the circular standard deviation</i>
------	---

Description

This function transforms the precision parameter kappa of the von Mises distribution to the circular standard deviation. Adapted from Matlab code by Paul Bays (<https://www.paulbays.com/code.php>)

Usage

```
k2sd(K)
```

Arguments

K numeric. A vector of kappa values.

Value

A vector of sd values.

Examples

```
kappas <- runif(1000, 0.01, 100)

# calculate SD (in radians)
SDs <- k2sd(kappas)

# transform SDs from radians to degrees
SDs_degrees <- SDs * 180 / pi

# plot the relationship between kappa and circular SD
plot(kappas, SDs)
plot(kappas, SDs_degrees)
```

mixture2p	<i>Two-parameter mixture model by Zhang and Luck (2008).</i>
-----------	--

Description

Two-parameter mixture model by Zhang and Luck (2008).

Usage

```
mixture2p(resp_error, ...)
```

Arguments

resp_error	The name of the variable in the provided dataset containing the response error. The response Error should code the response relative to the to-be-recalled target in radians. You can transform the response error in degrees to radian using the deg2rad function.
...	used internally for testing, ignore it

Details

- **Domain:** Visual working memory
- **Task:** Continuous reproduction
- **Name:** Two-parameter mixture model by Zhang and Luck (2008).
- **Citation:**
 - Zhang, W., & Luck, S. J. (2008). Discrete fixed-resolution representations in visual working memory. *Nature*, 453(7192), 233-235
- **Requirements:**
 - The response vairable should be in radians and represent the angular error relative to the target
- **Parameters:**
 - mu1: Location parameter of the von Mises distribution for memory responses (in radians). Fixed internally to 0 by default.
 - kappa: Concentration parameter of the von Mises distribution
 - thetat: Mixture weight for target responses
- **Fixed parameters:**
 - mu1 = 0
 - mu2 = 0
 - kappa2 = -100
- **Default parameter links:**
 - mu1 = tan_half; kappa = log; thetat = identity
- **Default priors:**
 - mu1:
 - * main: student_t(1, 0, 1)
 - kappa:
 - * main: normal(2, 1)
 - * effects: normal(0, 1)
 - thetat:
 - * main: logistic(0, 1)

Value

An object of class `bmmode1`

Examples

```
# generate artificial data
dat <- data.frame(y = rmixture2p(n=2000))

# define formula
ff <- bmmformula(kappa ~ 1, thetat ~ 1)

model <- mixture2p(resp_error = "y")

# fit the model
fit <- bmm(formula = ff,
           data = dat,
           model = model,
           cores = 4,
           iter = 500,
           backend = 'cmdstanr')
```

mixture2p_dist	<i>Distribution functions for the two-parameter mixture model (mixture2p)</i>
----------------	---

Description

Density, distribution, and random generation functions for the two-parameter mixture model with the location of μ , precision of memory representations κ and probability of recalling items from memory p_{mem} .

Usage

```
dmixture2p(x, mu = 0, kappa = 5, p_mem = 0.6, log = FALSE)
```

```
pmixture2p(q, mu = 0, kappa = 7, p_mem = 0.8)
```

```
qmixture2p(p, mu = 0, kappa = 5, p_mem = 0.6)
```

```
rmixture2p(n, mu = 0, kappa = 5, p_mem = 0.6)
```

Arguments

x	Vector of observed responses
mu	Vector of locations
kappa	Vector of precision values
p_mem	Vector of probabilities for memory recall
log	Logical; if TRUE, values are returned on the log scale.
q	Vector of quantiles
p	Vector of probability
n	Number of observations to generate data for

Value

dmixture2p gives the density of the two-parameter mixture model, pmixture2p gives the cumulative distribution function of the two-parameter mixture model, qmixture2p gives the quantile function of the two-parameter mixture model, and rmixture2p gives the random generation function for the two-parameter mixture model.

References

Zhang, W., & Luck, S. J. (2008). Discrete fixed-resolution representations in visual working memory. *Nature*, 453.

Examples

```
# generate random samples from the mixture2p model and overlay the density
r <- rmixture2p(10000, mu = 0, kappa = 4, p_mem = 0.8)
x <- seq(-pi,pi,length.out=10000)
d <- dmixture2p(x, mu = 0, kappa = 4, p_mem = 0.8)
hist(r, breaks=60, freq=FALSE)
lines(x,d,type="l", col="red")
```

mixture3p

Three-parameter mixture model by Bays et al (2009).

Description

Three-parameter mixture model by Bays et al (2009).

Usage

```
mixture3p(resp_error, nt_features, set_size, regex = FALSE, ...)
```

Arguments

resp_error	The name of the variable in the dataset containing the response error. The response error should code the response relative to the to-be-recalled target in radians. You can transform the response error in degrees to radians using the deg2rad function.
nt_features	A character vector with the names of the non-target feature values. The non_target feature values should be in radians and centered relative to the target. Alternatively, if regex=TRUE, a regular expression can be used to match the non-target feature columns in the dataset.
set_size	Name of the column containing the set size variable (if set_size varies) or a numeric value for the set_size, if the set_size is fixed.
regex	Logical. If TRUE, the nt_features argument is interpreted as a regular expression to match the non-target feature columns in the dataset.
...	used internally for testing, ignore it

Details

- **Domain:** Visual working memory
- **Task:** Continuous reproduction
- **Name:** Three-parameter mixture model by Bays et al (2009).
- **Citation:**
 - Bays, P. M., Catalao, R. F. G., & Husain, M. (2009). The precision of visual working memory is set by allocation of a shared resource. *Journal of Vision*, 9(10), 1-11
- **Requirements:**
 - The response variable should be in radians and represent the angular error relative to the target
- The non-target features should be in radians and be centered relative to the target
- **Parameters:**
 - mu1: Location parameter of the von Mises distribution for memory responses (in radians). Fixed internally to 0 by default.
 - kappa: Concentration parameter of the von Mises distribution
 - thetat: Mixture weight for target responses
 - thetant: Mixture weight for non-target responses
- **Fixed parameters:**
 - mu1 = 0
 - mu2 = 0
 - kappa2 = -100
- **Default parameter links:**
 - mu1 = tan_half; kappa = log; thetat = identity; thetant = identity
- **Default priors:**
 - mu1:
 - * main: student_t(1, 0, 1)
 - kappa:
 - * main: normal(2, 1)
 - * effects: normal(0, 1)
 - thetat:
 - * main: logistic(0, 1)
 - thetant:
 - * main: logistic(0, 1)

Value

An object of class `bmmode1`

Examples

```

# generate artificial data from the Bays et al (2009) 3-parameter mixture model
dat <- data.frame(
  y = rmixture3p(n=2000, mu = c(0,1,-1.5,2)),
  nt1_loc = 1,
  nt2_loc = -1.5,
  nt3_loc = 2
)

# define formula
ff <- bmmformula(
  kappa ~ 1,
  thetat ~ 1,
  thetant ~ 1
)

# specify the 3-parameter model with explicit column names for non-target features
model1 <- mixture3p(resp_error = "y", nt_features = paste0('nt',1:3, '_loc'), set_size = 4)

# fit the model
fit <- bmm(formula = ff,
  data = dat,
  model = model1,
  cores = 4,
  iter = 500,
  backend = 'cmdstanr')

# alternatively specify the 3-parameter model with a regular expression to match non-target features
# this is equivalent to the previous call, but more concise
model2 <- mixture3p(resp_error = "y", nt_features = "nt.*_loc", set_size = 4, regex = TRUE)

# fit the model
fit <- bmm(formula = ff,
  data = dat,
  model = model2,
  cores = 4,
  iter = 500,
  backend = 'cmdstanr')

```

mixture3p_dist

Distribution functions for the three-parameter mixture model (mixture3p)

Description

Density, distribution, and random generation functions for the three-parameter mixture model with the location of μ , precision of memory representations κ , probability of recalling items from memory p_{mem} , and probability of recalling non-targets p_{nt} .

Usage

```
dmixture3p(
  x,
  mu = c(0, 2, -1.5),
  kappa = 5,
  p_mem = 0.6,
  p_nt = 0.2,
  log = FALSE
)

pmixture3p(q, mu = c(0, 2, -1.5), kappa = 5, p_mem = 0.6, p_nt = 0.2)

qmixture3p(p, mu = c(0, 2, -1.5), kappa = 5, p_mem = 0.6, p_nt = 0.2)

rmixture3p(n, mu = c(0, 2, -1.5), kappa = 5, p_mem = 0.6, p_nt = 0.2)
```

Arguments

x	Vector of observed responses
mu	Vector of locations. First value represents the location of the target item and any additional values indicate the location of non-target items.
kappa	Vector of precision values
p_mem	Vector of probabilities for memory recall
p_nt	Vector of probabilities for swap errors
log	Logical; if TRUE, values are returned on the log scale.
q	Vector of quantiles
p	Vector of probability
n	Number of observations to generate data for

Value

dmixture3p gives the density of the three-parameter mixture model, pmixture3p gives the cumulative distribution function of the two-parameter mixture model, qmixture3p gives the quantile function of the two-parameter mixture model, and rmixture3p gives the random generation function for the two-parameter mixture model.

References

Bays, P. M., Catalao, R. F. G., & Husain, M. (2009). The precision of visual working memory is set by allocation of a shared resource. *Journal of Vision*, 9(10), 7.

Examples

```
# generate random samples from the mixture3p model and overlay the density
r <- rmixture3p(10000, mu = c(0, 2, -1.5), kappa = 4, p_mem = 0.6, p_nt = 0.2)
x <- seq(-pi, pi, length.out=10000)
d <- dmixture3p(x, mu = c(0, 2, -1.5), kappa = 4, p_mem = 0.6, p_nt = 0.2)
```

```
hist(r, breaks=60, freq=FALSE)
lines(x,d,type="l", col="red")
```

oberauer_lin_2017 *Data from Experiment 1 reported by Oberauer & Lin (2017)*

Description

Raw data of 19 subjects that completed a continuous reproduction task with set size 1 to 8 reported by Oberauer & Lin (2017).

Usage

```
oberauer_lin_2017
```

Format

oberauer_lin_2017:

A data frame with 15,200 rows and 19 columns:

ID Integer uniquely identifying different subjects

session Session number

trial Trial number within each session

set_size The set_size of the data in this row

dev_rad The response error, that is the difference between the response given and the target color in radians.

col_nt1, col_nt2, col_nt3, col_nt4, col_nt5, col_nt6, col_nt7 The non-target items' color value relative to the target.

dist_nt1, dist_nt2, dist_nt3, dist_nt4, dist_nt5, dist_nt6, dist_nt7, dist_nt8 The spatial distance between all non-target items and the target item in radians.

Source

<https://osf.io/m4shu>

restructure.bmmfit *Restructure Old bmmfit Objects*

Description

Restructure old `bmmfit` objects to work with the latest **bmm** version. This function is called internally when applying post-processing methods.

Usage

```
## S3 method for class 'bmmfit'  
restructure(x, ...)
```

Arguments

`x` An object of class `bmmfit`.
`...` Currently ignored.

Value

A `bmmfit` object compatible with the latest version of **bmm** and **brms**.

Examples

```
# Load an old bmmfit object  
old_fit <- readRDS("bmmfit_old.rds")  
new_fit <- restructure(old_fit)
```

sdm *Signal Discrimination Model (SDM) by Oberauer (2023)*

Description

Signal Discrimination Model (SDM) by Oberauer (2023)

Usage

```
sdm(resp_error, version = "simple", ...)  
  
sdmSimple(resp_error, version = "simple", ...)
```

Arguments

resp_error	The name of the variable in the dataset containing the response error. The response error should code the response relative to the to-be-recalled target in radians. You can transform the response error in degrees to radians using the deg2rad function.
version	Character. The version of the model to use. Currently only "simple" is supported.
...	used internally for testing, ignore it

Details

see [the online article](#) for a detailed description of the model and how to use it. * **Domain:** Visual working memory

- **Task:** Continuous reproduction
- **Name:** Signal Discrimination Model (SDM) by Oberauer (2023)
- **Citation:**
 - Oberauer, K. (2023). Measurement models for visual working memory - A factorial model comparison. *Psychological Review*, 130(3), 841-852
- **Version:** simple
- **Requirements:**
 - The response variable should be in radians and represent the angular error relative to the target
- **Parameters:**
 - mu: Location parameter of the SDM distribution (in radians; by default fixed internally to 0)
 - c: Memory strength parameter of the SDM distribution
 - kappa: Precision parameter of the SDM distribution
- **Fixed parameters:**
 - mu = 0
- **Default parameter links:**
 - mu = tan_half; c = log; kappa = log
- **Default priors:**
 - mu:
 - * main: student_t(1, 0, 1)
 - kappa:
 - * main: student_t(5, 1.75, 0.75)
 - * effects: normal(0, 1)
 - c:
 - * main: student_t(5, 2, 0.75)
 - * effects: normal(0, 1)

Value

An object of class `bmmmodel`

Examples

```
# simulate data from the model
dat <- data.frame(y = rsdm(n = 1000, c = 4, kappa = 3))

# specify formula
ff <- bmf(c ~ 1,
          kappa ~ 1)

# specify the model
fit <- bmm(formula = ff,
           data = dat,
           model = sdm(resp_error = 'y'),
           cores = 4,
           backend = 'cmdstanr')
```

SDMdist

Distribution functions for the Signal Discrimination Model (SDM)

Description

Density, distribution function, and random generation for the Signal Discrimination Model (SDM) Distribution with location μ , memory strength c , and precision κ . Currently only a single activation source is supported.

Usage

```
dsdm(x, mu = 0, c = 3, kappa = 3.5, log = FALSE, parametrization = "sqrtexp")

psdm(
  q,
  mu = 0,
  c = 3,
  kappa = 3.5,
  lower.tail = TRUE,
  log.p = FALSE,
  lower.bound = -pi,
  parametrization = "sqrtexp"
)

qsdm(p, mu = 0, c = 3, kappa = 3.5, parametrization = "sqrtexp")

rsdm(n, mu = 0, c = 3, kappa = 3.5, parametrization = "sqrtexp")
```

Arguments

x	Vector of quantiles
mu	Vector of location values in radians
c	Vector of memory strength values
kappa	Vector of precision values
log	Logical; if TRUE, values are returned on the log scale.
parametrization	Character; either "bessel" or "sqrtexp" (default). See the online article for details on the parameterization.
q	Vector of quantiles
lower.tail	Logical; If TRUE (default), return $P(X \leq x)$. Else, return $P(X > x)$
log.p	Logical; if TRUE, probabilities are returned on the log scale.
lower.bound	Numeric; Lower bound of integration for the cumulative distribution
p	Vector of probabilities
n	Number of observations to sample

Details**Parametrization**

See [the online article](#) for details on the parameterization. Oberauer (2023) introduced the SDM with the `bessel` parametrization. The `sqrtexp` parametrization is the default in the `bmm` package for numerical stability and efficiency. The two parametrizations are related by the functions `c_bessel2sqrtexp()` and `c_sqrtexp2bessel()`.

The cumulative distribution function

Since responses are on the circle, the cumulative distribution function requires you to choose a lower bound of integration. The default is $-\pi$, as for the `brms::pvon_mises()` function but you can choose any value in the argument `lower_bound` of `psdm`. Another useful choice is the mean of the response distribution minus π , e.g. `lower_bound = mu - pi`. This is the default in `circular::pvonmises()`, and it ensures that 50% of the cumulative probability mass is below the mean of the response distribution.

Value

`dsgm` gives the density, `psdm` gives the distribution function, `qsgm` gives the quantile function, `rsgm` generates random deviates, and `.dsgm_integrate` is a helper function for calculating the density of the SDM distribution.

References

Oberauer, K. (2023). Measurement models for visual working memory - A factorial model comparison. *Psychological Review*, 130(3), 841–852

Examples

```

# plot the density of the SDM distribution
x <- seq(-pi,pi,length.out=10000)
plot(x,dsdm(x,0,2,3),type="l", xlim=c(-pi,pi),ylim=c(0,1),
      xlab="Angle error (radians)",
      ylab="density",
      main="SDM density")
lines(x,dsdm(x,0,9,1),col="red")
lines(x,dsdm(x,0,2,8),col="green")
legend("topright",c("c=2, kappa=3.0, mu=0",
                   "c=9, kappa=1.0, mu=0",
                   "c=2, kappa=8, mu=1"),
      col=c("black","red","green"),lty=1, cex=0.8)

# plot the cumulative distribution function of the SDM distribution
p <- psdm(x, mu = 0, c = 3.1, kappa = 5)
plot(x,p,type="l")

# generate random deviates from the SDM distribution and overlay the density
r <- rsdm(10000, mu = 0, c = 3.1, kappa = 5)
d <- dsdm(x, mu = 0, c = 3.1, kappa = 5)
hist(r, breaks=60, freq=FALSE)
lines(x,d,type="l", col="red")

```

 softmax

Softmax and logsoftmax functions and their inverse functions

Description

softmax returns the value of the softmax function softmaxinv returns the value of the inverse-softmax function

Usage

```
softmax(eta, lambda = 1)
```

```
softmaxinv(p, lambda = 1)
```

Arguments

eta	A numeric vector input
lambda	Tuning parameter (a single positive value)
p	A probability vector (i.e., numeric vector of non-negative values that sum to one)

Details

The softmax function is a bijective function that maps a real vector with length $m-1$ to a probability vector with length m with all non-zero probabilities. The present functions define the softmax function and its inverse, both with a tuning parameter.

The current functions define the softmax as:

$$P(\eta_i) = \frac{e^{\lambda\eta_i}}{1 + \sum_{j=1}^m e^{\lambda\eta_j}}$$

Code adapted from the [utilities](#) package

Value

Value of the softmax function or its inverse

Examples

```
softmax(5:7)
softmaxinv(softmax(5:7))
```

stancode.bmmformula *Generate Stan code for bmm models*

Description

Given the model, the data and the formula for the model, this function will return the combined stan code generated by bmm and brms

Usage

```
## S3 method for class 'bmmformula'
stancode(object, data, model, prior = NULL, ...)
```

Arguments

object	A bmmformula object
data	An object of class data.frame, containing data of all variables used in the model. The names of the variables must match the variable names passed to the bmmmodel object for required arguments.
model	A description of the model to be fitted. This is a call to a bmmmodel such as mixture3p() function. Every model function has a number of required arguments which need to be specified within the function call. Call supported_models() to see the list of supported models and their required arguments
prior	One or more brmsprior objects created by brms::set_prior() or related functions and combined using the c method or the + operator. See also default_prior() for more help. Not necessary for the default model fitting, but you can provide prior constraints to model parameters

... Further arguments passed to `brms::stancode()`. See the description of `brms::stancode()` for more details

Value

A character string containing the fully commented Stan code to fit a bmm model.

See Also

[supported_models\(\)](#), [brms::stancode\(\)](#)

Examples

```
scode1 <- stancode(bmf(c ~ 1, kappa ~ 1),
  data = oberauer_lin_2017,
  model = sdm(resp_error = "dev_rad")
)
cat(scode1)
```

standata.bmmformula *Stan data for bmm models*

Description

Given the model, the data and the formula for the model, this function will return the combined stan data generated by bmm and brms

Usage

```
## S3 method for class 'bmmformula'
standata(object, data, model, prior = NULL, ...)
```

Arguments

object	A bmmformula object
data	An object of class data.frame, containing data of all variables used in the model. The names of the variables must match the variable names passed to the bmmmodel object for required arguments.
model	A description of the model to be fitted. This is a call to a bmmmodel such as <code>mixture3p()</code> function. Every model function has a number of required arguments which need to be specified within the function call. Call supported_models() to see the list of supported models and their required arguments
prior	One or more <code>brmsprior</code> objects created by brms::set_prior() or related functions and combined using the <code>c</code> method or the <code>+</code> operator. See also default_prior() for more help. Not necessary for the default model fitting, but you can provide prior constraints to model parameters
...	Further arguments passed to brms::standata() . See the description of brms::standata() for more details

Value

A named list of objects containing the required data to fit a bmm model with Stan.

See Also

[supported_models\(\)](#), [brms::standata\(\)](#)

Examples

```
sdata1 <- standata(bmf(c ~ 1, kappa ~ 1),
  data = oberauer_lin_2017,
  model = sdm(resp_error = "dev_rad")
)
str(sdata1)
```

summary.bmmfit

Create a summary of a fitted model represented by a bmmfit object

Description

Create a summary of a fitted model represented by a bmmfit object

Usage

```
## S3 method for class 'bmmfit'
summary(
  object,
  priors = FALSE,
  prob = 0.95,
  robust = FALSE,
  mc_se = FALSE,
  ...,
  backend = "bmm"
)
```

Arguments

object	An object of class brmsfit.
priors	Logical; Indicating if priors should be included in the summary. Default is FALSE.
prob	A value between 0 and 1 indicating the desired probability to be covered by the uncertainty intervals. The default is 0.95.
robust	If FALSE (the default) the mean is used as the measure of central tendency and the standard deviation as the measure of variability. If TRUE, the median and the median absolute deviation (MAD) are applied instead.

mc_se	Logical; Indicating if the uncertainty in Estimate caused by the MCMC sampling should be shown in the summary. Defaults to FALSE.
...	Other potential arguments
backend	Choose whether to display the <i>bmm</i> summary method (default), or to display the <i>brms</i> summary method.

Note

You can turn off the color output by setting the option `options(bmm.color_summary = FALSE)` or `bmm_options(color_summary = FALSE)`

See Also

[summary.brmsfit](#)

Examples

```
# generate artificial data from the Signal Discrimination Model
dat <- data.frame(y = rsdm(2000))

# define formula
ff <- bmmformula(c ~ 1, kappa ~ 1)

# fit the model
fit <- bmm(
  formula = ff,
  data = dat,
  model = sdm(resp_error = "y"),
  cores = 4,
  backend = "cmdstanr"
)

# summary of the model
summary(fit)
```

supported_models

Measurement models available in bmm

Description

Measurement models available in bmm

Usage

```
supported_models(print_call = TRUE)
```

Arguments

`print_call` Logical; If TRUE (default), the function will print information about how each model function should be called and its required arguments. If FALSE, the function will return a character vector with the names of the available models

Value

A character vector of measurement models available in bmm

Examples

```
supported_models()
```

<code>update.bmmfit</code>	<i>Update a bmm model</i>
----------------------------	---------------------------

Description

Update an existing bmm mode. This function calls `brms::update.brmsfit()`, but it applies the necessary bmm postprocessing to the model object before and after the update.

Usage

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

Arguments

`object` An object of class `bmmfit`

`formula.` A `bmmformula()`. If missing, the original formula is used. Currently you have to specify a full `bmmformula`

`newdata` An optional data frame containing the variables in the model

`recompile` Logical, indicating whether the Stan model should be recompiled. If NULL (the default), `update` tries to figure out internally, if recompilation is necessary. Setting it to FALSE will cause all Stan code changing arguments to be ignored.

`...` Further arguments passed to `brms::update.brmsfit()`

Details

When updating a `brmsfit` created with the `cmdstanr` backend in a different R session, a recompilation will be triggered because by default, `cmdstanr` writes the model executable to a temporary directory. To avoid that, set option `"cmdstanr_write_stan_file_dir"` to a nontemporary path of your choice before creating the original `bmmfit`.

For more information and examples, see `brms::update.brmsfit()`

Examples

```
# generate artificial data from the Signal Discrimination Model
# generate artificial data from the Signal Discrimination Model
dat <- data.frame(y = rsdm(2000))

# define formula
ff <- bmf(c ~ 1, kappa ~ 1)

# fit the model
fit <- bmm(formula = ff,
           data = dat,
           model = sdm(resp_error = "y"),
           cores = 4,
           backend = 'cmdstanr')

# update the model
fit <- update(fit, newdata = data.frame(y = rsdm(2000, kappa = 5)))
```

wrap

Wrap angles that extend beyond $(-\pi;\pi)$

Description

On the circular space, angles can be only in the range $(-\pi;\pi)$ or $(-180;180)$. When subtracting angles, this can result in values outside of this range. For example, when calculating the difference between a value of 10 degrees minus 340 degrees, this results in a difference of 330 degrees. However, the true difference between these two values is -30 degrees. This function wraps such values, so that they occur in the circle

Usage

```
wrap(x, radians = TRUE)
```

Arguments

x	A numeric vector, matrix or data.frame of angles to be wrapped. In radians (default) or degrees.
radians	Logical. Is x in radians (default=TRUE) or degrees (FALSE)

Value

An object of the same type as x

Examples

```
x <- runif(1000, -pi, pi)
y <- runif(1000, -pi, pi)
diff <- x - y
hist(diff)
wrapped_diff <- wrap(x - y)
hist(wrapped_diff)
```

zhang_luck_2008

Data from Experiment 2 reported by Zhang & Luck (2008)

Description

Raw data of 8 subjects for the response error in a continuous reproduction task with set size 1, 2, 3, and 6 reported by Zhang & Luck (2008).

Usage

zhang_luck_2008

Format

zhang_luck_2008:

A data frame with 4,000 rows and 9 columns:

subID Integer uniquely identifying different subjects

trial Trial identifier

setsize The set_size of the data in this row

response_error The response error, that is the difference between the response given and the target color in radians.

col_lure1, col_Lure2, col_Lure3, col_Lure4, col_Lure5 Color value of the lure items coded relative to the target color.

Source

<https://www.nature.com/articles/nature06860>

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