

Package ‘historicalborrow’

May 8, 2026

Title Non-Longitudinal Bayesian Historical Borrowing Models

Description Historical borrowing in clinical trials can improve precision and operating characteristics. This package supports a hierarchical model and a mixture model to borrow historical control data from other studies to better characterize the control response of the current study. It also quantifies the amount of borrowing through benchmark models (independent and pooled). Some of the methods are discussed by Viele et al. (2013) <[doi:10.1002/pst.1589](https://doi.org/10.1002/pst.1589)>.

Version 1.1.1

License MIT + file LICENSE

URL <https://wlandau.github.io/historicalborrow/>,
<https://github.com/wlandau/historicalborrow>

BugReports <https://github.com/wlandau/historicalborrow/issues>

Depends R (>= 3.5.0)

Imports dplyr, ggplot2, Matrix, posterior, rjags, rlang, stats,
tibble, tidyr, tidyselect, utils

Suggests knitr, markdown, rmarkdown, testthat (>= 3.0.0)

Encoding UTF-8

Language en-US

VignetteBuilder knitr

Config/testthat/edition 3

RoxygenNote 7.3.2

NeedsCompilation no

Author William Michael Landau [aut, cre] (ORCID:
<<https://orcid.org/0000-0003-1878-3253>>),
Albert Man [rev],
Eli Lilly and Company [cph]

Maintainer William Michael Landau <will.landau.oss@gmail.com>

Repository CRAN

Date/Publication 2026-02-09 15:50:02 UTC

Contents

historicalborrow-package	2
hb_convergence	2
hb_data	3
hb_ess	5
hb_mcmc_hierarchical	7
hb_mcmc_independent	10
hb_mcmc_mixture	12
hb_mcmc_mixture_hyperparameters	14
hb_mcmc_pool	17
hb_plot_borrow	19
hb_plot_group	20
hb_plot_tau	22
hb_sim_hierarchical	22
hb_sim_independent	25
hb_sim_mixture	26
hb_sim_pool	28
hb_summary	30

Index	34
--------------	-----------

historicalborrow-package

historicalborrow: Bayesian historical borrowing models for clinical studies.

Description

Bayesian historical borrowing models for clinical studies.

hb_convergence

Check convergence diagnostics

Description

Check the convergence diagnostics on a model.

Usage

```
hb_convergence(mcmc)
```

Arguments

mcmc	A wide data frame of posterior samples returned by <code>hb_mcmc_hierarchical()</code> or similar MCMC function.
------	--

Value

A data frame of summarized convergence diagnostics. `max_rhat` is the maximum univariate Gelman/Rubin potential scale reduction factor over all the parameters of the model, `min_ess_bulk` is the minimum bulk effective sample size over the parameters, and `min_ess_tail` is the minimum tail effective sample size. `max_rhat` should be below 1.01, and the ESS metrics should both be above 100 times the number of MCMC chains. If any of these conditions are not true, the MCMC did not converge, and it is recommended to try running the model for more saved iterations (and if `max_rhat` is high, possibly more warmup iterations).

See Also

Other mcmc: [hb_mcmc_hierarchical\(\)](#), [hb_mcmc_independent\(\)](#), [hb_mcmc_mixture\(\)](#), [hb_mcmc_mixture_hyperparameter\(\)](#), [hb_mcmc_pool\(\)](#)

Examples

```
data <- hb_sim_pool(n_continuous = 2)$data
mcmc <- hb_mcmc_pool(
  data,
  n_chains = 1,
  n_adapt = 100,
  n_warmup = 200,
  n_iterations = 200
)
hb_convergence(mcmc)
```

hb_data

Standardize data

Description

Standardize a tidy input dataset.

Usage

```
hb_data(
  data,
  response,
  study,
  study_reference,
  group,
  group_reference,
  patient,
  covariates
)
```

Arguments

data	A tidy data frame or tibble with the data.
response	Character of length 1, name of the column in data with the response/outcome variable. <code>data[[response]]</code> must be a continuous variable, and it <i>should</i> be the change from baseline of a clinical endpoint of interest, as opposed to just the raw response. Treatment differences are computed directly from this scale, please supply change from baseline unless you are absolutely certain that treatment differences computed directly from this quantity are clinically meaningful.
study	Character of length 1, name of the column in data with the study ID.
study_reference	Atomic of length 1, element of the study column that indicates the current study. (The other studies are historical studies.)
group	Character of length 1, name of the column in data with the group ID.
group_reference	Atomic of length 1, element of the group column that indicates the control group. (The other groups may be treatment groups.)
patient	Character of length 1, name of the column in data with the patient ID.
covariates	Character vector of column names in data with the columns with baseline covariates. These can be continuous, categorical, or binary. Regardless, <code>historicalborrow</code> derives the appropriate model matrix.

Details

Users do not normally need to call this function. It mainly serves exposes the indexing behavior of studies and group levels to aid in interpreting summary tables.

Value

A standardized tidy data frame with one row per patient and the following columns:

- `response`: continuous response/outcome variable. (Should be change from baseline of an outcome of interest.)
- `study_label`: human-readable label of the study.
- `study`: integer study index with the max index equal to the current study (at `study_reference`).
- `group_label`: human-readable group label (e.g. treatment arm name).
- `group`: integer group index with an index of 1 equal to the control group (at `group_reference`).
- `patient_label`: original patient ID.
- `patient`: integer patient index.
- `covariate_*`: baseline covariate columns.

Examples

```
data <- hb_sim_independent(n_continuous = 1, n_study = 2)$data
data <- dplyr::select(
  data,
  study,
  group,
  patient,
  response,
  tidyselect::everything()
)
colnames(data) <- c("trial", "arm", "subject", "change", "cov1", "cov2")
data$trial <- paste0("trial", data$trial)
data$arm <- paste0("arm", data$arm)
hb_data(
  data = data,
  response = "change",
  study = "trial",
  study_reference = "trial1",
  group = "arm",
  group_reference = "arm1",
  patient = "subject",
  covariates = c("cov1", "cov2")
)
```

hb_ess

Effective sample size (ESS)

Description

Quantify borrowing with effective sample size (ESS) as cited and explained in the methods vignette at <https://wlandau.github.io/historicalborrow/articles/methods.html>.

Usage

```
hb_ess(
  mcmc_pool,
  mcmc_hierarchical,
  data,
  response = "response",
  study = "study",
  study_reference = max(data[[study]]),
  group = "group",
  group_reference = min(data[[group]]),
  patient = "patient"
)
```

Arguments

mcmc_pool	A fitted model from hb_mcmc_pool() .
mcmc_hierarchical	A fitted model from hb_mcmc_hierarchical() .
data	A tidy data frame or tibble with the data.
response	Character of length 1, name of the column in data with the response/outcome variable. <code>data[[response]]</code> must be a continuous variable, and it <i>should</i> be the change from baseline of a clinical endpoint of interest, as opposed to just the raw response. Treatment differences are computed directly from this scale, please supply change from baseline unless you are absolutely certain that treatment differences computed directly from this quantity are clinically meaningful.
study	Character of length 1, name of the column in data with the study ID.
study_reference	Atomic of length 1, element of the study column that indicates the current study. (The other studies are historical studies.)
group	Character of length 1, name of the column in data with the group ID.
group_reference	Atomic of length 1, element of the group column that indicates the control group. (The other groups may be treatment groups.)
patient	Character of length 1, name of the column in data with the patient ID.

Value

A data frame with one row and the following columns:

- v_0 : posterior predictive variance of the control group mean of a hypothetical new study given the pooled model. Calculated as the mean over MCMC samples of $1 / \sum(\sigma_i^2)$, where each σ_i is the residual standard deviation of study i estimated from the pooled model.
- v_{τ} : posterior predictive variance of a hypothetical new control group mean under the hierarchical model. Calculated by averaging over predictive draws, where each predictive draw is from $rnorm(n = 1, mean = \mu_, sd = \tau_)$ and $\mu_$ and $\tau_$ are the μ and τ components of an MCMC sample.
- n : number of non-missing historical control patients.
- $weight$: strength of borrowing as a ratio of variances: v_0 / v_{τ} .
- ess : strength of borrowing as an effective sample size: $n v_0 / v_{\tau}$, where n is the number of non-missing historical control patients.

See Also

Other summary: [hb_summary\(\)](#)

Examples

```

data <- hb_sim_independent(n_continuous = 2)$data
data$group <- sprintf("group%s", data$group)
data$study <- sprintf("study%s", data$study)
pool <- hb_mcmc_pool(
  data,
  n_chains = 1,
  n_adapt = 100,
  n_warmup = 50,
  n_iterations = 50
)
hierarchical <- hb_mcmc_hierarchical(
  data,
  n_chains = 1,
  n_adapt = 100,
  n_warmup = 50,
  n_iterations = 50
)
hb_ess(
  mcmc_pool = pool,
  mcmc_hierarchical = hierarchical,
  data = data
)

```

hb_mcmc_hierarchical *Hierarchical model MCMC*

Description

Run the hierarchical model with MCMC.

Usage

```

hb_mcmc_hierarchical(
  data,
  response = "response",
  study = "study",
  study_reference = max(data[[study]]),
  group = "group",
  group_reference = min(data[[group]]),
  patient = "patient",
  covariates = grep("^covariate", colnames(data), value = TRUE),
  s_delta = 30,
  s_beta = 30,
  s_sigma = 30,
  s_mu = 30,
  s_tau = sd(data[[response]], na.rm = TRUE),
  d_tau = 1,

```

```

prior_tau = "half_t",
n_chains = 4,
n_adapt = 2000,
n_warmup = 4000,
n_iterations = 20000,
quiet = TRUE
)

```

Arguments

data	Tidy data frame with one row per patient, indicator columns for the response variable, study, group, and patient, and covariates. All columns must be atomic vectors (e.g. not lists). The data for the mixture and simple models should have just one study, and the others should have data from more than one study. The simple model can be used to get the historical data components of <code>m_omega</code> and <code>s_omega</code> for the mixture model.
response	Character of length 1, name of the column in data with the response/outcome variable. <code>data[[response]]</code> must be a continuous variable, and it <i>should</i> be the change from baseline of a clinical endpoint of interest, as opposed to just the raw response. Treatment differences are computed directly from this scale, please supply change from baseline unless you are absolutely certain that treatment differences computed directly from this quantity are clinically meaningful.
study	Character of length 1, name of the column in data with the study ID.
study_reference	Atomic of length 1, element of the <code>study</code> column that indicates the current study. (The other studies are historical studies.)
group	Character of length 1, name of the column in data with the group ID.
group_reference	Atomic of length 1, element of the <code>group</code> column that indicates the control group. (The other groups may be treatment groups.)
patient	Character of length 1, name of the column in data with the patient ID.
covariates	Character vector of column names in data with the columns with baseline covariates. These can be continuous, categorical, or binary. Regardless, <code>historicalborrow</code> derives the appropriate model matrix.
s_delta	Numeric of length 1, prior standard deviation of the study-by-group effect parameters <code>delta</code> .
s_beta	Numeric of length 1, prior standard deviation of the fixed effects <code>beta</code> .
s_sigma	Numeric of length 1, prior upper bound of the residual standard deviations.
s_mu	Numeric of length 1, prior standard deviation of <code>mu</code> .
s_tau	Non-negative numeric of length 1. If <code>prior_tau</code> is "half_t", then <code>s_tau</code> is the scale parameter of the Student t prior of <code>tau</code> and analogous to the <code>sigma</code> parameter of the Student-t parameterization given at https://mc-stan.org/docs/functions-reference/unbounded_continuous_distributions.html . # nolint If <code>prior_tau</code> is "uniform", then <code>s_tau</code> is the upper bound of <code>tau</code> . Upper bound on <code>tau</code> if <code>prior_tau</code> is "uniform".

In the case of `prior_tau` equal to "half_t", the defaults `s_tau = sd(data[[response]])`, `na.rm = TRUE`) and `d_tau = 1` specify a weakly informative scaled half-Cauchy distribution. This choice is only provisional. The prior on `tau` is extremely important, especially for small numbers of historical studies, and the user should set a reasonable value for the use case, ideally informed by the results of sensitivity analyses and simulations.

<code>d_tau</code>	Positive numeric of length 1. Degrees of freedom of the Student t prior of <code>tau</code> if <code>prior_tau</code> is "half_t". In the case of <code>prior_tau</code> equal to "half_t", the defaults <code>s_tau = sd(data[[response]])</code> , <code>na.rm = TRUE</code>) and <code>d_tau = 1</code> specify a weakly informative scaled half-Cauchy distribution. This choice is only provisional. The prior on <code>tau</code> is extremely important, especially for small numbers of historical studies, and the user should set a reasonable value for the use case, ideally informed by the results of sensitivity analyses and simulations.
<code>prior_tau</code>	Character string, family of the prior of <code>tau</code> . If <code>prior_tau</code> equals "uniform", then the prior on <code>tau</code> is a uniform prior with lower bound 0 and upper bound <code>s_tau</code> . If <code>prior_tau</code> equals "half_t", then the prior on <code>tau</code> is a half Student-t prior with center 0, lower bound 0, scale parameter <code>s_tau</code> , and degrees of freedom <code>d_tau</code> . The scale parameter <code>s_tau</code> is analogous to the sigma parameter of the Student-t parameterization given at https://mc-stan.org/docs/functions-reference/unbounded_continuous_distributions.html . # no-lint
<code>n_chains</code>	Number of MCMC chains to run.
<code>n_adapt</code>	Number of adaptation iterations to run.
<code>n_warmup</code>	Number of warmup iterations per chain to run.
<code>n_iterations</code>	Number of saved MCMC iterations per chain to run.
<code>quiet</code>	Logical of length 1, TRUE to suppress R console output.

Value

A tidy data frame of parameter samples from the posterior distribution. Columns `.chain`, `.iteration`, and `.draw` have the meanings documented in the `posterior` package.

See Also

Other `mcmc`: [hb_convergence\(\)](#), [hb_mcmc_independent\(\)](#), [hb_mcmc_mixture\(\)](#), [hb_mcmc_mixture_hyperparameters\(\)](#), [hb_mcmc_pool\(\)](#)

Examples

```
if (!identical(Sys.getenv("HB_TEST", unset = ""), "")) {
  data <- hb_sim_hierarchical(n_continuous = 2)$data
  hb_mcmc_hierarchical(
    data,
    n_chains = 1,
    n_adapt = 100,
    n_warmup = 50,
    n_iterations = 50
  )
}
```

```
)
}
```

hb_mcmc_independent *Independent-study model MCMC*

Description

Run the independent-study model with MCMC.

Usage

```
hb_mcmc_independent(
  data,
  response = "response",
  study = "study",
  study_reference = max(data[[study]]),
  group = "group",
  group_reference = min(data[[group]]),
  patient = "patient",
  covariates = grep("^covariate", colnames(data), value = TRUE),
  s_alpha = 30,
  s_delta = 30,
  s_beta = 30,
  s_sigma = 30,
  n_chains = 4,
  n_adapt = 2000,
  n_warmup = 4000,
  n_observations = 20000,
  quiet = TRUE
)
```

Arguments

data	Tidy data frame with one row per patient, indicator columns for the response variable, study, group, and patient, and covariates. All columns must be atomic vectors (e.g. not lists). The data for the mixture and simple models should have just one study, and the others should have data from more than one study. The simple model can be used to get the historical data components of m_{ω} and s_{ω} for the mixture model.
response	Character of length 1, name of the column in data with the response/outcome variable. <code>data[[response]]</code> must be a continuous variable, and it <i>should</i> be the change from baseline of a clinical endpoint of interest, as opposed to just the raw response. Treatment differences are computed directly from this scale, please supply change from baseline unless you are absolutely certain that treatment differences computed directly from this quantity are clinically meaningful.
study	Character of length 1, name of the column in data with the study ID.

study_reference	Atomic of length 1, element of the study column that indicates the current study. (The other studies are historical studies.)
group	Character of length 1, name of the column in data with the group ID.
group_reference	Atomic of length 1, element of the group column that indicates the control group. (The other groups may be treatment groups.)
patient	Character of length 1, name of the column in data with the patient ID.
covariates	Character vector of column names in data with the columns with baseline covariates. These can be continuous, categorical, or binary. Regardless, <code>historicalborrow</code> derives the appropriate model matrix.
s_alpha	Numeric of length 1, prior standard deviation of the study-specific control group mean parameters alpha.
s_delta	Numeric of length 1, prior standard deviation of the study-by-group effect parameters delta.
s_beta	Numeric of length 1, prior standard deviation of the fixed effects beta.
s_sigma	Numeric of length 1, prior upper bound of the residual standard deviations.
n_chains	Number of MCMC chains to run.
n_adapt	Number of adaptation iterations to run.
n_warmup	Number of warmup iterations per chain to run.
n_iterations	Number of saved MCMC iterations per chain to run.
quiet	Logical of length 1, TRUE to suppress R console output.

Value

A tidy data frame of parameter samples from the posterior distribution. Columns `.chain`, `.iteration`, and `.draw` have the meanings documented in the `posterior` package.

See Also

Other `mcmc`: [hb_convergence\(\)](#), [hb_mcmc_hierarchical\(\)](#), [hb_mcmc_mixture\(\)](#), [hb_mcmc_mixture_hyperparameters](#), [hb_mcmc_pool\(\)](#)

Examples

```
if (!identical(Sys.getenv("HB_TEST", unset = ""), "")) {
  data <- hb_sim_independent(n_continuous = 2)$data
  hb_mcmc_independent(
    data,
    n_chains = 1,
    n_adapt = 100,
    n_warmup = 50,
    n_iterations = 50
  )
}
```

hb_mcmc_mixture

Mixture model MCMC

Description

Run the mixture model with MCMC.

Usage

```
hb_mcmc_mixture(
  data,
  response = "response",
  study = "study",
  study_reference = max(data[[study]]),
  group = "group",
  group_reference = min(data[[group]]),
  patient = "patient",
  covariates = grep("^covariate", colnames(data), value = TRUE),
  s_delta = 30,
  s_beta = 30,
  s_sigma = 30,
  m_omega = c(0, 0),
  s_omega = c(30, 30),
  p_omega = 1/length(m_omega),
  n_chains = 4,
  n_adapt = 2000,
  n_warmup = 4000,
  n_iterations = 20000,
  quiet = TRUE
)
```

Arguments

data	Tidy data frame with one row per patient, indicator columns for the response variable, study, group, and patient, and covariates. All columns must be atomic vectors (e.g. not lists). The data for the mixture and simple models should have just one study, and the others should have data from more than one study. The simple model can be used to get the historical data components of <code>m_omega</code> and <code>s_omega</code> for the mixture model.
response	Character of length 1, name of the column in data with the response/outcome variable. <code>data[[response]]</code> must be a continuous variable, and it <i>should</i> be the change from baseline of a clinical endpoint of interest, as opposed to just the raw response. Treatment differences are computed directly from this scale, please supply change from baseline unless you are absolutely certain that treatment differences computed directly from this quantity are clinically meaningful.
study	Character of length 1, name of the column in data with the study ID.

study_reference	Atomic of length 1, element of the study column that indicates the current study. (The other studies are historical studies.)
group	Character of length 1, name of the column in data with the group ID.
group_reference	Atomic of length 1, element of the group column that indicates the control group. (The other groups may be treatment groups.)
patient	Character of length 1, name of the column in data with the patient ID.
covariates	Character vector of column names in data with the columns with baseline covariates. These can be continuous, categorical, or binary. Regardless, <code>historicalborrow</code> derives the appropriate model matrix.
s_delta	Numeric of length 1, prior standard deviation of the study-by-group effect parameters delta.
s_beta	Numeric of length 1, prior standard deviation of the fixed effects beta.
s_sigma	Numeric of length 1, prior upper bound of the residual standard deviations.
m_omega	Numeric with length equal to the number of supposed studies (but only the current study is in the data). <code>m_omega</code> is the prior control group mean of each study. The last element corresponds to the current study, and the others are for historical studies.
s_omega	Numeric with length equal to the number of supposed studies (but only the current study is in the data). <code>s_omega</code> is the prior control group standard deviation of each study. The last element corresponds to the current study, and the others are for historical studies.
p_omega	Numeric with length equal to the number of supposed studies (but only the current study is in the data). <code>p_omega</code> is the prior control group mixture proportion of each study. The last element corresponds to the current study, and the others are for historical studies.
n_chains	Number of MCMC chains to run.
n_adapt	Number of adaptation iterations to run.
n_warmup	Number of warmup iterations per chain to run.
n_iterations	Number of saved MCMC iterations per chain to run.
quiet	Logical of length 1, TRUE to suppress R console output.

Details

The study-specific components of the mixture prior are all fixed in advance. Mixture components are normal distributions with means in `m_omega` and standard deviations in `s_omega`. These vectors are ordered with historical studies first and the current study last. These mixture components can be computed using `hb_mcmc_mixture_hyperparameters()` on a full set of data (all the historical studies and the current study together). Then the `m_omega` and `s_omega` columns of the output can be plugged directly into `hb_mcmc_mixture()`. See the examples for a demonstration.

Value

A tidy data frame of parameter samples from the posterior distribution. Columns `.chain`, `.iteration`, and `.draw` have the meanings documented in the `posterior` package.

See Also

Other mcmc: [hb_convergence\(\)](#), [hb_mcmc_hierarchical\(\)](#), [hb_mcmc_independent\(\)](#), [hb_mcmc_mixture_hyperparameters\(\)](#), [hb_mcmc_pool\(\)](#)

Examples

```
data_all_studies <- hb_sim_independent(n_continuous = 2)$data
data_all_studies$study <- paste0("study", data_all_studies$study)
hyperparameters <- hb_mcmc_mixture_hyperparameters(
  data = data_all_studies,
  response = "response",
  study = "study",
  study_reference = "study5",
  group = "group",
  group_reference = 1,
  patient = "patient",
  n_chains = 1,
  n_adapt = 100,
  n_warmup = 50,
  n_iterations = 50
)
print(hyperparameters)
data_current_study <- dplyr::filter(data_all_studies, study == max(study))
hb_mcmc_mixture(
  data = data_current_study,
  response = "response",
  study = "study",
  study_reference = "study5",
  group = "group",
  group_reference = 1,
  patient = "patient",
  m_omega = hyperparameters$m_omega, # use hyperparams from historical data
  s_omega = hyperparameters$s_omega, # use hyperparams from historical data
  p_omega = rep(1 / nrow(hyperparameters), nrow(hyperparameters)),
  n_chains = 1,
  n_adapt = 100,
  n_warmup = 50,
  n_iterations = 50
)
```

hb_mcmc_mixture_hyperparameters

Mixture model MCMC hyperparameters

Description

Run a simple model separately on each historical study control group and compute hyperparameters for [hb_mcmc_mixture\(\)](#).

Usage

```

hb_mcmc_mixture_hyperparameters(
  data,
  response = "response",
  study = "study",
  study_reference = max(data[[study]]),
  group = "group",
  group_reference = min(data[[group]]),
  patient = "patient",
  m_mu = 0,
  s_mu = 30,
  s_sigma = 30,
  m_omega_current = 0,
  s_omega_current = 30,
  n_chains = 4,
  n_adapt = 2000,
  n_warmup = 4000,
  n_iterations = 20000,
  quiet = TRUE
)

```

Arguments

data	Tidy data frame with one row per patient, indicator columns for the response variable, study, group, and patient, and covariates. All columns must be atomic vectors (e.g. not lists). The data for the mixture and simple models should have just one study, and the others should have data from more than one study. The simple model can be used to get the historical data components of m_omega and s_omega for the mixture model.
response	Character of length 1, name of the column in data with the response/outcome variable. data[[response]] must be a continuous variable, and it <i>should</i> be the change from baseline of a clinical endpoint of interest, as opposed to just the raw response. Treatment differences are computed directly from this scale, please supply change from baseline unless you are absolutely certain that treatment differences computed directly from this quantity are clinically meaningful.
study	Character of length 1, name of the column in data with the study ID.
study_reference	Atomic of length 1, element of the study column that indicates the current study. (The other studies are historical studies.)
group	Character of length 1, name of the column in data with the group ID.
group_reference	Atomic of length 1, element of the group column that indicates the control group. (The other groups may be treatment groups.)
patient	Character of length 1, name of the column in data with the patient ID.
m_mu	Numeric of length 1, prior mean of the mean mu in the simple model.

<code>s_mu</code>	Numeric of length 1, prior standard deviation of the mean μ in the simple model.
<code>s_sigma</code>	Numeric of length 1, uniform prior upper bound of the residual standard deviation σ in the simple model.
<code>m_omega_current</code>	Numeric with length 1, <code>m_omega</code> value of the current study. Inserted as the final component of the <code>m_omega</code> column in the output.
<code>s_omega_current</code>	Numeric with length 1, <code>s_omega</code> value of the current study. Inserted as the final component of the <code>s_omega</code> column in the output.
<code>n_chains</code>	Number of MCMC chains to run.
<code>n_adapt</code>	Number of adaptation iterations to run.
<code>n_warmup</code>	Number of warmup iterations per chain to run.
<code>n_iterations</code>	Number of saved MCMC iterations per chain to run.
<code>quiet</code>	Logical of length 1, TRUE to suppress R console output.

Details

The model is a simple Bayesian model with a normal likelihood, an unknown mean μ , and an unknown standard deviation σ . For each historical study, the posterior mean of μ becomes the corresponding component of `m_omega` in the output, and the posterior standard deviation of μ becomes the corresponding component of `s_omega` in the output. See the examples in this help file for a demonstration. `m_omega` and `s_omega` define the components of the mixture prior in `hb_mcmc_mixture()` that act as the contribution of the historical studies to the model.

Value

A tidy data frame of hyperparameter values for `hb_mcmc_mixture()`. The first several rows are for historical studies, and the last row is for the current study. Studies/rows are sorted in the order `hb_mcmc_mixture()` sorts them, so you can use columns `m_omega` and `s_omega` for the same dataset and same values of other arguments directly in `hb_mcmc_mixture()`.

See Also

Other mcmc: `hb_convergence()`, `hb_mcmc_hierarchical()`, `hb_mcmc_independent()`, `hb_mcmc_mixture()`, `hb_mcmc_pool()`

Examples

```
data_all_studies <- hb_sim_independent(n_continuous = 2)$data
data_all_studies$study <- paste0("study", data_all_studies$study)
hyperparameters <- hb_mcmc_mixture_hyperparameters(
  data = data_all_studies,
  response = "response",
  study = "study",
  study_reference = "study5",
  group = "group",
  group_reference = 1,
```

```

    patient = "patient",
    n_chains = 1,
    n_adapt = 100,
    n_warmup = 50,
    n_iterations = 50
  )
print(hyperparameters)
data_current_study <- dplyr::filter(data_all_studies, study == max(study))
hb_mcmc_mixture(
  data = data_current_study,
  response = "response",
  study = "study",
  study_reference = "study5",
  group = "group",
  group_reference = 1,
  patient = "patient",
  m_omega = hyperparameters$m_omega, # use hyperparams from historical data
  s_omega = hyperparameters$s_omega, # use hyperparams from historical data
  p_omega = rep(1 / nrow(hyperparameters), nrow(hyperparameters)),
  n_chains = 1,
  n_adapt = 100,
  n_warmup = 50,
  n_iterations = 50
)

```

hb_mcmc_pool

Non-longitudinal pooled MCMC

Description

Run the non-longitudinal pooled model with MCMC.

Usage

```

hb_mcmc_pool(
  data,
  response = "response",
  study = "study",
  study_reference = max(data[[study]]),
  group = "group",
  group_reference = min(data[[group]]),
  patient = "patient",
  covariates = grep("^covariate", colnames(data), value = TRUE),
  s_alpha = 30,
  s_delta = 30,
  s_beta = 30,
  s_sigma = 30,
  n_chains = 4,
  n_adapt = 2000,

```

```

n_warmup = 4000,
n_iterations = 20000,
quiet = TRUE
)

```

Arguments

data	Tidy data frame with one row per patient, indicator columns for the response variable, study, group, and patient, and covariates. All columns must be atomic vectors (e.g. not lists). The data for the mixture and simple models should have just one study, and the others should have data from more than one study. The simple model can be used to get the historical data components of <code>m_omega</code> and <code>s_omega</code> for the mixture model.
response	Character of length 1, name of the column in data with the response/outcome variable. <code>data[[response]]</code> must be a continuous variable, and it <i>should</i> be the change from baseline of a clinical endpoint of interest, as opposed to just the raw response. Treatment differences are computed directly from this scale, please supply change from baseline unless you are absolutely certain that treatment differences computed directly from this quantity are clinically meaningful.
study	Character of length 1, name of the column in data with the study ID.
study_reference	Atomic of length 1, element of the study column that indicates the current study. (The other studies are historical studies.)
group	Character of length 1, name of the column in data with the group ID.
group_reference	Atomic of length 1, element of the group column that indicates the control group. (The other groups may be treatment groups.)
patient	Character of length 1, name of the column in data with the patient ID.
covariates	Character vector of column names in data with the columns with baseline covariates. These can be continuous, categorical, or binary. Regardless, <code>historicalborrow</code> derives the appropriate model matrix.
s_alpha	Numeric of length 1, prior standard deviation of the study-specific control group mean parameters alpha.
s_delta	Numeric of length 1, prior standard deviation of the study-by-group effect parameters delta.
s_beta	Numeric of length 1, prior standard deviation of the fixed effects beta.
s_sigma	Numeric of length 1, prior upper bound of the residual standard deviations.
n_chains	Number of MCMC chains to run.
n_adapt	Number of adaptation iterations to run.
n_warmup	Number of warmup iterations per chain to run.
n_iterations	Number of saved MCMC iterations per chain to run.
quiet	Logical of length 1, TRUE to suppress R console output.

Value

A tidy data frame of parameter samples from the posterior distribution. Columns `.chain`, `.iteration`, and `.draw` have the meanings documented in the `posterior` package.

See Also

Other mcmc: [hb_convergence\(\)](#), [hb_mcmc_hierarchical\(\)](#), [hb_mcmc_independent\(\)](#), [hb_mcmc_mixture\(\)](#), [hb_mcmc_mixture_hyperparameters\(\)](#)

Examples

```
if (!identical(Sys.getenv("HB_TEST", unset = ""), "")) {
  data <- hb_sim_pool(n_continuous = 2)$data
  hb_mcmc_pool(
    data,
    n_chains = 1,
    n_adapt = 100,
    n_warmup = 50,
    n_iterations = 50
  )
}
```

 hb_plot_borrow

Plot a borrowing model response against the benchmark models.

Description

Plot the response from a borrowing model (hierarchical or mixture) against the independent and pooled benchmark models.

Usage

```
hb_plot_borrow(borrow, pool, independent, outcome = c("response", "diff"))
```

Arguments

<code>borrow</code>	A data frame returned by hb_summary() for the mixture or hierarchical model.
<code>pool</code>	A data frame returned by hb_summary() for the pooled model.
<code>independent</code>	A data frame returned by hb_summary() for the independent model.
<code>outcome</code>	Character of length 1, either "response" or "diff", the quantity to plot on the vertical axis.

Value

A `ggplot` object

See Also

Other plot: [hb_plot_group\(\)](#), [hb_plot_tau\(\)](#)

Examples

```
if (!identical(Sys.getenv("HB_TEST", unset = ""), "")) {
  data <- hb_sim_independent(n_continuous = 2)$data
  mcmc_borrow <- hb_mcmc_hierarchical(
    data,
    n_chains = 1,
    n_adapt = 100,
    n_warmup = 100,
    n_iterations = 200
  )
  mcmc_pool <- hb_mcmc_pool(
    data,
    n_chains = 1,
    n_adapt = 100,
    n_warmup = 200,
    n_iterations = 200
  )
  mcmc_independent <- hb_mcmc_independent(
    data,
    n_chains = 1,
    n_adapt = 100,
    n_warmup = 200,
    n_iterations = 200
  )
  borrow <- hb_summary(mcmc_borrow, data)
  pool <- hb_summary(mcmc_pool, data)
  independent <- hb_summary(mcmc_independent, data)
  hb_plot_borrow(
    borrow = borrow,
    pool = pool,
    independent = independent
  )
}
```

 hb_plot_group

Plot the groups of a borrowing model and its benchmark models.

Description

Plot the groups against one another for a borrowing model (hierarchical or mixture) and the independent and pooled benchmark models.

Usage

```
hb_plot_group(borrow, pool, independent, outcome = c("response", "diff"))
```

Arguments

borrow	A data frame returned by <code>hb_summary()</code> for the mixture or hierarchical model.
pool	A data frame returned by <code>hb_summary()</code> for the pooled model.
independent	A data frame returned by <code>hb_summary()</code> for the independent model.
outcome	Character of length 1, either "response" or "diff", the quantity to plot on the vertical axis.

Value

A ggplot object

See Also

Other plot: `hb_plot_borrow()`, `hb_plot_tau()`

Examples

```
if (!identical(Sys.getenv("HB_TEST", unset = ""), "")) {
  data <- hb_sim_independent(n_continuous = 2)$data
  mcmc_borrow <- hb_mcmc_hierarchical(
    data,
    n_chains = 1,
    n_adapt = 100,
    n_warmup = 100,
    n_iterations = 200
  )
  mcmc_pool <- hb_mcmc_pool(
    data,
    n_chains = 1,
    n_adapt = 100,
    n_warmup = 200,
    n_iterations = 200
  )
  mcmc_independent <- hb_mcmc_independent(
    data,
    n_chains = 1,
    n_adapt = 100,
    n_warmup = 200,
    n_iterations = 200
  )
  borrow <- hb_summary(mcmc_borrow, data)
  pool <- hb_summary(mcmc_pool, data)
  independent <- hb_summary(mcmc_independent, data)
  hb_plot_group(
    borrow = borrow,
    pool = pool,
    independent = independent
  )
}
```

hb_plot_tau	<i>Plot tau</i>
-------------	-----------------

Description

Plot the tau parameter of a fitted hierarchical model.

Usage

```
hb_plot_tau(mcmc)
```

Arguments

mcmc Data frame of posterior samples generated by [hb_mcmc_hierarchical\(\)](#).

Value

A ggplot object

See Also

Other plot: [hb_plot_borrow\(\)](#), [hb_plot_group\(\)](#)

Examples

```
data <- hb_sim_independent(n_continuous = 2)$data
mcmc <- hb_mcmc_hierarchical(
  data,
  n_chains = 1,
  n_adapt = 100,
  n_warmup = 100,
  n_iterations = 200
)
hb_plot_tau(mcmc = mcmc)
```

hb_sim_hierarchical	<i>Non-longitudinal hierarchical simulations.</i>
---------------------	---

Description

Simulate from the non-longitudinal hierarchical model.

Usage

```

hb_sim_hierarchical(
  n_study = 5,
  n_group = 3,
  n_patient = 100,
  n_continuous = 0,
  n_binary = 0,
  s_delta = 1,
  s_beta = 1,
  s_sigma = 1,
  s_mu = 1,
  s_tau = 1,
  d_tau = 4,
  prior_tau = "half_t",
  alpha = NULL,
  delta = stats::rnorm(n = n_group - 1, mean = 0, sd = s_delta),
  beta = stats::rnorm(n = n_study * (n_continuous + n_binary), mean = 0, sd = s_delta),
  sigma = stats::runif(n = n_study, min = 0, max = s_sigma),
  mu = stats::rnorm(n = 1, mean = 0, sd = s_mu),
  tau = NULL
)

```

Arguments

n_study	Number of studies to simulate.
n_group	Number of groups (e.g. study arms) to simulate per study.
n_patient	Number of patients to simulate per study per group.
n_continuous	Number of continuous covariates to simulate (all from independent standard normal distributions).
n_binary	Number of binary covariates to simulate (all from independent Bernoulli distributions with $p = 0.5$).
s_delta	Numeric of length 1, prior standard deviation of the study-by-group effect parameters delta.
s_beta	Numeric of length 1, prior standard deviation of the fixed effects beta.
s_sigma	Numeric of length 1, prior upper bound of the residual standard deviations.
s_mu	Numeric of length 1, prior standard deviation of mu.
s_tau	Non-negative numeric of length 1. If prior_tau is "half_t", then s_tau is the scale parameter of the Student t prior of tau and analogous to the sigma parameter of the Student-t parameterization given at https://mc-stan.org/docs/functions-reference/unbounded_continuous_distributions.html . # nolint If prior_tau is "uniform", then s_tau is the upper bound of tau. Upper bound on tau if prior_tau is "uniform".
d_tau	Positive numeric of length 1. Degrees of freedom of the Student t prior of tau if prior_tau is "half_t".

prior_tau	Character string, family of the prior of tau. If prior_tau equals "uniform", then the prior on tau is a uniform prior with lower bound 0 and upper bound s_tau. If prior_tau equals "half_t", then the prior on tau is a half Student-t prior with center 0, lower bound 0, scale parameter s_tau, and degrees of freedom d_tau. The scale parameter s_tau is analogous to the sigma parameter of the Student-t parameterization given at https://mc-stan.org/docs/functions-reference/unbounded_continuous_distributions.html . # no-lint
alpha	Numeric vector of length 1 for the pooled and mixture models and length n_study for the independent and hierarchical models. alpha is the vector of control group mean parameters. alpha enters the model by multiplying with <code>\$matrices\$x_alpha</code> (see the return value). The control group in the data is the one with the group column equal to 1.
delta	Numeric vector of length n_group - 1 of treatment effect parameters. delta enters the model by multiplying with <code>\$matrices\$x_delta</code> (see the return value). The control (non-treatment) group in the data is the one with the group column equal to 1.
beta	Numeric vector of n_study * (n_continuous + n_binary) fixed effect parameters. Within each study, the first n_continuous betas are for the continuous covariates, and the rest are for the binary covariates. All the betas for one study appear before all the betas for the next study, and studies are arranged in increasing order of the sorted unique values in <code>\$data\$study</code> in the output. betas enters the model by multiplying with <code>\$matrices\$x_alpha</code> (see the return value).
sigma	Numeric vector of n_study study-specific residual standard deviations.
mu	Numeric of length 1, mean of the control group means alpha.
tau	Numeric of length 1, standard deviation of the control group means alpha.

Value

A list with the following elements:

- `data`: tidy long-form dataset with the patient-level data. one row per patient and indicator columns for the study, group (e.g. treatment arm), and patient ID. The response columns is the patient response. The other columns are baseline covariates. The control group is the one with the group column equal to 1, and the current study (non-historical) is the one with the maximum value of the study column. Only the current study has any non-control-group patients, the historical studies have only the control group.
- `parameters`: named list of model parameter values. See the model specification vignette for details.
- `matrices`: A named list of model matrices. See the model specification vignette for details.

See Also

Other simulate: `hb_sim_independent()`, `hb_sim_mixture()`, `hb_sim_pool()`

Examples

```
hb_sim_hierarchical()$data
```

```
hb_sim_independent      Non-longitudinal independent simulations.
```

Description

Simulate from the non-longitudinal independent model.

Usage

```
hb_sim_independent(
  n_study = 5,
  n_group = 3,
  n_patient = 100,
  n_continuous = 0,
  n_binary = 0,
  s_alpha = 1,
  s_delta = 1,
  s_beta = 1,
  s_sigma = 1,
  alpha = stats::rnorm(n = n_study, mean = 0, sd = s_alpha),
  delta = stats::rnorm(n = n_group - 1, mean = 0, sd = s_delta),
  beta = stats::rnorm(n = n_study * (n_continuous + n_binary), mean = 0, sd = s_delta),
  sigma = stats::runif(n = n_study, min = 0, max = s_sigma)
)
```

Arguments

n_study	Number of studies to simulate.
n_group	Number of groups (e.g. study arms) to simulate per study.
n_patient	Number of patients to simulate per study per group.
n_continuous	Number of continuous covariates to simulate (all from independent standard normal distributions).
n_binary	Number of binary covariates to simulate (all from independent Bernoulli distributions with $p = 0.5$).
s_alpha	Numeric of length 1, prior standard deviation of the study-specific control group mean parameters alpha.
s_delta	Numeric of length 1, prior standard deviation of the study-by-group effect parameters delta.
s_beta	Numeric of length 1, prior standard deviation of the fixed effects beta.
s_sigma	Numeric of length 1, prior upper bound of the residual standard deviations.

alpha	Numeric vector of length 1 for the pooled and mixture models and length n_study for the independent and hierarchical models. alpha is the vector of control group mean parameters. alpha enters the model by multiplying with $\$matrices\x_alpha (see the return value). The control group in the data is the one with the group column equal to 1.
delta	Numeric vector of length n_group - 1 of treatment effect parameters. delta enters the model by multiplying with $\$matrices\x_delta (see the return value). The control (non-treatment) group in the data is the one with the group column equal to 1.
beta	Numeric vector of n_study * (n_continuous + n_binary) fixed effect parameters. Within each study, the first n_continuous betas are for the continuous covariates, and the rest are for the binary covariates. All the betas for one study appear before all the betas for the next study, and studies are arranged in increasing order of the sorted unique values in $\$data\$study$ in the output. betas enters the model by multiplying with $\$matrices\x_alpha (see the return value).
sigma	Numeric vector of n_study study-specific residual standard deviations.

Value

A list with the following elements:

- data: tidy long-form dataset with the patient-level data. one row per patient and indicator columns for the study, group (e.g. treatment arm), and patient ID. The response columns is the patient response. The other columns are baseline covariates. The control group is the one with the group column equal to 1, and the current study (non-historical) is the one with the maximum value of the study column. Only the current study has any non-control-group patients, the historical studies have only the control group.
- parameters: named list of model parameter values. See the model specification vignette for details.
- matrices: A named list of model matrices. See the model specification vignette for details.

See Also

Other simulate: [hb_sim_hierarchical\(\)](#), [hb_sim_mixture\(\)](#), [hb_sim_pool\(\)](#)

Examples

```
hb_sim_independent()$data
```

hb_sim_mixture *Non-longitudinal mixture simulations.*

Description

Simulate from the non-longitudinal mixture model.

Usage

```

hb_sim_mixture(
  n_study = 5,
  n_group = 3,
  n_patient = 100,
  n_continuous = 0,
  n_binary = 0,
  s_delta = 1,
  s_beta = 1,
  s_sigma = 1,
  m_omega = 0,
  s_omega = 1,
  p_omega = 1/n_study,
  alpha = omega[pi],
  delta = stats::rnorm(n = n_group - 1, mean = 0, sd = s_delta),
  beta = stats::rnorm(n = n_continuous + n_binary, mean = 0, sd = s_delta),
  sigma = stats::runif(n = 1, min = 0, max = s_sigma),
  pi = sample.int(n = n_study, size = 1, prob = p_omega),
  omega = stats::rnorm(n = n_study, mean = m_omega, sd = s_omega)
)

```

Arguments

n_study	Number of studies to simulate.
n_group	Number of groups (e.g. study arms) to simulate per study.
n_patient	Number of patients to simulate per study per group.
n_continuous	Number of continuous covariates to simulate (all from independent standard normal distributions).
n_binary	Number of binary covariates to simulate (all from independent Bernoulli distributions with $p = 0.5$).
s_delta	Numeric of length 1, prior standard deviation of the study-by-group effect parameters delta.
s_beta	Numeric of length 1, prior standard deviation of the fixed effects beta.
s_sigma	Numeric of length 1, prior upper bound of the residual standard deviations.
m_omega	Numeric of length 1 or n_study, prior control group mean of each study. If length n_study, then the last element corresponds to the current study, and the others are for historical studies.
s_omega	Numeric of length 1 or n_study, prior control group standard deviation of each study. If length n_study, the the last element corresponds to the current study, and the others are for historical studies.
p_omega	Numeric of length n_study, prior mixture proportion of each study. If length n_study, then the last element corresponds to the current study, and the others are for historical studies.
alpha	Numeric vector of length 1 for the pooled and mixture models and length n_study for the independent and hierarchical models. alpha is the vector of control

	group mean parameters. <code>alpha</code> enters the model by multiplying with <code>\$matrices\$x_alpha</code> (see the return value). The control group in the data is the one with the group column equal to 1.
<code>delta</code>	Numeric vector of length <code>n_group - 1</code> of treatment effect parameters. <code>delta</code> enters the model by multiplying with <code>\$matrices\$x_delta</code> (see the return value). The control (non-treatment) group in the data is the one with the group column equal to 1.
<code>beta</code>	Numeric vector of <code>n_continuous + n_binary</code> fixed effect parameters. The first <code>n_continuous</code> betas are for the continuous covariates, and the rest are for the binary covariates. <code>betas</code> enters the model by multiplying with <code>\$matrices\$x_alpha</code> (see the return value).
<code>sigma</code>	Numeric vector of <code>n_study</code> study-specific residual standard deviations.
<code>pi</code>	Integer of length 1, index of the mixture component randomly chosen for <code>alpha</code> .
<code>omega</code>	Numeric of length <code>n_study</code> , Candidate placebo mean parameters drawn from each of the mixture components.

Value

A list with the following elements:

- `data`: tidy long-form dataset with the patient-level data. one row per patient and indicator columns for the study, group (e.g. treatment arm), and patient ID. The response column is the patient response. The other columns are baseline covariates. The control group is the one with the group column equal to 1, and the current study (non-historical) is the one with the maximum value of the study column. Only the current study has any non-control-group patients, the historical studies have only the control group.
- `parameters`: named list of model parameter values. See the model specification vignette for details.
- `matrices`: A named list of model matrices. See the model specification vignette for details.

See Also

Other simulate: [hb_sim_hierarchical\(\)](#), [hb_sim_independent\(\)](#), [hb_sim_pool\(\)](#)

Examples

```
hb_sim_mixture()$data
```

```
hb_sim_pool
```

Non-longitudinal pooled simulations.

Description

Simulate from the non-longitudinal pooled model.

Usage

```

hb_sim_pool(
  n_study = 5,
  n_group = 3,
  n_patient = 100,
  n_continuous = 0,
  n_binary = 0,
  s_alpha = 1,
  s_delta = 1,
  s_beta = 1,
  s_sigma = 1,
  alpha = stats::rnorm(n = 1, mean = 0, sd = s_alpha),
  delta = stats::rnorm(n = n_group - 1, mean = 0, sd = s_delta),
  beta = stats::rnorm(n = n_study * (n_continuous + n_binary), mean = 0, sd = s_delta),
  sigma = stats::runif(n = n_study, min = 0, max = s_sigma)
)

```

Arguments

n_study	Number of studies to simulate.
n_group	Number of groups (e.g. study arms) to simulate per study.
n_patient	Number of patients to simulate per study per group.
n_continuous	Number of continuous covariates to simulate (all from independent standard normal distributions).
n_binary	Number of binary covariates to simulate (all from independent Bernoulli distributions with $p = 0.5$).
s_alpha	Numeric of length 1, prior standard deviation of the study-specific control group mean parameters alpha.
s_delta	Numeric of length 1, prior standard deviation of the study-by-group effect parameters delta.
s_beta	Numeric of length 1, prior standard deviation of the fixed effects beta.
s_sigma	Numeric of length 1, prior upper bound of the residual standard deviations.
alpha	Numeric vector of length 1 for the pooled and mixture models and length n_study for the independent and hierarchical models. alpha is the vector of control group mean parameters. alpha enters the model by multiplying with <code>\$matrices\$x_alpha</code> (see the return value). The control group in the data is the one with the group column equal to 1.
delta	Numeric vector of length n_group - 1 of treatment effect parameters. delta enters the model by multiplying with <code>\$matrices\$x_delta</code> (see the return value). The control (non-treatment) group in the data is the one with the group column equal to 1.
beta	Numeric vector of n_study * (n_continuous + n_binary) fixed effect parameters. Within each study, the first n_continuous betas are for the continuous covariates, and the rest are for the binary covariates. All the betas for one study appear before all the betas for the next study, and studies are arranged

in increasing order of the sorted unique values in `$data$study` in the output. `betas` enters the model by multiplying with `$matrices$x_alpha` (see the return value).

`sigma` Numeric vector of `n_study` study-specific residual standard deviations.

Value

A list with the following elements:

- `data`: tidy long-form dataset with the patient-level data. one row per patient and indicator columns for the study, group (e.g. treatment arm), and patient ID. The response column is the patient response. The other columns are baseline covariates. The control group is the one with the group column equal to 1, and the current study (non-historical) is the one with the maximum value of the study column. Only the current study has any non-control-group patients, the historical studies have only the control group.
- `parameters`: named list of model parameter values. See the model specification vignette for details.
- `matrices`: A named list of model matrices. See the model specification vignette for details.

See Also

Other simulate: [hb_sim_hierarchical\(\)](#), [hb_sim_independent\(\)](#), [hb_sim_mixture\(\)](#)

Examples

```
hb_sim_pool(n_continuous = 1)$data
```

<code>hb_summary</code>	<i>Model summary</i>
-------------------------	----------------------

Description

Summarize a fitted model in a table.

Usage

```
hb_summary(
  mcmc,
  data,
  response = "response",
  study = "study",
  study_reference = max(data[[study]]),
  group = "group",
  group_reference = min(data[[group]]),
  patient = "patient",
  covariates = grep("^covariate", colnames(data), value = TRUE),
  eoi = 0,
  direction = "<"
)
```

Arguments

mcmc	A wide data frame of posterior samples returned by <code>hb_mcmc_hierarchical()</code> or similar MCMC function.
data	Tidy data frame with one row per patient, indicator columns for the response variable, study, group, and patient, and covariates. All columns must be atomic vectors (e.g. not lists). The data for the mixture and simple models should have just one study, and the others should have data from more than one study. The simple model can be used to get the historical data components of <code>m_omega</code> and <code>s_omega</code> for the mixture model.
response	Character of length 1, name of the column in data with the response/outcome variable. <code>data[[response]]</code> must be a continuous variable, and it <i>should</i> be the change from baseline of a clinical endpoint of interest, as opposed to just the raw response. Treatment differences are computed directly from this scale, please supply change from baseline unless you are absolutely certain that treatment differences computed directly from this quantity are clinically meaningful.
study	Character of length 1, name of the column in data with the study ID.
study_reference	Atomic of length 1, element of the study column that indicates the current study. (The other studies are historical studies.)
group	Character of length 1, name of the column in data with the group ID.
group_reference	Atomic of length 1, element of the group column that indicates the control group. (The other groups may be treatment groups.)
patient	Character of length 1, name of the column in data with the patient ID.
covariates	Character vector of column names in data with the columns with baseline covariates. These can be continuous, categorical, or binary. Regardless, <code>historicalborrow</code> derives the appropriate model matrix.
eoi	Numeric of length at least 1, vector of effects of interest (EOIs) for critical success factors (CSFs).
direction	Character of length <code>length(eoi)</code> indicating how to compare the treatment effect to each EOI. ">" means <code>Prob(treatment effect > EOI)</code> , and "<" means <code>Prob(treatment effect < EOI)</code> . All elements of <code>direction</code> must be either ">" or "<".

Details

The `hb_summary()` function post-processes the results from the model. It estimates marginal means of the response, treatment effect, and other quantities of interest.

Value

A tidy data frame with one row per group (e.g. treatment arm) and the columns in the following list. Unless otherwise specified, the quantities are calculated at the group level. Some are calculated for the current (non-historical) study only, while others pertain to the combined dataset which includes all historical studies. The mixture model is an exception because the `data` argument only includes the current study, so other quantities that include historical information will need to borrow from an `hb_summary()` call on one of the other models.

- `group`: group label.
- `data_mean`: observed mean response specific to the current study.
- `data_sd`: observed standard deviation of the response specific to the current study.
- `data_lower`: lower bound of a simple frequentist 95% confidence interval of the observed mean specific to the current study.
- `data_upper`: upper bound of a simple frequentist 95% confidence interval of the observed mean specific to the current study.
- `data_n`: number of non-missing observations in the combined dataset with all studies.
- `data_N`: total number of observations (missing and non-missing) in the combined dataset with all studies.
- `data_n_study_*`: number of non-missing observations separately for each study. The suffixes of these column names are integer study indexes. Call `dplyr::distinct(hb_data(your_data), study, study_label)` to see which study labels correspond to these integer indexes. Note: the combined dataset for the mixture model is just the current study. If all the `data_n_study_*` results across all studies are desired, then call `hb_summary()` on a different model (e.g. pooled).
- `data_N_study_*`: same as `data_n_study_*` except both missing and non-missing observations are counted (total number of observations).
- `response_mean`: Estimated posterior mean of the response from the model specific to the current study. Typically, the raw response is change from baseline, in which case `response_mean` is estimating change from baseline.
- `response_sd`: Estimated posterior standard deviation of the mean response from the model specific to the current study.
- `response_variance`: Estimated posterior variance of the mean response from the model specific to the current study.
- `response_lower`: Lower bound of a 95% posterior interval on the mean response from the model specific to the current study.
- `response_upper`: Upper bound of a 95% posterior interval on the mean response from the model specific to the current study.
- `response_mean_mcse`: Monte Carlo standard error of `response_mean`.
- `response_sd_mcse`: Monte Carlo standard error of `response_sd`.
- `response_lower_mcse`: Monte Carlo standard error of `response_lower`.
- `response_upper_mcse`: Monte Carlo standard error of `response_upper`.
- `diff_mean`: Estimated treatment effect from the model specific to the current study.
- `diff_lower`: Lower bound of a 95% posterior interval on the treatment effect from the model specific to the current study..
- `diff_upper`: Upper bound of a 95% posterior interval on the treatment effect from the model specific to the current study..
- `diff_mean_mcse`: Monte Carlo standard error of `diff_mean`.
- `diff_lower_mcse`: Monte Carlo standard error of `diff_lower`.
- `diff_upper_mcse`: Monte Carlo standard error of `diff_upper`.
- $P(\text{diff} > \text{EOI})$, $P(\text{diff} < \text{EOI})$: CSF probabilities on the treatment effect specified with the `eoi` and `direction` arguments. Specific to the current study.

- `effect_mean`: Estimated posterior mean of effect size (treatment difference divided by residual standard deviation). Specific to the current study.
- `effect_lower`: Lower bound of a 95% posterior interval of effect size from the model. Specific to the current study.
- `effect_upper`: Upper bound of a 95% posterior interval of effect size from the model. Specific to the current study.
- `precision_ratio`: For the hierarchical model only, a model-based mean of the precision ratio. Specific to the current study.
- `precision_ratio_lower`: For the hierarchical model only, lower bound of a model-based 95% posterior interval of the precision ratio. Specific to the current study.
- `precision_ratio_upper`: For the hierarchical model only, upper bound of a model-based 95% posterior interval of the precision ratio. Specific to the current study.
- `mix_prop_*`: For the mixture model only, posterior mixture proportions of each of the mixture components. The last one is for the current study and the first ones are for the historical studies. The suffixes of these column names are the integer study indexes. Call `dplyr::distinct(hb_data(your_data), study, study_label)` to see which study labels correspond to these integer indexes.

See Also

Other summary: [hb_ess\(\)](#)

Examples

```
if (!identical(Sys.getenv("HB_TEST", unset = ""), "")) {
  data <- hb_sim_pool(n_continuous = 2)$data
  data$group <- sprintf("group%s", data$group)
  mcmc <- hb_mcmc_pool(
    data,
    n_chains = 1,
    n_adapt = 100,
    n_warmup = 50,
    n_iterations = 50
  )
  hb_summary(mcmc, data)
}
```

Index

- * **data**
 - hb_data, 3
- * **help**
 - historicalborrow-package, 2
- * **mcmc**
 - hb_convergence, 2
 - hb_mcmc_hierarchical, 7
 - hb_mcmc_independent, 10
 - hb_mcmc_mixture, 12
 - hb_mcmc_mixture_hyperparameters, 14
 - hb_mcmc_pool, 17
- * **plot**
 - hb_plot_borrow, 19
 - hb_plot_group, 20
 - hb_plot_tau, 22
- * **simulate**
 - hb_sim_hierarchical, 22
 - hb_sim_independent, 25
 - hb_sim_mixture, 26
 - hb_sim_pool, 28
- * **summary**
 - hb_ess, 5
 - hb_summary, 30

hb_convergence, 2, 9, 11, 14, 16, 19
hb_data, 3
hb_ess, 5, 33
hb_mcmc_hierarchical, 3, 7, 11, 14, 16, 19
hb_mcmc_hierarchical(), 2, 6, 22, 31
hb_mcmc_independent, 3, 9, 10, 14, 16, 19
hb_mcmc_mixture, 3, 9, 11, 12, 16, 19
hb_mcmc_mixture(), 13, 14, 16
hb_mcmc_mixture_hyperparameters, 3, 9, 11, 14, 14, 19
hb_mcmc_mixture_hyperparameters(), 13
hb_mcmc_pool, 3, 9, 11, 14, 16, 17
hb_mcmc_pool(), 6
hb_plot_borrow, 19, 21, 22
hb_plot_group, 20, 20, 22
hb_plot_tau, 20, 21, 22
hb_sim_hierarchical, 22, 26, 28, 30
hb_sim_independent, 24, 25, 28, 30
hb_sim_mixture, 24, 26, 26, 30
hb_sim_pool, 24, 26, 28, 28
hb_summary, 6, 30
hb_summary(), 19, 21
historicalborrow-package, 2