

Package ‘baggr’

September 2, 2021

Type Package

Title Bayesian Aggregate Treatment Effects

Version 0.6.9

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Description Running and comparing meta-analyses of data with hierarchical Bayesian models in Stan, including convenience functions for formatting data, plotting and pooling measures specific to meta-analysis. This implements many models from Meager (2019) <[doi:10.1257/app.20170299](https://doi.org/10.1257/app.20170299)>.

License GPL (>= 3)

Encoding UTF-8

LazyData true

Biarch true

Depends R (>= 3.5.0), Rcpp (>= 0.12.17)

Imports rstan (>= 2.18.1), rstantools (>= 2.1.1), bayesplot, crayon, forestplot, ggplot2, gridExtra, utils, stats, testthat, quantreg, methods

LinkingTo StanHeaders (>= 2.18.1), rstan (>= 2.18.1), BH (>= 1.66.0-1), Rcpp (>= 0.12.17), RcppParallel (>= 5.0.1), RcppEigen (>= 0.3.3.4.0)

SystemRequirements GNU make

NeedsCompilation yes

RoxygenNote 7.1.1

Suggests knitr, covr, rmarkdown

VignetteBuilder knitr

URL <https://github.com/wiecek/baggr>

BugReports <https://github.com/wiecek/baggr/issues>

Language en-GB

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Repository CRAN

Date/Publication 2021-09-02 06:50:02 UTC

R topics documented:

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| | |
|---------------|---|
| baggr-package | <i>baggr - a package for Bayesian meta-analysis</i> |
|---------------|---|

Description

This is *baggr* (pronounced as *bagger* or *badger*), a Bayesian meta-analysis package for R that uses **Stan** to fit the models. *Baggr* is intended to be user-friendly and transparent so that it's easier to understand the models you are building and criticise them.

Details

Baggr package provides a suite of models that work with both summary data and full data sets, to synthesise evidence collected from different groups, contexts or time periods. The `baggr` command automatically detects the data type and, by default, fits a partial pooling model (which you may know as **random effects models**) with weakly informative priors by calling **Stan** to carry out Bayesian inference. Modelling of variances or quantiles, standardisation and transformation of data are also possible.

Getting help

This is only a simple package help file. For documentation of the main function for conducting analyses see `baggr`. For description of models, data types and priors available in the package, try the built-in vignette (`vignette("baggr")`).

References

Stan Development Team (2020). RStan: the R interface to Stan. R package version 2.21.2. <https://mc-stan.org>

 baggr

Bayesian aggregate treatment effects model

Description

Bayesian inference on parameters of an average treatment effects model that's appropriate to the supplied individual- or group-level data, using Hamiltonian Monte Carlo in Stan. (For overall package help file see [baggr-package](#))

Usage

```
baggr(
  data,
  model = NULL,
  pooling = "partial",
  effect = NULL,
  covariates = c(),
  prior_hypermean = NULL,
  prior_hypersd = NULL,
  prior_hypercor = NULL,
  prior_beta = NULL,
  prior_control = NULL,
  prior_control_sd = NULL,
  prior = NULL,
  ppd = FALSE,
  pooling_control = "none",
  test_data = NULL,
  quantiles = seq(0.05, 0.95, 0.1),
  outcome = "outcome",
  group = "group",
  treatment = "treatment",
  silent = FALSE,
  warn = TRUE,
  ...
)
```

Arguments

| | |
|---------|--|
| data | data frame with summary or individual level data to meta-analyse |
| model | if NULL, detected automatically from input data otherwise choose from "rubin", "mutau", "rubin_full", "quantiles" (see Details). |
| pooling | Type of pooling; choose from "none", "partial" (default) and "full". If you are not familiar with the terms, consult the vignette; "partial" can be understood as random effects and "full" as fixed effects |
| effect | Label for effect. Will default to "mean" in most cases, "log OR" in logistic model, quantiles in quantiles model etc. These labels are used in various print |

and plot outputs. Comparable models (e.g. in [baggr_compare](#)) should have same effect.

| | |
|------------------|--|
| covariates | Character vector with column names in data. The corresponding columns are used as covariates (fixed effects) in the meta-regression model (in case of aggregate data). In the case of individual level data the model does not differentiate between group-level variables (same values of the covariate for all rows related to a given group) and individual-level covariates. |
| prior_hypermean | prior distribution for hypermean; you can use "plain text" notation like <code>prior_hypermean=normal(0, 100)</code> or <code>uniform(-10, 10)</code> . See <i>Details:Priors</i> section below for more possible specifications. If unspecified, the priors will be derived automatically based on data (and printed out in the console). |
| prior_hypersd | prior for hyper-standard deviation, used by Rubin and "mutau" models; same rules apply as for <code>_hypermean</code> ; |
| prior_hypercor | prior for hypercorrelation matrix, used by the "mutau" model |
| prior_beta | prior for regression coefficients if covariates are specified; will default to experimental <code>normal(0, 10^2)</code> distribution |
| prior_control | prior for the mean in the control arm (baseline), currently used in "logit" model only; if <code>pooling_control = "partial"</code> , the prior is hyperprior for all baselines, if "none", then it is an independent prior for all baselines |
| prior_control_sd | prior for the SD in the control arm (baseline), currently used in "logit" model only; this can only be used if <code>pooling_control = "partial"</code> |
| prior | alternative way to specify all priors as a named list with <code>hypermean</code> , <code>hypersd</code> , <code>hypercor</code> , <code>beta</code> , analogous to <code>prior_</code> arguments above, e.g. <code>prior = list(hypermean = normal(0, 10), beta = uniform(-50, 50))</code> |
| ppd | logical; use prior predictive distribution? (<i>p.p.d.</i>) If <code>ppd=TRUE</code> , Stan model will sample from the prior distribution(s) and ignore data in inference. However, <code>data</code> argument might still be used to infer the correct model (if <code>model=NULL</code>) and to set the default priors, therefore you must specify it. |
| pooling_control | Pooling for group-specific control mean terms (currently only in <code>logit</code>). Either "none" or "partial". |
| test_data | data for cross-validation; <code>NULL</code> for no validation, otherwise a data frame with the same columns as <code>data</code> argument. See "Cross-validation" section below. |
| quantiles | if <code>model = "quantiles"</code> , a vector indicating which quantiles of data to use (with values between 0 and 1) |
| outcome | character; column name in (individual-level) data with outcome variable values |
| group | character; column name in data with grouping factor; it's necessary for individual-level data, for summarised data it will be used as labels for groups when displaying results |
| treatment | character; column name in (individual-level) data with treatment factor; |
| silent | Whether to silence messages about prior settings and about other automatic behaviour. |

warn print an additional warning if Rhat exceeds 1.05
 ... extra options passed to Stan function, e.g. `control = list(adapt_delta = 0.99)`, number of iterations etc.

Details

Running baggr requires 1/ data preparation, 2/ choice of model, 3/ choice of priors. All three are discussed in depth in the package vignette (`vignette("baggr")`).

Data. For aggregate data models you need a data frame with columns `tau` and `se` or `tau, mu, se.tau, se.mu`. An additional column can be used to provide labels for each group (by default column `group` is used if available, but this can be customised – see the example below). For individual level data three columns are needed: `outcome`, `treatment`, `group`. These are identified by using the `outcome`, `treatment` and `group` arguments.

Many data preparation steps can be done through a helper function `prepare_ma`. It can convert individual to summary-level data, calculate odds/risk ratios (with/without corrections) in binary data, standardise variables and more. Using it will automatically format data inputs to work with `baggr()`.

Models. Available models are:

- for the **continuous variable** means: "rubin" model for average treatment effect (using summary data), "mutau" version which takes into account means of control groups (also using summary data), "rubin_full", which is the same model as "rubin" but works with individual-level data
- for **continuous variable quantiles**: "quantiles" model (see Meager, 2019 in references)
- for *mixture data*: "sslabs" (experimental)
- for **binary data**: "logit" model can be used on individual-level data; you can also analyse continuous statistics such as log odds ratios and logs risk ratios using the models listed above; see `vignette("baggr_binary")` for tutorial with examples

If no model is specified, the function tries to infer the appropriate model automatically. Additionally, the user must specify type of pooling. The default is always partial pooling.

Covariates. Both aggregate and individual-level data can include extra columns, given by `covariates` argument (specified as a character vector of column names) to be used in regression models. We also refer to impact of these covariates as *fixed effects*.

Two types of covariates may be present in your data:

- In "rubin" and "mutau" models, covariates that **change according to group unit**. In that case, the model accounting for the group covariates is a **meta-regression** model. It can be modelled on summary-level data.
- In "logit" and "rubin_full" models, covariates that **change according to individual unit**. Then, such a model is commonly referred to as a **mixed model**. It has to be fitted to individual-level data. Note that meta-regression is a special case of a mixed model for individual-level data.

Priors. It is optional to specify priors yourself, as the package will try propose an appropriate prior for the input data if you do not pass a `prior` argument. To set the priors yourself, use `prior_` arguments. For specifying many priors at once (or re-using between models), a single `prior =`

`list(...)` argument can be used instead. Meaning of the prior parameters may slightly change from model to model. Details and examples are given in `vignette("baggr")`. Setting `ppd=TRUE` can be used to obtain prior predictive distributions, which is useful for understanding the prior assumptions, especially useful in conjunction with `effect_plot`. You can also `baggr_compare` different priors by setting `baggr_compare(..., compare="prior")`.

Cross-validation. When `test_data` are specified, an extra parameter, the log predictive density, will be returned by the model. (The fitted model itself is the same regardless of whether there are `test_data`.) To understand this parameter, see documentation of `loocv`, a function that can be used to assess out of sample prediction of the model using all available data. If using individual-level data model, `test_data` should only include treatment arms of the groups of interest. (This is because in cross-validation we are not typically interested in the model's ability to fit heterogeneity in control arms, but only heterogeneity in treatment arms.) For using aggregate level data, there is no such restriction.

Outputs. By default, some outputs are printed. There is also a plot method for `baggr` objects which you can access via `baggr_plot` (or simply `plot()`). Other standard functions for working with `baggr` object are

- `treatment_effect` for distribution of hyperparameters
- `group_effects` for distributions of group-specific parameters
- `fixed_effects` for coefficients in (meta-)regression
- `effect_draw` and `effect_plot` for posterior predictive distributions
- `baggr_compare` for comparing multiple `baggr` models
- `loocv` for cross-validation
- `pp_check` for posterior predictive checks

Value

`baggr` class structure: a list including Stan model fit alongside input data, pooling metrics, various model properties. If test data is used, mean value of $-2 \cdot \text{lpd}$ is reported as `mean_lpd`

Author(s)

Witold Wiecek, Rachael Meager

Examples

```
df_pooled <- data.frame("tau" = c(1, -1, .5, -.5, .7, -.7, 1.3, -1.3),
                       "se" = rep(1, 8),
                       "state" = datasets::state.name[1:8])
baggr(df_pooled) #baggr automatically detects the input data
# same model, but with correct labels,
# different pooling & passing some options to Stan
baggr(df_pooled, group = "state", pooling = "full", iter = 500)
# model with non-default (and very informative) priors

baggr(df_pooled, prior_hypersd = normal(0, 2))
```

```
# "mu & tau" model, using a built-in dataset
# prepare_ma() can summarise individual-level data
ms <- microcredit_simplified
microcredit_summary_data <- prepare_ma(ms, outcome = "consumption")
baggr(microcredit_summary_data, model = "mutau",
      iter = 500, #this is just for illustration -- don't set it this low normally!
      pooling = "partial", prior_hypercor = lkj(1),
      prior_hypersd = normal(0,10),
      prior_hypermean = multinormal(c(0,0),matrix(c(10,3,3,10),2,2)))
```

baggr_compare

(Run and) compare multiple baggr models

Description

Compare multiple [baggr](#) models by either providing multiple already existing models as (named) arguments or passing parameters necessary to run a [baggr](#) model.

Usage

```
baggr_compare(
  ...,
  what = "pooling",
  compare = "groups",
  transform = NULL,
  plot = FALSE
)
```

Arguments

| | |
|-----------|--|
| ... | Either some (at least 1) objects of class <code>baggr</code> (you should name your objects, see the example below) or the same arguments you'd pass to baggr . In the latter case you must specify what to compare. |
| what | One of "pooling" (comparison between no, partial and full pooling) or "prior" (comparison between prior and posterior predictive). If pre-existing <code>baggr</code> models are passed to ..., this argument is ignored. |
| compare | When plotting, choose between comparison of "groups" (default) or (hyper-) "effects". The former is not available when what = "prior". |
| transform | a function (e.g. <code>exp()</code> , <code>log()</code>) to apply to the values of group (and hyper, if hyper=TRUE) effects before plotting; when working with effects that are on log scale, exponent transform is used automatically, you can plot on log scale by setting <code>transform = identity</code> |
| plot | logical; calls plot.baggr_compare when running <code>baggr_compare</code> |

Details

If you pass parameters to the function you must specify what kind of comparison you want, either "pooling", which will run fully/partially/un-pooled models and then compare them, or "prior" which will generate estimates without the data and compare them to the model with the full data. For more details see [baggr](#), specifically the `ppd` argument.

Value

an object of class `baggr_compare`

Author(s)

Witold Wiecek, Brice Green

See Also

[plot.baggr_compare](#) and [print.baggr_compare](#) for working with results of this function

Examples

```
# Most basic comparison between no, partial and full pooling
# (This will run the models)
# run model with just prior and then full data for comparison
# with the same arguments that are passed to baggr
prior_comparison <-
  baggr_compare(schools,
                model = 'rubin',
                #this is just for illustration -- don't set it this low normally!
                iter = 500,
                prior_hypermean = normal(0, 3),
                prior_hypersd = normal(0,2),
                prior_hypercor = lkj(2),
                what = "prior")
# print the aggregated treatment effects
prior_comparison
# plot the comparison of the two distributions
plot(prior_comparison)
# Now compare different types of pooling for the same model
pooling_comparison <-
  baggr_compare(schools,
                model = 'rubin',
                #this is just for illustration -- don't set it this low normally!
                iter = 500,
                prior_hypermean = normal(0, 3),
                prior_hypersd = normal(0,2),
                prior_hypercor = lkj(2),
                what = "pooling",
                # You can automatically plot:
                plot = TRUE)
# Compare existing models (you don't have to, but best to name them):
```

```

bg1 <- baggr(schools, pooling = "partial")
bg2 <- baggr(schools, pooling = "full")
baggr_compare("Partial pooling model" = bg1, "Full pooling" = bg2)

#' ...or simply draw from prior predictive dist (note ppd=T)
bg1 <- baggr(schools, ppd=TRUE)
bg2 <- baggr(schools, prior_hypermean = normal(0, 5), ppd=TRUE)
baggr_compare("Prior A, p.p.d."=bg1,
              "Prior B p.p.d."=bg2,
              compare = "effects")

# Compare how posterior predictive effect varies with e.g. choice of prior
bg1 <- baggr(schools, prior_hypersd = uniform(0, 20))
bg2 <- baggr(schools, prior_hypersd = normal(0, 5))
baggr_compare("Uniform prior on SD"=bg1,
              "Normal prior on SD"=bg2,
              compare = "effects", plot = TRUE)

# Models don't have to be identical. Compare different subsets of input data:
bg1_small <- baggr(schools[1:6,], pooling = "partial")
baggr_compare("8 schools model" = bg1, "First 6 schools" = bg1_small,
              plot = TRUE)

```

baggr_plot

Plotting method in baggr package

Description

Extracts study effects from the baggr model and sends them to one of bayesplot package plotting functions.

Usage

```

baggr_plot(
  bg,
  hyper = FALSE,
  style = "intervals",
  transform = NULL,
  prob = 0.5,
  prob_outer = 0.95,
  vline = FALSE,
  order = TRUE,
  ...
)

```

Arguments

| | |
|------------|--|
| bg | object of class baggr |
| hyper | logical; show hypereffect as the last row of the plot? |
| style | either "intervals" or "areas" |
| transform | a function (e.g. <code>exp()</code> , <code>log()</code>) to apply to the values of group (and hyper, if hyper=TRUE) effects before plotting; when working with effects that are on log scale, exponent transform is used automatically, you can plot on log scale by setting <code>transform = identity</code> |
| prob | Probability mass for the inner interval in visualisation |
| prob_outer | Probability mass for the outer interval in visualisation |
| vline | logical; show vertical line through 0 in the plot? |
| order | logical; sort groups by magnitude of treatment effect? |
| ... | extra arguments to pass to the bayesplot functions |

Value

ggplot2 object

Author(s)

Witold Wiecek; the visual style is based on *bayesplot* package

See Also

[bayesplot::MCMC-intervals](#) for more information about *bayesplot* functionality; [forest_plot](#) for a typical meta-analysis alternative; [effect_plot](#) for plotting treatment effects for a new group

Examples

```
fit <- baggr(schools, pooling = "none")
plot(fit)
plot(fit, style = "areas", order = FALSE)
```

baggr_theme_set *Set, get, and replace themes for baggr plots*

Description

These functions get, set, and modify the ggplot2 themes of the baggr plots. `baggr_theme_get()` returns a ggplot2 theme function for adding themes to a plot. `baggr_theme_set()` assigns a new theme for all plots of baggr objects. `baggr_theme_update()` edits a specific theme element for the current theme while holding the theme's other aspects constant. `baggr_theme_replace()` is used for wholesale replacing aspects of a plot's theme (see [ggplot2::theme_get\(\)](#)).

Usage

```
baggr_theme_set(new = bayesplot::theme_default())  
  
baggr_theme_get()  
  
baggr_theme_update(...)  
  
baggr_theme_replace(...)
```

Arguments

| | |
|-----|--------------------------------------|
| new | New theme to use for all baggr plots |
| ... | A named list of theme settings |

Details

Under the hood, many of the visualizations rely on the bayesplot package, and thus these leverage the `bayesplot::bayesplot_theme_get()` functions. By default, these match the bayesplot's package theme to make it easier to form cohesive graphs across this package and others. The trickiest of these to use is `baggr_theme_replace`; 9 times out of 10 you want `baggr_theme_update`.

Value

The get method returns the current theme, but all of the others invisibly return the old theme.

See Also

[bayesplot::bayesplot_theme_get](#)

Examples

```
# make plot look like default ggplots  
  
library(ggplot2)  
  
fit <- baggr(schools)  
baggr_theme_set(theme_grey())  
baggr_plot(fit)  
  
# use baggr_theme_get to return theme elements for current theme  
qplot(mtcars$mpg) + baggr_theme_get()  
  
# update specific aspect of theme you are interested in  
baggr_theme_update(text = element_text(family = "mono"))  
  
# undo that silliness  
baggr_theme_update(text = element_text(family = "serif"))  
  
# update and replace are similar, but replace overwrites the
```

```
# whole element, update just edits the aspect of the element
# that you give it
# this will error:
# baggr_theme_replace(text = element_text(family = "Times"))
# baggr_plot(fit)
# because it deleted everything else to do with text elements
```

binary_to_individual *Generate individual-level binary outcome data from an aggregate statistics*

Description

This is a helper function that is typically used automatically by some of *baggr* functions, such as when running `model="logit"` in [baggr](#), when summary-level data are supplied.

Usage

```
binary_to_individual(data, group = "group")
```

Arguments

| | |
|--------------------|--|
| <code>data</code> | A data frame with columns <code>a</code> , <code>c</code> and <code>b/n1</code> , <code>d/n2</code> . See <code>vignette("baggr_binary")</code> for an example of use and notation details. |
| <code>group</code> | Column name storing group |

Value

A data frame with columns `group`, `outcome` and `treatment`.

See Also

[prepare_ma](#) uses this function

Examples

```
df_yusuf <- read.table(text="
trial a n1 c n2i
Balcon 14 56 15 58
Clausen 18 66 19 64
Multicentre 15 100 12 95
Barber 10 52 12 47
Norris 21 226 24 228
Kahler 3 38 6 31
Ledwich 2 20 3 20
", header=TRUE)
bti <- binary_to_individual(df_yusuf, group = "trial")
head(bti)
```

```
# to go back to summary-level data
prepare_ma(bti, effect = "logOR")
# the last operation is equivalent to simply doing
prepare_ma(df_yusuf, group="trial", effect="logOR")
```

| | |
|--------|--|
| chicks | <i>Chickens: impact of electromagnetic field on calcium ion efflux in chicken brains</i> |
|--------|--|

Description

An experiment conducted by Blackman et al. (1988) and documented in the following [GitHub repository](#) by Vakar and Gelman. The dataset consists of a large number of experiments (tau, se.tau) repeated at varying wave frequencies. Sham experiments (mu, se.mu) are also included, allowing us to compare performance of models with and without control measurements.

Usage

```
chicks
```

Format

An object of class `data.frame` with 38 rows and 7 columns.

References

Blackman, C. F., S. G. Benane, D. J. Elliott, D. E. House, and M. M. Pollock. "Influence of Electromagnetic Fields on the Efflux of Calcium Ions from Brain Tissue in Vitro: A Three-Model Analysis Consistent with the Frequency Response up to 510 Hz." *Bioelectromagnetics* 9, no. 3 (1988): 215–27.

| | |
|----------------|--|
| convert_inputs | <i>Convert inputs for baggr models</i> |
|----------------|--|

Description

Converts data to Stan inputs, checks integrity of data and suggests default model if needed. Typically all of this is done automatically by [baggr](#), **this function is only for debugging** or running models "by hand".

Usage

```

convert_inputs(
  data,
  model,
  quantiles,
  group = "group",
  outcome = "outcome",
  treatment = "treatment",
  covariates = c(),
  test_data = NULL,
  silent = FALSE
)

```

Arguments

| | |
|------------|--|
| data | ‘data.frame’ with desired modelling input |
| model | valid model name used by baggr; see baggr for allowed models if model = NULL, this function will try to find appropriate model automatically |
| quantiles | vector of quantiles to use (only applicable if model = "quantiles") |
| group | name of the column with grouping variable |
| outcome | name of column with outcome variable (designated as string) |
| treatment | name of column with treatment variable |
| covariates | Character vector with column names in data. The corresponding columns are used as covariates (fixed effects) in the meta-regression model. |
| test_data | same format as data argument, gets left aside for testing purposes (see baggr) |
| silent | Whether to print messages when evaluated |

Details

Typically this function is only called within [baggr](#) and you do not need to use it yourself. It can be useful to understand inputs or to run models which you modified yourself.

Value

R structure that’s appropriate for use by [baggr](#) Stan models; group_label, model and n_groups are included as attributes and are necessary for [baggr](#) to work correctly

Author(s)

Witold Wiecek

Examples

```

# simple meta-analysis example,
# this is the formatted input for Stan models in baggr():
convert_inputs(schools, "rubin")

```

| | |
|------------|---|
| data_spike | <i>Spike & slab example dataset</i> |
|------------|---|

Description

Spike & slab example dataset

Usage

```
data_spike
```

Format

An object of class `data.frame` with 1500 rows and 4 columns.

| | |
|-------------|---|
| effect_draw | <i>Make predictive draws from baggr model</i> |
|-------------|---|

Description

This function takes the samples of hyperparameters from a `baggr` model (typically hypermean and hyper-SD, which you can see using `treatment_effect`) and draws values of new realisations of treatment effect, i.e. an additional draw from the "population of studies". This can be used for both prior and posterior draws, depending on `baggr` model.

Usage

```
effect_draw(
  x,
  n,
  transform = NULL,
  summary = FALSE,
  message = TRUE,
  interval = 0.95
)
```

Arguments

| | |
|------------------------|--|
| <code>x</code> | A <code>baggr</code> class object. |
| <code>n</code> | How many values to draw? The default is as long as the number of samples in the <code>baggr</code> object (see <i>Details</i>). |
| <code>transform</code> | a transformation (an R function) to apply to the result of a draw. |
| <code>summary</code> | logical; if TRUE returns summary statistics rather than samples from the distribution; |
| <code>message</code> | logical; use to disable messages prompted by using with no pooling models |
| <code>interval</code> | uncertainty interval width (numeric between 0 and 1), if <code>summary=TRUE</code> |

Details

The predictive distribution can be used to "combine" heterogeneity between treatment effects and uncertainty in the mean treatment effect. This is useful both in understanding impact of heterogeneity (see Riley et al, 2011, for a simple introduction) and for study design e.g. as priors in analysis of future data (since the draws can be seen as an expected treatment effect in a hypothetical study).

The default number of samples is the same as what is returned by Stan model implemented in [baggr](#), (depending on such options as `iter`, `chains`, `thin`). If `n` is larger than what is available in Stan model, we draw values with replacement. This is not recommended and warning is printed in these cases.

Under default settings in [baggr](#), a *posterior* predictive distribution is obtained. But `effect_draw` can also be used for *prior* predictive distributions when setting `ppd=T` in [baggr](#). The two outputs work exactly the same way.

Value

A vector (with `n` values) for models with one treatment effect parameter, a matrix (`n` rows and same number of columns as number of parameters) otherwise.

References

Riley, Richard D., Julian P. T. Higgins, and Jonathan J. Deeks. "Interpretation of Random Effects Meta-Analyses". *BMJ* 342 (10 February 2011)..

See Also

[treatment_effect](#) returns samples of hypermean and hyper-SD which are used by this function

effect_plot

Plot predictive draws from baggr model

Description

This function plots values from [effect_draw](#), the predictive distribution (under default settings, *posterior* predictive), for one or more [baggr](#) objects.

Usage

```
effect_plot(..., transform = NULL)
```

Arguments

| | |
|-----------|---|
| ... | Object(s) of class baggr . If there is more than one, a comparison will be plotted and names of objects will be used as a plot legend (see examples). |
| transform | a transformation to apply to the result, should be an R function; (this is commonly used when calling <code>group_effects</code> from other plotting or printing functions) |

Details

Under default settings in `baggr` posterior predictive is obtained. But `effect_plot` can also be used for *prior* predictive distributions when setting `ppd=T` in `baggr`. The two outputs work exactly the same, but labels will change to indicate this difference.

Value

A ggplot object.

See Also

`effect_draw` documents the process of drawing values; `baggr_compare` can be used as a shortcut for `effect_plot` with argument `compare = "effects"`

Examples

```
# A single effects plot
bg1 <- baggr(schools, prior_hypersd = uniform(0, 20))
effect_plot(bg1)

# Compare how posterior depends on the prior choice
bg2 <- baggr(schools, prior_hypersd = normal(0, 5))
effect_plot("Uniform prior on SD"=bg1,
           "Normal prior on SD"=bg2)

# Compare the priors themselves (ppd=T)
bg1_ppd <- baggr(schools, prior_hypersd = uniform(0, 20), ppd=TRUE)
bg2_ppd <- baggr(schools, prior_hypersd = normal(0, 5), ppd=TRUE)
effect_plot("Uniform prior on SD"=bg1_ppd,
           "Normal prior on SD"=bg2_ppd)
```

fixed_effects

Effects of covariates on outcome in baggr models

Description

Effects of covariates on outcome in baggr models

Usage

```
fixed_effects(bg, summary = FALSE, transform = NULL, interval = 0.95)
```

Arguments

| | |
|-----------|--|
| bg | a baggr model |
| summary | logical; if TRUE returns summary statistic instead of all MCMC samples |
| transform | a transformation (R function) to apply to the result; (this is commonly used when calling from other plotting or printing functions) |
| interval | uncertainty interval width (numeric between 0 and 1), if summary=TRUE |

Value

A list with 2 vectors (corresponding to MCMC samples) tau (mean effect) and sigma_tau (SD). If summary=TRUE, both vectors are summarised as mean and lower/upper bounds according to interval

See Also

[treatment_effect](#) for overall treatment effect across groups, [group_effects](#) for effects within each group, [effect_draw](#) and [effect_plot](#) for predicted treatment effect in new group

forest_plot

Draw a forest plot for a baggr model

Description

The forest plot functionality in *baggr* is a simple interface for calling the [forestplot](#) function. By default the forest plot displays raw (unpooled) estimates for groups and the treatment effect estimate underneath. This behaviour can be modified to display pooled group estimates.

Usage

```
forest_plot(
  bg,
  show = c("inputs", "posterior", "both", "covariates"),
  print = show,
  prob = 0.95,
  digits = 3,
  ...
)
```

Arguments

| | |
|-------|---|
| bg | a baggr class object |
| show | if "inputs", then plotted points and lines correspond to raw inputs for each group; if "posterior" – to posterior distribution; you can also plot "both" inputs and posteriors; if "covariates", then fixed effect coefficients are plotted |
| print | which values to print next to the plot: values of "inputs" or "posterior" means? (if show="covariates", it must be "posterior") |

| | |
|--------|---|
| prob | width of the intervals (lines) for the plot |
| digits | number of digits to display when printing out mean and SD in the plot |
| ... | other arguments passed to forestplot |

See Also

[forestplot](#) function and its vignette for examples; [effect_plot](#) and [baggr_plot](#) for non-forest plots of baggr results

Examples

```
bg <- baggr(schools, iter = 500)
forest_plot(bg)
forest_plot(bg, show = "posterior", print = "inputs", digits = 2)
```

| | |
|---------------|--|
| get_n_samples | <i>Extract number of samples from a baggr object</i> |
|---------------|--|

Description

Extract number of samples from a baggr object

Usage

```
get_n_samples(x)
```

Arguments

| | |
|---|-------------------------------|
| x | baggr fit to get samples from |
|---|-------------------------------|

Details

Checks for number of iterations and number of Markov chains, returns maximum number of valid samples

| | |
|-----------|--|
| get_order | <i>Separate out ordering so we can test directly</i> |
|-----------|--|

Description

Separate out ordering so we can test directly

Usage

```
get_order(df_groups, hyper)
```

Arguments

| | |
|-----------|--|
| df_groups | data.frame of group effects used in plot.baggr_compare |
| hyper | show parameter estimate? same as in plot.baggr_compare |

Details

Given a set of effects measured by models, identifies the model which has the biggest range of estimates and ranks groups by those estimates, returning the order

| | |
|---------------|------------------------------------|
| group_effects | <i>Extract baggr study effects</i> |
|---------------|------------------------------------|

Description

Given a baggr object, returns the raw MCMC draws of the posterior for each group's effect or a summary of these draws. If there are no covariates in the model, this effect is a single random variable. If there are covariates, the group effect is a sum of effect of covariates (fixed effects) and the study-specific random variable (random effects). This is an internal function currently used as a helper for plotting and printing of results.

Usage

```
group_effects(  
  bg,  
  summary = FALSE,  
  transform = NULL,  
  interval = 0.95,  
  random_only = FALSE  
)
```

Arguments

| | |
|-------------|--|
| bg | baggr object |
| summary | logical; if TRUE returns summary statistics as explained below. |
| transform | a transformation to apply to the result, should be an R function; (this is commonly used when calling group_effects from other plotting or printing functions) |
| interval | uncertainty interval width (numeric between 0 and 1), if summarising |
| random_only | logical; for meta-regression models, should fixed_effects be included in the returned group effect? |

Details

If `summary = TRUE`, the returned object contains, for each study or group, the following 5 values: the posterior medians, the lower and upper bounds of the uncertainty intervals using the central posterior credible interval of width specified in the argument `interval`, the posterior mean, and the posterior standard deviation.

Value

Either an array with MCMC samples (if `summary = FALSE`) or a summary of these samples (if `summary = TRUE`). For arrays the three dimensions are: N samples, N groups and N effects (equal to 1 for the basic models).

See Also

[fixed_effects](#) for effects of covariates on outcome. To extract random effects when covariates are present, you can use either [random_effects](#) or, equivalently, `group_effects(random_only=TRUE)`.

Examples

```
fit1 <- baggr(schools)
group_effects(fit1, summary = TRUE, interval = 0.5)
```

is.baggr_cv

Check if something is a baggr_cv object

Description

Check if something is a baggr_cv object

Usage

```
is.baggr_cv(x)
```

Arguments

| | |
|---|-----------------|
| x | object to check |
|---|-----------------|

labbe

L'Abbe plot for binary data

Description

This plot shows relationship between proportions of events in control and treatment groups in binary data.

Usage

```
labbe(  
  data,  
  group = "group",  
  plot_model = FALSE,  
  labels = TRUE,  
  shade_se = c("rr", "or", "none")  
)
```

Arguments

| | |
|------------|--|
| data | a data frame with binary data (must have columns a, c, b/n1, d/n2) |
| group | a character string specifying group names (e.g. study names), used for labels; |
| plot_model | if TRUE, then odds ratios and risk ratios baggr models are estimated (using default settings) and their mean estimates of effects are plotted as lines |
| labels | if TRUE, names from the group column are displayed |
| shade_se | if "none", nothing is plotted, if "or" or "rr", a shaded area corresponding to inverse of effect's (OR or RR) SE is added to each data point; the default is "rr" |

Value

A ggplot object

See Also

`vignette("baggr_binary")` for an illustrative example

loocv *Leave one group out cross-validation for baggr models*

Description

Performs exact leave-one-group-out cross-validation on a baggr model.

Usage

```
loocv(data, return_models = FALSE, ...)
```

Arguments

| | |
|---------------|---|
| data | Input data frame - same as for baggr function. |
| return_models | logical; if FALSE, summary statistics will be returned and the models discarded; if TRUE, a list of models will be returned alongside summaries |
| ... | Additional arguments passed to baggr . |

Details

The values returned by `loocv()` can be used to understand how excluding any one group affects the overall result, as well as how well the model predicts the omitted group. LOO-CV approaches are a good general practice for comparing Bayesian models, not only in meta-analysis.

This function automatically runs K baggr models, where K is number of groups (e.g. studies), leaving out one group at a time. For each run, it calculates *expected log predictive density* (ELPD) for that group (see Gelman et al 2013). (In the logistic model, where the proportion in control group is unknown, each of the groups is divided into data for controls, which is kept for estimation, and data for treated units, which is not used for estimation but only for calculating predictive density. This is akin to fixing the baseline risk and only trying to infer the odds ratio.)

The main output is the cross-validation information criterion, or -2 times the ELPD averaged over K models. This is related to, and often approximated by, the Watanabe-Akaike Information Criterion. When comparing models, smaller values mean a better fit. For more information on cross-validation see [this overview article](#)

For running more computation-intensive models, consider setting the `mc.cores` option before running `loocv`, e.g. `options(mc.cores = 4)` (by default baggr runs 4 MCMC chains in parallel). As a default, rstan runs "silently" (`refresh=0`). To see sampling progress, please set e.g. `loocv(data, refresh = 500)`.

Value

log predictive density value, an object of class `baggr_cv`; full model, prior values and *lpd* of each model are also returned. These can be examined by using `attributes()` function.

Author(s)

Witold Wiecek

References

Gelman, Andrew, Jessica Hwang, and Aki Vehtari. “Understanding Predictive Information Criteria for Bayesian Models.” *Statistics and Computing* 24, no. 6 (November 2014): 997–1016.

See Also

[loo_compare](#) for comparison of many LOO CV results

Examples

```
## Not run:
# even simple examples may take a while
cv <- loocv(schools, pooling = "partial")
print(cv)      # returns the lpd value
attributes(cv) # more information is included in the object

## End(Not run)
```

 loo_compare

Compare LOO CV models

Description

Given multiple [loocv](#) outputs, calculate differences in their expected log predictive density.

Usage

```
loo_compare(x, ...)
```

Arguments

`x` An object of class `baggr_cv` or a list of such objects.
`...` Additional objects of class `"baggr_cv"`

See Also

[loocv](#) for fitting LOO CV objects and explanation of the procedure

Examples

```
## Not run:
# 2 models with more/less informative priors -- this will take a while to run
cv_1 <- loocv(schools, model = "rubin", pooling = "partial")
cv_2 <- loocv(schools, model = "rubin", pooling = "partial",
             prior_hypermean = normal(0, 5), prior_hypersd = cauchy(0,4))
loo_compare(cv_1, cv_2)

## End(Not run)
```

| | |
|-------------|--|
| microcredit | <i>7 studies on effect of microcredit supply</i> |
|-------------|--|

Description

This dataframe contains the data used in Meager (2019) to estimate hierarchical models on the data from 7 randomized controlled trials of expanding access to microcredit.

Usage

```
microcredit
```

Format

A data frame with 40267 rows, 7 study identifiers and 7 outcomes

Details

The columns include the group indicator which gives the name of the lead author on each of the respective studies, the value of the 6 outcome variables of most interest (consumer durables spending, business expenditures, business profit, business revenues, temptation goods spending and consumption spending) all of which are standardised to USD PPP in 2009 dollars per two weeks (these are flow variables), and finally a treatment assignment status indicator.

The dataset has not otherwise been cleaned and therefore includes NAs and other issues common to real-world datasets.

For more information on how and why these variables were chosen and standardised, see Meager (2019) or consult the associated code repository which includes the standardisation scripts: [link](#)

References

Meager, Rachael (2019) Understanding the average impact of microcredit expansions: A Bayesian hierarchical analysis of seven randomized experiments. *American Economic Journal: Applied Economics*, 11(1), 57-91.

| | |
|------------------------|---|
| microcredit_simplified | <i>Simplified version of the microcredit dataset.</i> |
|------------------------|---|

Description

This dataframe contains the data used in Meager (2019) to estimate hierarchical models on the data from 7 randomized controlled trials of expanding access to microcredit.

Usage

```
microcredit_simplified
```

Format

A data frame with 14224 rows, 7 study identifiers and 1 outcome

Details

The columns include the group indicator which gives the name of the lead author on each of the respective studies, the value of the household consumption spending standardised to USD PPP in 2009 dollars per two weeks (these are flow variables), and finally a treatment assignment status indicator.

The dataset has not otherwise been cleaned and therefore includes NAs and other issues common to real data.

For more information on how and why these variables were chosen and standardised, see Meager (2019) or consult the associated code repository: [link](#)

This dataset includes only complete cases and only the consumption outcome variable.

References

Meager, Rachael (2019) Understanding the average impact of microcredit expansions: A Bayesian hierarchical analysis of seven randomized experiments. *American Economic Journal: Applied Economics*, 11(1), 57-91.

| | |
|------|---|
| mint | <i>"Mean and interval" function, including other summaries, calculated for matrix (by column) or vector</i> |
|------|---|

Description

This function is just a convenient shorthand for getting typical summary statistics.

Usage

```
mint(y, int = 0.95, digits = NULL, median = FALSE, sd = FALSE)
```

Arguments

| | |
|--------|--|
| y | matrix or a vector; for matrices, mint is done by-column |
| int | probability interval (default is 95 percent) to calculate |
| digits | number of significant digits to round values by. |
| median | return median value? |
| sd | return SD? |

Examples

```
mint(rnorm(100, 12, 5))
```

| | |
|-----------|--|
| mutau_cor | <i>Correlation between mu and tau in a baggr model</i> |
|-----------|--|

Description

Correlation between mu and tau in a baggr model

Usage

```
mutau_cor(bg, summary = FALSE, interval = 0.95)
```

Arguments

| | |
|----------|--|
| bg | a baggr model where model = "mutau" |
| summary | logical; if TRUE returns summary statistics as explained below. |
| interval | uncertainty interval width (numeric between 0 and 1), if summarising |

Value

a vector of values

| | |
|------------|---------------------------------------|
| plot.baggr | <i>Generic plot for baggr package</i> |
|------------|---------------------------------------|

Description

Using generic plot() on baggr output invokes [baggr_plot](#) visual. See therein for customisation options. Note that plot output is ggplot2 object.

Usage

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

Arguments

| | |
|-----|------------------------------------|
| x | object of class baggr |
| ... | optional arguments, see baggr_plot |

Value

ggplot2 object from baggr_plot

Author(s)

Witold Wiecek

plot.baggr_compare *Plot method for baggr_compare models*

Description

Allows plots that compare multiple baggr models that were passed for comparison purposes to baggr compare or run automatically by baggr_compare

Usage

```
## S3 method for class 'baggr_compare'
plot(
  x,
  compare = x$compare,
  style = "areas",
  grid_models = FALSE,
  grid_parameters = TRUE,
  interval = 0.95,
  hyper = TRUE,
  transform = NULL,
  order = F,
  vline = FALSE,
  ...
)
```

Arguments

| | |
|-----------------|---|
| x | baggr_compare model to plot |
| compare | When plotting, choose between comparison of "groups" (default) or (hyper-) "effects". The former is not available when what = "prior". |
| style | What kind of plot to display (if grid_models = TRUE), passed to the style argument in baggr_plot . |
| grid_models | If FALSE (default), generate a single comparison plot; if TRUE, display each model (using individual baggr_plot 's) side-by-side. |
| grid_parameters | if TRUE, uses ggplot-style faceting when plotting models with many parameters (especially "quantiles", "sslab"); if FALSE, returns separate plot for each parameter |
| interval | probability level used for display of posterior interval |
| hyper | Whether to plot pooled treatment effect in addition to group treatment effects |
| transform | a function (e.g. exp(), log()) to apply to the values of group (and hyper, if hyper=TRUE) effects before plotting; when working with effects that are on log scale, exponent transform is used automatically, you can plot on log scale by setting transform = identity |

| | |
|-------|--|
| order | Whether to sort by median treatment effect by group. If yes, medians from the model with largest range of estimates are used for sorting. If not, groups are shown alphabetically. |
| vline | logical; show vertical line through 0 in the plot? |
| ... | ignored for now, may be used in the future |

| | |
|----------------|-----------------------|
| plot_quantiles | <i>plot_quantiles</i> |
|----------------|-----------------------|

Description

Plot results for baggr quantile models. Displays results faceted per group. Results are ggplot2 plots and can be modified.

Usage

```
plot_quantiles(fit, ncol, hline = TRUE)
```

Arguments

| | |
|-------|--|
| fit | an object of class baggr |
| ncol | number of columns for the plot; defaults to half of number of groups |
| hline | logical; plots a line through 0 |

Value

ggplot2 object

Examples

```
## Not run:
bg <- baggr(microcredit_simplified, model = "quantiles",
            quantiles = c(0.25, 0.50, 0.75),
            iter = 1000, refresh = 0,
            outcome = "consumption")
#vanilla plot
plot_quantiles(bg)[[1]]
plot_quantiles(bg, hline = TRUE)[[2]] +
  ggplot2::coord_cartesian(ylim = c(-2, 5)) +
  ggplot2::ggtitle("Works like a ggplot2 plot!")

## End(Not run)
```

| | |
|---------|----------------------------------|
| pooling | <i>Pooling metrics for baggr</i> |
|---------|----------------------------------|

Description

Compute statistics relating to heterogeneity (whole model) and pooling (for each group) given a [baggr](#) meta-analysis model. The statistics are the pooling metric by Gelman & Pardoe (2006) or its complement, the *I-squared* statistic.

Usage

```
pooling(bg, type = c("groups", "total"), summary = TRUE)
```

```
heterogeneity(bg, summary = TRUE)
```

Arguments

| | |
|---------|--|
| bg | output of a baggr() function |
| type | In pooling calculation is done for each of the "groups" (default) or for "total" hypereffect(s). See <i>Details</i> section for how calculation is done. |
| summary | logical; if FALSE a whole vector of pooling values is returned, otherwise only the means and intervals |

Details

Pooling statistic describes the extent to which group-level estimates of treatment effect are "pooled" (or pulled!) closer to average treatment effect in the meta-analysis model. If pooling = "none" or "full" in [baggr](#), then the values are always 0 or 1, respectively. If pooling = "partial", the value is somewhere between 0 and 1.

Formulae for the calculations below are provided in main package vignette.

Value

Matrix with mean and intervals for chosen pooling metric, each row corresponding to one meta-analysis group.

Group pooling

This is the calculation done by pooling() if type = "groups" (default). See vignette("baggr") for more details on pooling calculations.

In a partial pooling model (see [baggr](#)), group k (e.g. study) has standard error of treatment effect estimate, se_k . The treatment effect (across k groups) is variable across groups, with hyper-SD parameter $\sigma(\tau)$.

The quantity of interest is ratio of variation in treatment effects to the total variation. By convention, we subtract it from 1, to obtain a *pooling metric* p .

$$p = 1 - (\sigma(\tau))^2 / (\sigma(\tau))^2 + se_k^2$$

- If $p < 0.5$, the variation across studies is higher than variation within studies.
- Values close to 1 indicate nearly full pooling. Variation across studies dominates.
- Values close to 0 indicate no pooling. Variation within studies dominates.

Note that, since σ_τ^2 is a Bayesian parameter (rather than a single fixed value), p is also a parameter. It is typical for p to have very high dispersion, as in many cases we cannot precisely estimate σ_τ . To obtain the whole distribution of `_p_` (rather than summarised values), set `summary=FALSE`.

Overall pooling in the model

Typically researchers want to report a single measure from the model, relating to heterogeneity across groups. This is calculated by either `pooling(mymodel, type = "total")` or simply `heterogeneity(mymodel)`

In many contexts, i.e. medical statistics, it is typical to report I^2 , called I^2 (see Higgins and Thompson, 2002; sometimes another statistic, $H^2 = 1/P$, is used). Higher values of I^2 indicate higher heterogeneity; Von Hippel (2015) provides useful details for I^2 calculations.

To obtain such single estimate we need to substitute average variability of group-specific treatment effects and then calculate the same way we would calculate p . By default we use the mean across k se_k^2 values. Typically, implementations of I^2 in statistical packages use a different calculation for this quantity, which may make I^2 's not comparable when different studies have different SE's.

Same as for group-specific estimates, P is a Bayesian parameter and its dispersion can be high.

Relationship to R-squared statistic

See Gelman & Pardoe (2006) Section 1.1 for a short explanation of how R^2 statistic relates to the pooling metric.

References

Gelman, Andrew, and Iain Pardoe. "Bayesian Measures of Explained Variance and Pooling in Multilevel (Hierarchical) Models." *Technometrics* 48, no. 2 (May 2006): 241-51.

Higgins, Julian P. T., and Simon G. Thompson. "Quantifying Heterogeneity in a Meta-Analysis." *Statistics in Medicine*, vol. 21, no. 11, June 2002, pp. 1539–58.

Hippel, Paul T von. "The Heterogeneity Statistic I^2 Can Be Biased in Small Meta-Analyses." *BMC Medical Research Methodology* 15 (April 14, 2015).

pp_check.baggr

Posterior predictive checks for baggr model

Description

Performs posterior predictive checks with the **bayesplot** package.

Usage

```
## S3 method for class 'baggr'
pp_check(x, type = "dens_overlay", nsamples = 40)
```


Arguments

| | |
|----------|---|
| x | Model to check |
| type | type of pp_check. For a list see here . |
| nsamples | number of samples to compare |

Details

For a detailed explanation of each of the ppc functions, see the [PPC](#) documentation of the bayesplot package.

| | |
|---------------|---|
| predict.baggr | <i>Predict method for baggr objects</i> |
|---------------|---|

Description

Predict method for baggr objects

Usage

```
## S3 method for class 'baggr'
predict(object, nsamples, newdata = NULL, allow_new_levels = TRUE, ...)
```

Arguments

| | |
|------------------|---|
| object | model to predict from |
| nsamples | Number of samples to draw from the posterior. Cannot exceed the number of samples in the fitted model. |
| newdata | optional, new data to predict observations from |
| allow_new_levels | whether to allow the model to make predictions about unobserved groups. Without additional group-level information the model will use the unconditional, pooled estimate. |
| ... | other arguments to pass to predict function (currently not used) |

| | |
|---------------|--|
| predict_mutau | <i>Predict function for the mu & tau model</i> |
|---------------|--|

Description

Predict function for the mu & tau model

Usage

```
predict_mutau(x, nsamples, newdata = NULL, allow_new_levels = TRUE)
```

Arguments

| | |
|------------------|--|
| x | model to predict from |
| nsamples | number of samples to predict |
| newdata | new data to predict, defaults to NULL |
| allow_new_levels | allow the predictive of new, unobserved groups |

| | |
|-------------------|---|
| predict_quantiles | <i>Predict function for the quantiles model</i> |
|-------------------|---|

Description

Predict function for the quantiles model

Usage

```
predict_quantiles(x, nsamples, newdata = NULL, allow_new_levels = TRUE)
```

Arguments

| | |
|------------------|--|
| x | model to predict from |
| nsamples | number of samples to predict |
| newdata | new data to predict, defaults to NULL |
| allow_new_levels | allow the predictive of new, unobserved groups |

| | |
|---------------|---|
| predict_rubin | <i>Predict function for the rubin model</i> |
|---------------|---|

Description

Predict function for the rubin model

Usage

```
predict_rubin(x, nsamples, newdata = NULL, allow_new_levels = TRUE)
```

Arguments

| | |
|------------------|--|
| x | model to predict from |
| nsamples | number of samples to predict |
| newdata | new data to predict, defaults to NULL |
| allow_new_levels | allow the predictive of new, unobserved groups |

| | |
|-----------------|--|
| predict_unknown | <i>Predict method for model that is unknown or not implemented</i> |
|-----------------|--|

Description

Predict method for model that is unknown or not implemented

Usage

```
predict_unknown(x)
```

Arguments

| | |
|---|--|
| x | baggr model to generate predictions from |
|---|--|

prepare_ma

*Convert from individual to summary data in meta-analyses***Description**

Allows for one-way conversion from full to summary data or for calculation of effects for binary data. Input must be pre-formatted appropriately.

Usage

```
prepare_ma(
  data,
  effect = c("mean", "logOR", "logRR"),
  rare_event_correction = 0.25,
  log = FALSE,
  cfb = FALSE,
  summarise = TRUE,
  treatment = "treatment",
  baseline = NULL,
  group = "group",
  outcome = "outcome"
)
```

Arguments

| | |
|-----------------------|--|
| data | either a data.frame of individual-level observations with columns for outcome (numeric), treatment (values 0 and 1) and group (numeric, character or factor); or , a data frame with binary data (must have columns a, c, b/n1, d/n2). |
| effect | what effect to calculate? a mean (and SE) of outcome in groups or (for binary data) logOR (odds ratio), logRR (risk ratio); |
| rare_event_correction | If effect is logOR or logRR, this correction is used when working with binary data only. The value of correction is added to all arms in trials where some arms had 0 events. Using corrections may bias results but is the only alternative to avoid infinite values. |
| log | logical; log-transform the outcome variable? |
| cfb | logical; calculate change from baseline? If yes, the outcome variable is taken as a difference between values in outcome and baseline columns |
| summarise | logical; TRUE by default, but you can disable it to obtain converted (e.g. logged) data with columns renamed |
| treatment | name of column with treatment variable |
| baseline | name of column with baseline variable |
| group | name of the column with grouping variable |
| outcome | name of column with outcome variable |

Details

The conversions done by this function are not typically needed and may happen automatically when data is given to [baggr](#). However, this function can be used to explicitly convert from full to reduced (summarised) data without analysing it in any model. It can be useful for examining your data and generating summary tables.

If multiple operations are performed, they are taken in this order:

1. conversion to log scale,
2. calculating change from baseline,
3. summarising data (using appropriate effect)

Value

- If you summarise: a data.frame with columns for group, tau and se.tau (for effect = "mean", also baseline means; for "logRR" or "logOR" also a, b, c, d, which correspond to typical contingency table notation, that is: a = events in exposed; b = no events in exposed, c = events in unexposed, d = no events in unexposed).
- If you do not summarise data, individual level data will be returned, but some columns may be renamed or transformed (see the arguments above).

Author(s)

Witold Wiecek

See Also

[convert_inputs](#) for how any type of data is (internally) converted into a list of Stan inputs;

prepare_prior

Prepare prior values for Stan models in baggr

Description

This is an internal function called by [baggr](#). You can use it for debugging or to run modified models. It extracts and prepares priors passed by the user. Then, if any necessary priors are missing, it sets them automatically and notifies user about these automatic choices.

Usage

```
prepare_prior(  
  prior,  
  data,  
  stan_data,  
  model,  
  pooling,  
  covariates,
```

```

    quantiles = c(),
    silent = FALSE
  )

```

Arguments

| | |
|------------|---|
| prior | prior argument passed from baggr call |
| data | data another argument in baggr |
| stan_data | list of inputs that will be used by sampler this is already pre-obtained through convert_inputs |
| model | same as in baggr |
| pooling | same as in baggr |
| covariates | same as in baggr |
| quantiles | same as in baggr |
| silent | same as in baggr |

Value

A named list with prior values that can be appended to `stan_data` and passed to a Stan model.

| | |
|-------------|--|
| print.baggr | <i>S3 print method for objects of class baggr (model fits)</i> |
|-------------|--|

Description

This prints a concise summary of the main [baggr](#) model features. More info is included in the summary of the model and its attributes.

Usage

```

## S3 method for class 'baggr'
print(x, exponent = FALSE, digits = 2, group, fixed = TRUE, ...)

```

Arguments

| | |
|----------|---|
| x | object of class baggr |
| exponent | if TRUE, results (for means) are converted to exp scale |
| digits | Number of significant digits to print. |
| group | logical; print group effects? If unspecified, they are printed only if less than 20 groups are present |
| fixed | logical: print fixed effects? |
| ... | currently unused by this package: further arguments passed to or from other methods (print requirement) |

print.baggr_compare *Print method for baggr_compare models*

Description

Print method for baggr_compare models

Usage

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

Arguments

| | |
|--------|---|
| x | baggr_compare model |
| digits | number of significant digits for effect estimates |
| ... | other parameters passed to print |

print.baggr_cv *Print baggr cv objects nicely*

Description

Print baggr cv objects nicely

Usage

```
## S3 method for class 'baggr_cv'  
print(x, digits = 3, ...)
```

Arguments

| | |
|--------|---|
| x | baggr_cv object to print |
| digits | number of digits to print |
| ... | additional arguments for s3 consistency |

```
print.compare_baggr_cv
```

Print baggr_cv comparisons

Description

Print baggr_cv comparisons

Usage

```
## S3 method for class 'compare_baggr_cv'  
print(x, digits = 3, ...)
```

Arguments

| | |
|--------|---|
| x | baggr_cv comparison to print |
| digits | number of digits to print |
| ... | additional arguments for s3 consistency |

```
print_dist
```

Output a distribution as a string

Description

Used for printing nicely formatted outputs when reporting results etc.

Usage

```
print_dist(dist)
```

Arguments

| | |
|------|--|
| dist | distribution name, one of priors |
|------|--|

Value

Character string like normal(0, 10^2).

priors

Prior distributions in baggr

Description

This page provides a list of all available distributions that can be used to specify priors in `baggr()`. These convenience functions are designed to allow the user to write the priors in the most "natural" way when implementing them in `baggr`. Apart from passing on the arguments, their only other role is to perform a rudimentary check if the distribution is specified correctly.

Usage

```
multinormal(location, Sigma)
```

```
lkj(shape, order = NULL)
```

```
normal(location, scale)
```

```
cauchy(location, scale)
```

```
uniform(lower, upper)
```

Arguments

| | |
|----------|--|
| location | Mean for normal and multivariate normal (in which case <code>location</code> is a vector), and median for Cauchy distributions |
| Sigma | Variance-covariance matrix for multivariate normal. |
| shape | Shape parameter for LKJ |
| order | Order of LKJ matrix (typically it does not need to be specified, as it is inferred directly in the model) |
| scale | SD for Normal, scale for Cauchy |
| lower | Lower bound for Uniform |
| upper | Upper bound for Uniform |

Details

The prior choice in `baggr` is always done via 3 distinct arguments: `prior_hypermean`, `prior_hypersd`, and `prior_hypercor`.

These respectively refer to the priors on the average of the effects across the groups (`hypermean`), the standard deviation of the effects across the groups (`hypersd`), and the correlation in the distribution of parameters across groups when the model allows multivariate shrinkage (say on control group means and effects).

Notation for priors is "plain-text", in that you can write the distributions as `normal(5, 10)`, `uniform(0, 100)` etc. As with any other argument one has the option to simply input the prior directly, e.g. `prior_hypermean`

= $\text{normal}(0, 1)$, or by creating a named list of custom priors and then inputting the list to the argument priors. See the examples below for more.

Different parameters admit different priors:

- prior_hypermean will take "normal", "uniform" and "cauchy" input for a scalar mean. For a vector mean, it will take any of these arguments and apply them independently to each component of the vector, or it can also take a "multinormal" argument (see the example below).
- prior_hypersd will take "normal" and "uniform"
- prior_hypercor allows "lkj" input

Author(s)

Witold Wiecek, Rachael Meager

References

Lewandowski, Daniel, Dorota Kurowicka, and Harry Joe. "Generating Random Correlation Matrices Based on Vines and Extended Onion Method." *Journal of Multivariate Analysis* 100, no. 9 (October 1, 2009): 1989-2001.

Examples

```
# (these are not the recommended priors -- for syntax illustration only)

# change the priors for 8 schools:
baggr(schools, model = "rubin", pooling = "partial",
      prior_hypermean = normal(5,5),
      prior_hypersd = normal(0,20))

# passing priors as a list
custom_priors <- list(hypercor = lkj(1), hypersd = normal(0,10),
                    hypermean = multinormal(c(0,0),matrix(c(10,3,3,10),2,2)))
microcredit_summary_data <- prepare_ma(microcredit, outcome = "consumption")
baggr(microcredit_summary_data, model = "mutau",
      pooling = "partial", prior = custom_priors)
```

random_effects

Extract random effects from a baggr model

Description

This is a shortcut for writing `group_effects(random_only=TRUE, ...)`

Usage

```
random_effects(...)
```

Arguments

... arguments passed to [group_effects](#)

| | |
|------------|---|
| rubin_data | <i>Make model matrix for the rubin data</i> |
|------------|---|

Description

Make model matrix for the rubin data

Usage

```
rubin_data(x, newdata = NULL, allow_new_levels = TRUE)
```

Arguments

| | |
|------------------|--|
| x | model to get data from |
| newdata | new data to use with model |
| allow_new_levels | whether to allow for unobserved groups |

| | |
|---------|--------------------------|
| schools | <i>8 schools example</i> |
|---------|--------------------------|

Description

A classic example of aggregate level continuous data in Bayesian hierarchical modelling. This dataframe contains a column of estimated treatment effects of an SAT prep program implemented in 8 different schools in the US, and a column of estimated standard errors.

Usage

```
schools
```

Format

An object of class `data.frame` with 8 rows and 3 columns.

Details

See Gelman et al (1995), Chapter 5, for context and applied example.

References

Gelman, Andrew, John B. Carlin, Hal S. Stern, and Donald B. Rubin. Bayesian Data Analysis. Taylor & Francis, 1995.

| | |
|---------------|---|
| set_prior_val | <i>Add prior values to Stan input for baggr</i> |
|---------------|---|

Description

Add prior values to Stan input for baggr

Usage

```
set_prior_val(target, name, prior, p = 1)
```

Arguments

| | |
|--------|--|
| target | list object (Stan input) to which prior will be added |
| name | prior name, like hypermean, hypersd, hypercor |
| prior | one of prior distributions allowed by baggr like normal |
| p | number of repeats of the prior, i.e. when P i.i.d. priors are set for P dimensional parameter as in "mu & tau" type of model |

| | |
|------------------|---|
| single_comp_plot | <i>Plot single comparison plot in baggr_compare style</i> |
|------------------|---|

Description

Plot single comparison plot in baggr_compare style

Usage

```
single_comp_plot(df, title = "", legend = "top", ylab = "", grid = F)
```

Arguments

| | |
|--------|---|
| df | data.frame with columns 'group', 'median', 'lci', 'uci', 'model' and optionally 'parameter' |
| title | 'ggtitle' |
| legend | 'legend.position' |
| ylab | Y axis label |
| grid | logical; if TRUE, facets by 'parameter' column |

Value

a ggplot2 object

stop_not_implemented *Stop with informative error*

Description

Stop with informative error

Usage

```
stop_not_implemented()
```

summarise_quantiles_data

summarise_quantiles_data

Description

Given individual level data, return list of summary statistics Of quantile means and Sigma's, as well as K, N. This operation is done automatically inside [baggr](#) function but it can be used for exploring input data.

Usage

```
summarise_quantiles_data(
  data,
  quantiles,
  outcome = "outcome",
  group = "group",
  treatment = "treatment",
  means_only = FALSE
)
```

Arguments

| | |
|------------|--|
| data | a data.frame with grouping, outcome, treatment variables |
| quantiles | a vector of quantiles (between 0 and 1) |
| outcome | character: column name in data for outcome variable |
| group | character: column name in data for grouping variable |
| treatment | character: column name in data for treatment variable |
| means_only | logical; if true, covariances and extra outputs are not returned, but only means (y_0 and y_1) |

Details

This function is intended for data exploration outside of `baggr` function. When calling `baggr`, individual-level data should be supplied – they will be summarised automatically. (See `baggr` for documentation of how columns should be formatted.)

Estimates are obtained externally via `quantreg::rq` function of the `quantreg` package. This implementation is experimental and will change in the future versions of the package.

Value

A list with mean estimates for control and treatment (y_{θ} and y_1) and variance-covariance matrices ($\text{Sigma}_{y_k\theta}$ and Sigma_{y_k1}).

Examples

```
summarise_quantiles_data(microcredit_simplified, c(.2, .4, .6),
  outcome = "consumption")
```

| | |
|------------------|--|
| treatment_effect | <i>Average treatment effect in a baggr model</i> |
|------------------|--|

Description

Average treatment effect in a baggr model

Usage

```
treatment_effect(
  bg,
  summary = FALSE,
  transform = NULL,
  interval = 0.95,
  message = TRUE
)
```

Arguments

| | |
|------------------------|--|
| <code>bg</code> | a <code>baggr</code> model |
| <code>summary</code> | logical; if TRUE returns summary statistics as explained below. |
| <code>transform</code> | a transformation to apply to the result, should be an R function; (this is commonly used when calling <code>treatment_effect</code> from other plotting or printing functions) |
| <code>interval</code> | uncertainty interval width (numeric between 0 and 1), if summarising |
| <code>message</code> | logical; use to disable messages prompted by using with no pooling models |

Value

A list with 2 vectors (corresponding to MCMC samples) tau (mean effect) and sigma_tau (SD).
If summary=TRUE, both vectors are summarised as mean and lower/upper bounds according to interval

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