

Package ‘easyRaschBayes’

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Title Bayesian Rasch Analysis Using 'brms'

Version 0.3.0

Description Reproduces classic Rasch psychometric analysis features using Bayesian item response theory models fitted with 'brms' following Bürkner (2021) <[doi:10.18637/jss.v100.i05](https://doi.org/10.18637/jss.v100.i05)> and Bürkner (2020) <[doi:10.3390/jintelligence8010005](https://doi.org/10.3390/jintelligence8010005)>. Supports both dichotomous and polytomous Rasch models. Features include posterior predictive item fit, conditional infit, item-restscore associations, person fit, differential item functioning, local dependence assessment via Q3 residual correlations, dimensionality assessment with residual principal components analysis, person-item targeting plots, item category probability curves, and reliability using relative measurement uncertainty following Bignardi et al. (2025) <[doi:10.31234/osf.io/h54k8_v1](https://doi.org/10.31234/osf.io/h54k8_v1)>.

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dif_statistic

*Differential Item Functioning (DIF) Analysis for Bayesian IRT Models***Description**

Tests for differential item functioning (DIF) in Bayesian Rasch-family models fitted with **brms** by comparing item parameters across subgroups defined by an exogenous variable. The function fits a DIF model that includes group-by-item interactions and summarizes the posterior distribution of the DIF effects.

Usage

```
dif_statistic(
  model,
  group_var,
  item_var = item,
  person_var = id,
  data = NULL,
  dif_type = c("uniform", "non-uniform"),
  prob = 0.95,
  rope = 0.5,
  refit = TRUE,
  ...
)
```

Arguments

model	A fitted brmsfit object from the baseline (no-DIF) model.
group_var	An unquoted variable name identifying the grouping variable for DIF testing (e.g., gender). Must be a factor or character variable with exactly 2 levels in the current implementation.
item_var	An unquoted variable name identifying the item grouping variable. Default is <code>item</code> .
person_var	An unquoted variable name identifying the person grouping variable. Default is <code>id</code> .
data	An optional data frame containing all variables needed for the DIF model, including the group variable. If <code>NULL</code> (the default), the function attempts to use <code>model\$data</code> . Since the baseline model formula typically does not include the group variable, brms will have dropped it from the stored model data. In that case, you must supply the original data frame here.
dif_type	Character. For polytomous ordinal models only. "uniform" (the default) tests for a uniform location shift per item via a <code>group:item</code> fixed-effect interaction. "non-uniform" fits group-specific thresholds per item and computes per-threshold DIF effects as the difference between groups. Ignored for dichotomous models.

prob	Numeric in (0, 1). Width of the credible intervals. Default is 0.95.
rope	Numeric. Half-width of the Region of Practical Equivalence (ROPE) around zero for DIF effects, on the logit scale. Default is 0.5, corresponding to a practically negligible DIF effect. Set to 0 to skip ROPE analysis.
refit	Logical. If TRUE (the default), the DIF model is fitted automatically by updating the baseline model via update , which reuses the compiled Stan code for faster sampling. If FALSE, only the DIF model formula is returned (useful for manual fitting with custom settings).
...	Additional arguments passed to update.brmsfit when refitting the DIF model (e.g., cores, control).

Details

For polytomous models, two types of DIF can be tested:

Uniform DIF (`dif_type = "uniform"`, **default**) A single location shift per item across groups, modelled as a `group:item` fixed-effect interaction. This tests whether the average item difficulty differs between groups.

Non-uniform / threshold-level DIF (`dif_type = "non-uniform"`) Each item receives group-specific thresholds via `thres(gr = interaction(item, group))`. DIF effects are computed as the difference in each threshold between groups, revealing whether DIF affects specific response categories.

The function constructs a DIF model by adding a group-by-item interaction to the baseline model:

- **Dichotomous models** (`family = bernoulli()`): The baseline response $\sim 1 + (1 \mid \text{item}) + (1 \mid \text{id})$ becomes response $\sim 1 + \text{group} + (1 + \text{group} \mid \text{item}) + (1 \mid \text{id})$, where the group slope varying by item captures item-specific DIF.
- **Polytomous uniform DIF** (`dif_type = "uniform"`): The baseline response $\mid \text{thres}(\text{gr} = \text{item}) \sim 1 + (1 \mid \text{id})$ becomes response $\mid \text{thres}(\text{gr} = \text{item}) \sim 1 + \text{group}:\text{item} + (1 \mid \text{id})$.
- **Polytomous non-uniform DIF** (`dif_type = "non-uniform"`): The baseline becomes response $\mid \text{thres}(\text{gr} = \text{item_group}) \sim 1 + (1 \mid \text{id})$, where `item_group = interaction(item, group)`. Each `item × group` combination gets its own thresholds. DIF effects are the differences between group-specific thresholds for each item, computed draw-by-draw from the posterior.

DIF effects are summarized using:

Probability of Direction (pd) The proportion of the posterior on the dominant side of zero. Values > 0.975 indicate strong directional evidence.

ROPE The Region of Practical Equivalence (Kruschke, 2018). If $> 95\%$ the DIF effect is practically negligible. If $> 95\%$ outside, the effect is practically significant.

Credible Interval If the CI excludes zero, there is evidence of DIF at the specified credibility level.

Value

A list with the following elements:

summary A [tibble](#) with one row per item (for uniform DIF) or per item $\times$ threshold (for non-uniform DIF) containing: item, optionally threshold, dif_estimate (posterior mean), dif_lower, dif_upper (credible interval), dif_sd (posterior SD), pd (probability of direction), rope_percentage (proportion inside ROPE), and flag (classification).

dif_draws A matrix of posterior draws for the DIF effects (draws $\times$ effects), for further analysis.

dif_model The fitted DIF brmsfit object (if refit = TRUE), or NULL.

dif_formula The brmsformula used for the DIF model.

baseline_model The original baseline model.

plot A [ggplot](#) forest plot of DIF effects with credible intervals and ROPE.

References

Kruschke, J. K. (2018). Rejecting or accepting parameter values in Bayesian estimation. *Advances in Methods and Practices in Psychological Science*, 1(2), 270–280.

Bürkner, P.-C. (2021). Bayesian Item Response Modeling in R with brms and Stan. *Journal of Statistical Software*, 100, 1–54. doi:10.18637/jss.v100.i05

See Also

[infit_statistic](#) for item fit, [q3_statistic](#) for local dependence, [brm](#), [hypothesis](#).

Examples

```
library(brms)
library(dplyr)
library(tidyr)
library(tibble)

# --- Dichotomous Rasch with DIF testing ---

set.seed(123)
df <- expand.grid(id = 1:200, item = paste0("I", 1:10)) %>%
  mutate(
    gender = rep(sample(c("M", "F"), 200, TRUE), each = 10),
    theta = rep(rnorm(200), each = 10),
    delta = rep(seq(-2, 2, length.out = 10), 200),
    dif = ifelse(item == "I3" & gender == "F", 1.0,
                 ifelse(item == "I7" & gender == "F", -0.8, 0)),
    p = plogis(theta - delta - dif),
    response = rbinom(n(), 1, p)
  )

fit_base <- brm(
  response ~ 1 + (1 | item) + (1 | id),
  data = df,
  family = bernoulli(),
  chains = 4,
  cores = 2, # use more cores if you have
  iter = 1000 # use at least 2000
)
```

```

dif_result <- dif_statistic(
  model      = fit_base,
  group_var  = gender,
  data       = df
)

dif_result$summary
dif_result$plot

# --- Partial Credit Model: uniform DIF ---

df_pcm <- eRm::pcmdat2 %>%
  mutate(across(everything(), ~ .x + 1)) %>%
  rownames_to_column("id") %>%
  mutate(gender = sample(c("M", "F"), n(), TRUE)) %>%
  pivot_longer(!c(id, gender),
    names_to = "item", values_to = "response")

fit_pcm <- brm(
  response | thres(gr = item) ~ 1 + (1 | id),
  data     = df_pcm,
  family   = acat,
  chains    = 4,
  cores     = 2, # use more cores if you have
  iter      = 1000 # use at least 2000
)

# Uniform DIF (default): one shift per item
dif_uni <- dif_statistic(fit_pcm, group_var = gender, data = df_pcm)
dif_uni$plot

# Non-uniform DIF: threshold-level effects
dif_nu <- dif_statistic(fit_pcm, group_var = gender, data = df_pcm,
  dif_type = "non-uniform")

dif_nu$summary
dif_nu$plot

```

fit_statistic_pcm

Posterior Predictive Item Fit Statistic for Bayesian IRT Models

Description

Computes posterior predictive item (or person) fit statistics for Bayesian IRT models fitted with **brms**. For each posterior draw, observed and replicated data are compared via a user-supplied criterion function, grouped by item, person, or any other variable. Posterior predictive p-values can then be derived from the output to assess fit.

Usage

```
fit_statistic_pcm(model, criterion, group, ndraws_use = NULL)
```

Arguments

<code>model</code>	A fitted <code>brmsfit</code> object.
<code>criterion</code>	A function with signature <code>function(y, p)</code> that computes a pointwise fit criterion. For ordinal and categorical models, <code>y</code> is the observed (or replicated) response category and <code>p</code> is the model-predicted probability of that category. For binary models, <code>y</code> is the binary response and <code>p</code> is the predicted probability of success.
<code>group</code>	An unquoted variable name (e.g., <code>item</code> or <code>id</code>) indicating the grouping variable over which the fit statistic is aggregated. Typically <code>item</code> for item fit or <code>id</code> for person fit.
<code>ndraws_use</code>	Optional positive integer. If specified, a random subset of posterior draws of this size is used, which can speed up computation for large models. If <code>NULL</code> (the default), all draws are used.

Details

The function implements the posterior predictive checking approach for item fit described in Bürkner (2020). The procedure works as follows:

1. Draw posterior expected category probabilities via `posterior_epred` and posterior predicted responses via `posterior_predict`.
2. For ordinal or categorical models (3D array output from `posterior_epred`), extract the probability assigned to the observed response category and to the replicated response category for each draw and observation.
3. Apply the user-supplied `criterion` function to compute pointwise fit values for both observed and replicated data.
4. Aggregate (sum) the criterion values within each level of group and each posterior draw.

A common choice for ordinal IRT models is the categorical log-likelihood `criterion(y, p) log(p)`. For binary (e.g., dichotomous Rasch) models, the Bernoulli log-likelihood `criterion(y, p) y * log(p) + (1 - y) * log(1 - p)` may be used instead.

Value

A `tibble` with the following columns:

group The grouping variable (e.g., item name or person id).

draw Integer index of the posterior draw.

crit The observed fit statistic (criterion applied to observed data) summed within each group and draw.

crit_rep The replicated fit statistic (criterion applied to posterior predicted data) summed within each group and draw.

crit_diff The difference `crit_rep - crit`.

The output is grouped by the grouping variable. Posterior predictive p-values can be obtained by computing `mean(crit_rep > crit)` within each group.

References

- Bürkner, P.-C. (2020). Analysing Standard Progressive Matrices (SPM-LS) with Bayesian Item Response Models. *Journal of Intelligence*, 8(1). doi:10.3390/jintelligence8010005
- Bürkner, P.-C. (2021). Bayesian Item Response Modeling in R with brms and Stan. *Journal of Statistical Software*, 100, 1–54. doi:10.18637/jss.v100.i05

See Also

[fit_statistic_rm](#) for dichotomous Rasch models, [posterior_epred](#) for expected predictions, [posterior_predict](#) for posterior predictive samples, [pp_check](#) for graphical posterior predictive checks.

Examples

```
library(brms)
library(dplyr)
library(tidyr)
library(tibble)

# --- Polytomous Rasch (Partial Credit Model) ---

# Prepare data in long format
df_pcm <- eRm::pcmdat2 %>%
  mutate(across(everything(), ~ .x + 1)) %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

# Fit a Partial Credit Model using the adjacent category family
fit_pcm <- brm(
  response | thres(gr = item) ~ 1 + (1 | id),
  data      = df_pcm,
  family    = acat,
  chains    = 4,
  cores     = 1, # use more cores if you have
  iter      = 500 # use at least 2000
)

# Categorical log-likelihood criterion (for polytomous models)
ll_categorical <- function(y, p) log(p)

# Compute item fit statistics
item_fit <- fit_statistic_pcm(
  model      = fit_pcm,
  criterion  = ll_categorical,
  group      = item,
  ndraws_use = 100 # use at least 500
```

```

)

# Summarise: posterior predictive p-values per item
item_fit %>%
  group_by(item) %>%
  summarise(
    observed = mean(crit),
    replicated = mean(crit_rep),
    ppp = mean(crit_rep > crit)
  )

# Use ggplot2 to make a histogram
library(ggplot2)
item_fit %>%
  ggplot(aes(crit_diff)) +
  geom_histogram(aes(fill = ifelse(crit_diff > 0, "above", "below"))) +
  facet_wrap("item") +
  theme_bw() +
  theme(legend.position = "none")

# Compute person fit statistics
person_fit <- fit_statistic_pcm(
  model = fit_pcm,
  criterion = ll_categorical,
  group = id,
  ndraws_use = 100 # use at least 500
)

```

fit_statistic_rm

Posterior Predictive Item Fit Statistic for Binary Bayesian IRT Models

Description

Computes posterior predictive item (or person) fit statistics for dichotomous Bayesian IRT models fitted with **brms**. For each posterior draw, observed and replicated data are compared via a user-supplied criterion function, grouped by item, person, or any other variable. Posterior predictive p-values can then be derived from the output to assess fit.

Usage

```
fit_statistic_rm(model, criterion, group, ndraws_use = NULL)
```

Arguments

model A fitted **brmsfit** object with a binary response (e.g., family = bernoulli()).

criterion	A function with signature <code>function(y, p)</code> that computes a pointwise fit criterion, where <code>y</code> is the binary response (0 or 1) and <code>p</code> is the predicted probability of success. A common choice is the Bernoulli log-likelihood: <code>function(y, p) y * log(p) + (1 - y) * log(1 - p)</code> .
group	An unquoted variable name (e.g., <code>item</code> or <code>id</code>) indicating the grouping variable over which the fit statistic is aggregated. Typically <code>item</code> for item fit or <code>id</code> for person fit.
ndraws_use	Optional positive integer. If specified, a random subset of posterior draws of this size is used, which can speed up computation for large models. If <code>NULL</code> (the default), all draws are used.

Details

This function is the binary-response counterpart of `fit_statistic_pcm`, which handles polytomous (ordinal / categorical) models. For dichotomous models, `posterior_epred()` returns a 2D matrix (`S x N`) of success probabilities, so the criterion function receives the observed binary response and the corresponding probability directly.

The procedure follows the posterior predictive checking approach described in Bürkner (2020):

1. Draw posterior expected success probabilities via `posterior_epred` and posterior predicted binary responses via `posterior_predict`.
2. Apply the user-supplied criterion function pointwise to both observed and replicated data paired with the predicted probabilities.
3. Aggregate (sum) the criterion values within each level of group and each posterior draw.

The standard criterion for binary models is the Bernoulli log-likelihood:

$$\ell(y, p) = y \log(p) + (1 - y) \log(1 - p).$$

Value

A `tibble` with the following columns:

group The grouping variable (e.g., item name or person id).

draw Integer index of the posterior draw.

crit The observed fit statistic (criterion applied to observed data) summed within each group and draw.

crit_rep The replicated fit statistic (criterion applied to posterior predicted data) summed within each group and draw.

crit_diff The difference `crit_rep - crit`.

The output is grouped by the grouping variable. Posterior predictive p-values can be obtained by computing `mean(crit_rep > crit)` within each group.

References

- Bürkner, P.-C. (2020). Analysing Standard Progressive Matrices (SPM-LS) with Bayesian Item Response Models. *Journal of Intelligence*, 8(1). doi:10.3390/jintelligence8010005
- Bürkner, P.-C. (2021). Bayesian Item Response Modeling in R with brms and Stan. *Journal of Statistical Software*, 100, 1–54. doi:10.18637/jss.v100.i05

See Also

[fit_statistic_pcm](#) for polytomous (ordinal/categorical) models, [posterior_epred](#) for expected predictions, [posterior_predict](#) for posterior predictive samples, [pp_check](#) for graphical posterior predictive checks.

Examples

```
library(brms)
library(dplyr)
library(tidyr)
library(tibble)

# --- Dichotomous Rasch Model ---

# Prepare binary response data in long format
df_rm <- eRm::raschdat3 %>%
  as.data.frame() %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

# Fit a dichotomous Rasch model
fit_rm <- brm(
  response ~ 1 + (1 | item) + (1 | id),
  data = df_rm,
  family = bernoulli(),
  chains = 4,
  cores = 1, # use more cores if you have
  iter = 500 # use at least 2000
)

# Bernoulli log-likelihood criterion
ll_bernoulli <- function(y, p) y * log(p) + (1 - y) * log(1 - p)

# Compute item fit statistics
item_fit <- fit_statistic_rm(
  model = fit_rm,
  criterion = ll_bernoulli,
  group = item,
  ndraws_use = 100 # use at least 500
)

# Summarise: posterior predictive p-values per item
item_fit %>%
  group_by(item) %>%
  summarise(
    observed = mean(crit),
    replicated = mean(crit_rep),
    ppp = mean(crit_rep > crit)
  )

# Use ggplot2 to make a histogram
library(ggplot2)
```

```

item_fit %>%
  ggplot(aes(crit_diff)) +
  geom_histogram(aes(fill = ifelse(crit_diff > 0, "above", "below"))) +
  facet_wrap("item") +
  theme_bw() +
  theme(legend.position = "none")

# Compute person fit statistics
person_fit <- fit_statistic_rm(
  model      = fit_rm,
  criterion  = ll_bernoulli,
  group      = id,
  ndraws_use = 100 # use at least 500
)

person_fit %>%
  group_by(id) %>%
  summarise(
    observed   = mean(crit),
    replicated = mean(crit_rep),
    ppp        = mean(crit_rep > crit)
  )

# --- 1PL model with item-specific intercepts ---

# Alternative parameterisation with fixed item effects
fit_1pl <- brm(
  response ~ 0 + item + (1 | id),
  data     = df_rm,
  family   = bernoulli(),
  chains   = 4,
  cores    = 1, # use more cores if you have
  iter     = 500 # use at least 2000
)

item_fit_1pl <- fit_statistic_rm(
  model      = fit_1pl,
  criterion  = ll_bernoulli,
  group      = item,
  ndraws_use = 100 # use at least 500
)

item_fit_1pl %>%
  group_by(item) %>%
  summarise(
    observed   = mean(crit),
    replicated = mean(crit_rep),
    ppp        = mean(crit_rep > crit)
  )

```

hurdle_acat

*Hurdle Partial Credit Model Custom brms Family***Description**

A custom **brms** family implementing a hurdle Rasch partial credit model (hPCM). A Bernoulli logit gate models $P(Y = 0)$; conditional on $Y > 0$, an acat-logit partial credit process governs transitions among the positive categories $1, \dots, K - 1$.

Usage

```
hurdle_acat()
```

Details

The two submodels can be interpreted as a *hurdle* (presence / absence of a symptom or behaviour, sometimes called "susceptibility") and a *partial credit* severity model (frequency / intensity given presence), in the spirit of Magnus and Garnier-Villarreal (2022). Person random effects on the two submodels can be modelled as correlated (via brms's `(1 | g| id)` syntax), allowing the data to inform whether susceptibility and severity are distinct latent constructs or essentially the same trait.

Value

A customfamily object suitable for the family argument of `brm`. The companion stanvars (the Stan code for the custom lpmf) must be passed via the `stanvars` argument; see [hurdle_acat_stanvars](#).

Usage

```
library(brms)
fit <- brm(
  bf(
    response | thres(gr = item) ~ 1 + (1 | g| id),
    hu ~ 0 + factor(item) + (1 | g| id)
  ),
  data      = dat,
  family    = hurdle_acat(),
  stanvars  = hurdle_acat_stanvars(),
  ...
)
```

brms compatibility note

The custom family relies on brms's native ordinal infrastructure (`specials = "ordinal" + thres(gr = item)`). On CRAN brms 2.23.x this combination emits invalid Stan code for custom ordinal families with grouped thresholds. A patched branch is available:

```
devtools::install_github(
  "rpsychologist/brms@fix-custom-ordinal-grouped-thres"
)
```

Sign convention

The brms formula $hu \sim \dots + (1 \mid id)$ adds the person random effect to the logit of $hu = P(Y = 0)$, so higher values of the person random effect on hu mean *more* zeros (lower susceptibility). This is the opposite sign of a Stan-style parameterisation in which higher `theta_gate` means fewer zeros (higher susceptibility). The two parameterisations imply identical likelihoods; only the sign of the recovered correlation between the two person random effects is flipped relative to a "susceptibility x severity" labelling. No inferences about person ranking, gate probabilities, or severity probabilities are affected.

Author(s)

Kristoffer Magnusson

References

Magnus, B. E. & Garnier-Villareal, M. (2022). A multidimensional zero-inflated graded response model for ordinal symptom data. *Psychological Methods*, 27(2), 261-279. doi:10.1037/met0000395

See Also

[hurdle_acat_stanvars](#) for the Stan function block, [infit_statistic_hpcm](#) for submodel-specific item infit, [q3_statistic_hpcm](#) for submodel-specific Q3 residual correlations.

infit_post

Summarize and Plot Posterior Predictive Infit Statistics

Description

Postprocesses the output of [infit_statistic](#) to produce summary tables of posterior predictive infit statistics and a combined slab + interval plot comparing observed infit values to the posterior predictive distribution.

Usage

```
infit_post(infit_draws, ci = 0.84)
```

Arguments

<code>infit_draws</code>	A data frame (or tibble) as returned by infit_statistic containing at minimum the columns <code>item</code> , <code>infit</code> (observed infit per draw), and <code>infit_rep</code> (replicated infit per draw).
<code>ci</code>	Numeric in (0, 1). Width of the credible interval used for the posterior predictive HDI and the slab display. Default is 0.84.

Details

Two complementary summary tables are provided:

summary Reports the posterior mean observed and replicated infit values alongside the posterior predictive p-value (ppp). The ppp is the proportion of draws where the replicated infit exceeds the observed infit. Under good fit, the ppp should be near 0.5. A ppp near 0 indicates the observed infit is consistently larger than expected (underfit); a ppp near 1 indicates it is consistently smaller (overfit).

hdi Reports the probability that the observed infit falls above (underfit) or below (overfit) the HDI of the replicated distribution. This provides a more distributional assessment than the ppp alone.

The plot uses two layers from the **ggdist** package:

stat_slab Displays the posterior predictive (replicated) infit distribution as a filled density slab per item, shaded by credible interval level.

stat_slabinterval Displays the observed infit distribution per item as a semi-transparent slab with point and interval summaries.

Under good model fit, the observed infit distribution should overlap substantially with the replicated distribution. Items where the observed distribution sits systematically above the replicated HDI indicate underfit (more variation than expected); items below indicate overfit (less variation than expected).

Value

A list with three elements:

summary A **tibble** with one row per item containing: `item`, `infit_obs` (posterior mean of observed infit), `infit_rep` (posterior mean of replicated infit), and `infit_ppp` (posterior predictive p-value: proportion of draws where the replicated infit exceeds the observed infit). Values near 0.5 indicate good fit; values near 0 suggest underfit; values near 1 suggest overfit.

hdi A **tibble** with one row per item containing: `item`, `underfit` (posterior probability that the observed infit exceeds the upper HDI bound of the replicated distribution), and `overfit` (posterior probability that the observed infit falls below the lower HDI bound).

plot A **ggplot** object showing the posterior predictive distribution of replicated infit (grey filled slab) overlaid with the observed infit distribution (coloured slab + interval), with a dashed reference line at 1 (perfect fit).

See Also

[infit_statistic](#), [item_restscore_post](#).

Examples

```
## Not run:
library(brms)
library(ggplot2)
```

```
# Assuming fit_pcm is a fitted brmsfit object
infit_draws <- infit_statistic(fit_pcm)

result <- infit_post(infit_draws)
result$summary
result$hdi
result$plot

## End(Not run)
```

infit_statistic

*Posterior Predictive Infit Statistic for Bayesian IRT Models***Description**

Computes a Bayesian analogue of the conditional item infit statistic (as described in Christensen, Kreiner & Mesbah, 2013) for Rasch-family models fitted with **brms**. For each posterior draw, expected values and variances are derived from the category probabilities returned by [posterior_epred](#), and variance-weighted standardised residuals are computed for both observed and replicated data. The result can be summarised into posterior predictive p-values to assess item fit.

Usage

```
infit_statistic(
  model,
  item_var = item,
  person_var = id,
  ndraws_use = NULL,
  outfit = FALSE
)
```

Arguments

model	A fitted brmsfit object from an ordinal IRT model (e.g., family = acat for a partial credit model or family = bernoulli() for a dichotomous Rasch model).
item_var	An unquoted variable name identifying the item grouping variable in the model data (e.g., item).
person_var	An unquoted variable name identifying the person grouping variable in the model data (e.g., id).
ndraws_use	Optional positive integer. If specified, a random subset of posterior draws of this size is used. If NULL (the default), all draws are used.
outfit	Logical. If TRUE, outfit statistics are computed alongside infit. Default is FALSE (infit only), since outfit is highly sensitive to outliers and rarely recommended for Rasch diagnostics.

Details

The procedure adapts the conditional infit/outfit statistics (Christensen et al., 2013; Kreiner & Christensen, 2011; Müller, 2020) to the Bayesian framework:

1. For each posterior draw s , category probabilities $P^{(s)}(X_{vi} = c)$ are obtained from `posterior_epred`.
2. The conditional expected value and variance for each observation are computed as:

$$E_{vi}^{(s)} = \sum_c c \cdot P^{(s)}(X_{vi} = c)$$

$$Var_{vi}^{(s)} = \sum_c (c - E_{vi}^{(s)})^2 \cdot P^{(s)}(X_{vi} = c)$$

3. Standardised squared residuals are:

$$Z_{vi}^{2(s)} = (X_{vi} - E_{vi}^{(s)})^2 / Var_{vi}^{(s)}$$

4. The infit statistic for item i is the variance-weighted mean of Z^2 across persons:

$$Infit_i^{(s)} = \frac{\sum_v Var_{vi}^{(s)} Z_{vi}^{2(s)}}{\sum_v Var_{vi}^{(s)}}$$

5. If requested, the outfit is the unweighted mean of Z^2 .

Value

A `tibble` with the following columns:

item The item identifier.

draw Integer index of the posterior draw.

infit The observed infit statistic for that item and draw.

infit_rep The replicated infit statistic (based on posterior predicted data) for that item and draw.

outfit (Only if `outfit = TRUE`) The observed outfit statistic for that item and draw.

outfit_rep (Only if `outfit = TRUE`) The replicated outfit statistic for that item and draw.

The output is grouped by the item variable. Posterior predictive p-values can be obtained by computing, e.g., `mean(infit_rep > infit)` within each item.

References

- Christensen, K. B., Kreiner, S. & Mesbah, M. (Eds.) (2013). *Rasch Models in Health*. Iste and Wiley, pp. 86–90.
- Kreiner, S. & Christensen, K. B. (2011). Exact evaluation of Bias in Rasch model residuals. *Advances in Mathematics Research*, 12, 19–40.
- Müller, M. (2020). Item fit statistics for Rasch analysis: can we trust them? *Journal of Statistical Distributions and Applications*, 7(1). doi:10.1186/s40488020001087

See Also

[fit_statistic_pcm](#) for a general-purpose posterior predictive fit statistic with user-supplied criterion functions, [fit_statistic_rm](#) for a general-purpose posterior predictive fit statistic with user-supplied criterion functions, [posterior_epred](#), [posterior_predict](#).

Examples

```
library(brms)
library(dplyr)
library(tidyr)
library(tibble)

# --- Partial Credit Model (polytomous) ---

df_pcm <- eRm::pcmdat2 %>%
  mutate(across(everything(), ~ .x + 1)) %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_pcm <- brm(
  response | thres(gr = item) ~ 1 + (1 | id),
  data = df_pcm,
  family = acat,
  chains = 4,
  cores = 1, # use more cores if you have
  iter = 500 # use at least 2000
)

# Compute infit per item
item_infit <- infit_statistic(
  model = fit_pcm,
  ndraws_use = 100 # use at least 500
)

# Post-process draws
infit_results <- infit_post(item_infit)
infit_results$summary
infit_results$hdi
infit_results$plot

# --- Dichotomous Rasch Model ---

df_rm <- eRm::raschdat3 %>%
  as.data.frame() %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_rm <- brm(
  response ~ 1 + (1 | item) + (1 | id),
  data = df_rm,
  family = bernoulli(),
  chains = 4,
```

```

cores = 1, # use more cores if you have
iter  = 500 # use at least 2000
)

item_infit_rm <- infit_statistic(
  model      = fit_rm,
  ndraws_use = 100 # use at least 500
)

# Post-process draws
infit_results <- infit_post(item_infit_rm)
infit_results$summary
infit_results$hdi
infit_results$plot

```

infit_statistic_hpcm *Posterior Predictive Infit Statistic for the Hurdle Partial Credit Model*

Description

Computes conditional item infit statistics separately for the two submodels of a hurdle partial credit model fitted with **brms** using the [hurdle_acat](#) custom family: (i) the *hurdle* submodel for $P(Y > 0)$ (Bernoulli) and (ii) the *partial credit* severity submodel for $P(Y = k \mid Y > 0)$ on the positive categories. For each posterior draw, expected values and variances are derived from the submodel-specific category probabilities, and variance-weighted standardised residuals are computed for both observed and replicated data.

Usage

```

infit_statistic_hpcm(
  model,
  item_var = item,
  person_var = id,
  ndraws_use = NULL,
  outfit = FALSE
)

```

Arguments

model	A fitted brmsfit object using the hurdle_acat custom family (i.e., <code>posterior_epred</code> returns an $S \times N \times K_{\text{total}}$ array whose first category is the hurdle / zero probability).
item_var	An unquoted variable name identifying the item grouping variable in the model data (e.g., <code>item</code>).

person_var	An unquoted variable name identifying the person grouping variable in the model data (e.g., id).
ndraws_use	Optional positive integer. If specified, a random subset of posterior draws of this size is used. If NULL (the default), all draws are used.
outfit	Logical. If TRUE, outfit statistics are computed alongside infit. Default is FALSE.

Details

The hurdle PCM splits the generative process into:

1. A Bernoulli hurdle with $hu = P(Y = 0)$.
2. A partial credit / acat-logit severity process over the positive categories $1, \dots, K - 1$, applied only when the hurdle is crossed.

posterior_epred for the hurdle_acat family returns an $S \times N \times K_{\text{total}}$ array whose first category is hu and whose remaining categories are $(1 - hu) \cdot P_{sev}(k)$. The two submodel infits are computed as follows:

Hurdle submodel. All observations contribute. The Bernoulli moments are $E_{hurdle} = 1 - hu$ and $Var_{hurdle} = hu \cdot (1 - hu)$. Observed and replicated scores are $1[Y_{obs} > 0]$ and $1[Y_{rep} > 0]$ with Y_{rep} obtained from [posterior_predict](#).

Partial credit submodel. Only observations with $Y_{obs} > 0$ contribute. Conditional probabilities are

$$P(Y = k \mid Y > 0) = \text{epred}[, k + 1] / (1 - hu), \quad k = 1, \dots, K - 1.$$

The conditional expected value and variance use category scores $1, \dots, K - 1$. Replicated severity values are drawn independently for each (draw, observation) from these conditional probabilities via inverse-CDF sampling, so the partial credit PPC is not contaminated by hurdle misfit.

Within each submodel the infit per item is

$$\text{Infit}_i^{(s)} = \sum_v (X_{vi} - E_{vi}^{(s)})^2 / \sum_v Var_{vi}^{(s)},$$

with the sum restricted to the rows the submodel applies to (all rows for the hurdle; rows with $Y_{obs} > 0$ for partial credit).

Value

A list with two elements, each a [tibble](#) in the same format as the output of [infit_statistic](#) (and directly compatible with [infit_post](#)):

hurdle Item infit for the Bernoulli hurdle submodel on $1[Y > 0]$, evaluated on *all* observations.

pcm Item infit for the partial credit severity submodel on $P(Y = k \mid Y > 0)$, evaluated only on the observations with $Y_{obs} > 0$.

References

- Christensen, K. B., Kreiner, S. & Mesbah, M. (Eds.) (2013). *Rasch Models in Health*. Iste and Wiley, pp. 86–90.
- Kreiner, S. & Christensen, K. B. (2011). Exact evaluation of bias in Rasch model residuals. *Advances in Mathematics Research*, 12, 19–40.
- Magnus, B. E. & Garnier-Villarreal, M. (2022). A multidimensional zero-inflated graded response model for ordinal symptom data. *Psychological Methods*, 27(2), 261–279. doi:10.1037/met0000395

See Also

[infit_statistic](#) for the single-submodel version, [infit_post](#) for summarising and plotting the draws, [q3_statistic_hpcm](#) for hPCM Q3 residual correlations.

item_parameters	<i>Extract Item Parameters from a Bayesian Rasch Model</i>
-----------------	--

Description

Extracts item difficulty (threshold) parameters from a fitted Bayesian Rasch model. Returns a simple location table in both long and wide formats, a full summary with posterior SEs and HDCIs, item-level information, threshold ordering diagnostics, and optionally the full posterior draws matrix.

Usage

```
item_parameters(
  model,
  item_var = item,
  person_var = id,
  draws = FALSE,
  center = TRUE,
  prob = 0.95
)
```

Arguments

model	A fitted brmsfit object. Supported parameterisations: Polytomous ordinal (PCM) e.g., family = acat with thres(gr = item), producing item-specific thresholds. Dichotomous Rasch (random items) e.g., response ~ 1 + (1 item) + (1 id) with family = bernoulli(). Dichotomous 1PL (fixed items) e.g., response ~ 0 + item + (1 id) with family = bernoulli().
item_var	An unquoted variable name identifying the item grouping variable in the model data. Default is item.

person_var	An unquoted variable name identifying the person grouping variable in the model data. Default is id.
draws	Logical. If TRUE, a draws matrix of full posterior draws is included in the output. Default is FALSE.
center	Logical. If TRUE (the default), item parameters are shifted so that the grand mean of all threshold locations is zero, matching the frequentist CML convention and the centering used in plot_targeting .
prob	Numeric in (0,1). Width of the highest density continuous interval (HDCI) reported in the summary. Default is 0.95.

Details

Dichotomous models with random item effects ($\text{response} \sim 1 + (1 \mid \text{item}) + (1 \mid \text{id})$) parameterise item difficulty as $\delta_i = -(b_0 + r_i)$ where b_0 is the global intercept and r_i is the item random effect. Models with fixed item effects ($\text{response} \sim 0 + \text{item} + (1 \mid \text{id})$) parameterise difficulty as $\delta_i = -b_i$.

Polytomous (acat/PCM) models with grouped thresholds ($\text{thres}(\text{gr} = \text{item})$) directly estimate item-specific threshold parameters. Each row in the long output represents one threshold within one item; each row in the wide output represents one item.

Item information is computed from the posterior mean item parameters using the standard Rasch/PCM information formulae:

Dichotomous $I_i(\theta) = P_i(\theta)Q_i(\theta)$ where $P_i = \text{logistic}(\theta - \delta_i)$.

Polytomous (PCM) $I_i(\theta) = \sum_c (c - E_i)^2 P_{ic}(\theta)$ where $E_i = \sum_c c \cdot P_{ic}$ is the expected score for item i .

Threshold ordering: In the partial credit model, disordered thresholds ($\tau_{k+1} \leq \tau_k$) indicate that the probability of responding in the intermediate category never exceeds both adjacent categories — the category is empirically "absorbed". This does not necessarily indicate misfit (see Adams et al., 2012), but may suggest response categories should be collapsed. The `prob_disordered` column reports the posterior probability of at least one disordered pair per item, providing a Bayesian alternative to post-hoc threshold checks.

Value

A list with the following elements:

locations A [tibble](#) in long format with one row per item (dichotomous) or per item-threshold (polytomous), containing `item`, `threshold` (integer, always 1 for dichotomous), and `location` (posterior mean on the logit scale).

locations_wide A [tibble](#) in wide format with one row per item, sorted by mean location. For polytomous models, threshold columns are named `t1`, `t2`, etc., and a `location` column gives the mean across thresholds. For dichotomous models, only `item` and `location` columns are present.

summary A [tibble](#) extending the long `locations` with: `se` (posterior SD), `hdc_lower` and `hdc_upper` (highest density continuous interval bounds at the level specified by `prob`), and `n_eff` (effective sample size for the parameter).

item_information A [tibble](#) with one row per item containing: item, location (mean item location, i.e. mean of thresholds for polytomous items), info_at_location (Fisher information at the item's own location), and max_info (maximum Fisher information across theta). For Rasch dichotomous items, both are 0.25. For polytomous items, information depends on the number and spacing of thresholds.

threshold_order (Polytomous models only) A [tibble](#) with one row per item containing: item, n_thresholds, ordered (logical: are all thresholds in ascending order?), and prob_disordered (posterior probability that at least one pair of adjacent thresholds is disordered, i.e. $\tau_{k+1} \leq \tau_k$). NULL for dichotomous models.

person_sd A [tibble](#) with one row containing: mean, sd, hdci_lower, hdci_upper — the posterior summary of the person-level standard deviation parameter σ_θ .

draws_matrix (Only if draws = TRUE) A numeric matrix with rows = thresholds (named "item[threshold]") and columns = posterior draws. For dichotomous models, row names are item labels only.

References

Adams, R. J., Wu, M. L., & Wilson, M. (2012). The Rasch rating model and the disordered threshold controversy. *Educational and Psychological Measurement*, 72(4), 547–573. doi:10.1177/0013164411432166

Bürkner, P.-C. (2021). Bayesian Item Response Modeling in R with brms and Stan. *Journal of Statistical Software*, 100, 1–54. doi:10.18637/jss.v100.i05

See Also

[person_parameters](#), [plot_targeting](#), [plot_ipf](#), [posterior_to_prior](#).

Examples

```
library(brms)
library(dplyr)
library(tidyr)
library(tibble)

# --- Partial Credit Model ---

df_pcm <- eRm::pcmdat2 %>%
  mutate(across(everything(), ~ .x + 1)) %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_pcm <- brm(
  response | thres(gr = item) ~ 1 + (1 | id),
  data = df_pcm,
  family = acat,
  chains = 4, cores = 2, iter = 1000 # use more iter and cores
)

ip <- item_parameters(fit_pcm)

# Long format: one row per threshold
```

```

ip$locations

# Wide format: one row per item, easy to scan
ip$locations_wide

# Full summary with SE and HDCI
ip$summary

# Item information
ip$item_information

# Threshold ordering diagnostic
ip$threshold_order

# Person SD
ip$person_sd

# With full posterior draws
ip_draws <- item_parameters(fit_pcm, draws = TRUE)
dim(ip_draws$draws_matrix)

# --- Dichotomous Rasch ---

df_rm <- eRm::raschdat3 %>%
  as.data.frame() %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_rm <- brm(
  response ~ 1 + (1 | item) + (1 | id),
  data = df_rm,
  family = bernoulli(),
  chains = 4, cores = 2, iter = 1000 # use more iter and cores
)

ip_rm <- item_parameters(fit_rm)
# Wide and long are equivalent for dichotomous:
ip_rm$locations
ip_rm$locations_wide
ip_rm$summary

```

item_parameters_hpcm *Extract Item Parameters from a Hurdle Partial Credit Model*

Description

Extracts item parameters from a fitted hurdle Rasch partial credit model (`family = hurdle_acat()`), returning a per-submodel breakdown: hurdle item difficulties (Bernoulli logit on $P(Y > 0)$) and

partial credit thresholds. Each submodel's output mirrors the shape of `item_parameters` so existing plotting and downstream functions can be applied directly to `res$hurdle` or `res$pcm`.

Usage

```
item_parameters_hpcm(
  model,
  item_var = item,
  person_var = id,
  draws = FALSE,
  center = TRUE,
  prob = 0.95
)
```

Arguments

model	A fitted <code>brmsfit</code> object using the <code>hurdle_acat</code> custom family. The recommended formula is <pre>bf(response thres(gr = item) ~ 1 + (1 g id), hu ~ 0 + factor(item) + (1 g id))</pre>
item_var	An unquoted variable name identifying the item grouping variable in the model data. Default is <code>item</code> .
person_var	An unquoted variable name identifying the person grouping variable in the model data. Default is <code>id</code> .
draws	Logical. If TRUE, a draws matrix of full posterior draws is included in each submodel's output. Default is FALSE.
center	Logical. If TRUE (the default), item parameters are shifted within each submodel so their mean is zero, matching the convention in <code>item_parameters</code> . Person parameters reported by <code>person_parameters_hpcm</code> use the same shifts.
prob	Numeric in (0, 1). Width of the highest density continuous interval (HDCI) reported in the summary. Default is 0.95.

Details

Hurdle submodel. The hurdle linear predictor is $\text{logit}(hu) = \delta_{\text{hurdle},i} + \tilde{r}_v$ with $hu = P(Y = 0)$; higher $\delta_{\text{hurdle},i}$ means more zeros, so this is reported directly as the item "location" (harder = higher value, standard Rasch convention for the Bernoulli on $P(Y > 0)$). Hurdle item information is the Bernoulli variance $p(1 - p)$ evaluated at $\theta = \delta_{\text{hurdle},i}$, which is exactly $1/4$.

Partial credit submodel. Thresholds τ_{ik} are the per-item PCM threshold parameters. Item information uses the standard PCM formula; threshold ordering diagnostics are the same as for `item_parameters`.

Sign of the trait correlation. The brms model reports `cor_id__Intercept__hu_Intercept`, which is the correlation between the brms random effects `r_id[, Intercept]` and `r_id[, hu_Intercept]`. The latter has the opposite sign of the conventional "susceptibility" person trait θ_{hurdle} (because

higher values of the brms random effect mean more zeros, i.e., lower susceptibility). The correlation reported here is therefore $-\text{cor}_{brms}$, so that positive values mean higher susceptibility goes with higher severity. The marginal SDs are unchanged by the sign flip.

Centering. When `center = TRUE`, the hurdle item difficulties are shifted by their mean and the PCM thresholds are shifted by the grand mean of all PCM thresholds. The same shifts are applied to the corresponding person traits in `person_parameters_hpcm`, preserving the underlying likelihood.

Value

A list with three elements:

- `hurdle` A list with the same structure as the output of `item_parameters` applied to a dichotomous Rasch model: `locations`, `locations_wide`, `summary`, `item_information`, `person_sd`, optionally `draws_matrix`. `location` is the brms posterior mean of `b_hu_factoritem<i>`; higher values mean more zeros (a harder hurdle to cross).
- `pcm` A list with the same structure as `item_parameters` applied to a PCM: `locations` (long), `locations_wide` (`t1`, `t2`, ..., `location`), `summary`, `item_information`, `threshold_order`, `person_sd`, optionally `draws_matrix`. `location` columns are posterior means of `b_Intercept[<item>`, `<k>]`.
- `correlation` A `tibble` with `mean`, `sd`, `hdci_lower`, `hdci_upper` summarising the posterior of $\rho(\theta_{hurdle}, \theta_{pcm})$. This is the brms correlation `cor_id__Intercept__hu_Intercept` with its sign flipped to match the "higher = more presence" convention used for the hurdle person trait (see Details).

References

- Magnus, B. E. & Garnier-Villareal, M. (2022). A multidimensional zero-inflated graded response model for ordinal symptom data. *Psychological Methods*, 27(2), 261-279. doi:10.1037/met0000395
- Bürkner, P.-C. (2021). Bayesian Item Response Modeling in R with brms and Stan. *Journal of Statistical Software*, 100, 1–54. doi:10.18637/jss.v100.i05

See Also

`item_parameters` for the single-submodel version, `person_parameters_hpcm` for the person-side counterpart, `hurdle_acat` for the custom brms family.

item_restscore_post	Summarize and Plot Posterior Predictive Item-Restscore Associations
---------------------	---

Description

Postprocesses the output of `item_restscore_statistic` to produce a summary table and a slab plot comparing observed item-restscore gamma associations to the posterior predictive distribution.

Usage

```
item_restscore_post(item_restscore)
```

Arguments

- `item_restscore` A list as returned by `item_restscore_statistic`, containing at minimum:
- result** A data frame with columns including `item` and summary statistics (first 5 columns are used).
 - draws** A data frame with columns `item`, `gamma` (observed gamma per draw), and `gamma_rep` (replicated gamma per draw).

Details

The item-restscore gamma association measures the strength of the relationship between each item's responses and the rest score (total score excluding that item). Under good fit, the observed gamma should fall within the posterior predictive distribution.

The plot displays:

Grey slab The posterior predictive distribution of replicated gamma values, shaded by 84% interval levels.

Orange diamonds The observed gamma values per draw, plotted as points on top of the slab.

Items where the observed gamma (orange) falls consistently outside the replicated distribution (grey) indicate poor fit in terms of item discrimination.

Value

A list with two elements:

summary A `tibble` with the first 5 columns of the result table, rounded to 3 decimal places and sorted by item.

plot A `ggplot` object showing the posterior predictive distribution of replicated gamma (grey filled slab) with the observed gamma values overlaid as orange diamond points.

See Also

`item_restscore_statistic`, `infit_post`.

Examples

```
## Not run:
library(brms)
library(ggplot2)

# Assuming fit_pcm is a fitted brmsfit object
irs <- item_restscore_statistic(fit_pcm)

result <- item_restscore_post(irs)
result$summary
result$plot

## End(Not run)
```

item_restscore_statistic

Posterior Predictive Item-Restscore Association for Bayesian IRT Models

Description

Computes a Bayesian analogue of the item-restscore association test (Kreiner, 2011) for Rasch-family models fitted with **brms**. For each posterior draw, the Goodman-Kruskal gamma coefficient between each item's score and the rest-score (total score minus that item) is computed for both observed and replicated data. Posterior predictive p-values indicate whether the observed association is stronger than the model predicts, which signals violations of local independence or unidimensionality.

Usage

```
item_restscore_statistic(
  model,
  item_var = item,
  person_var = id,
  ndraws_use = NULL
)
```

Arguments

model	A fitted brmsfit object from an ordinal IRT model (e.g., family = acat for a partial credit model) or a dichotomous model (family = bernoulli()).
item_var	An unquoted variable name identifying the item grouping variable in the model data (e.g., item).
person_var	An unquoted variable name identifying the person grouping variable in the model data (e.g., id).
ndraws_use	Optional positive integer. If specified, a random subset of posterior draws of this size is used. If NULL (the default), all draws are used.

Details

The item-restscore association is a key diagnostic in Rasch measurement. Under the Rasch model, each item should relate to the latent trait (and hence the rest-score) only through the modelled relationship. Goodman-Kruskal's gamma is a rank-based measure of association for ordinal cross-tabulations that is well-suited for this purpose (Kreiner, 2011).

The procedure for each posterior draw s is:

1. Obtain replicated responses $Y^{rep(s)}$ from **posterior_predict**.
2. For each item i and each person v , compute the rest-score: $R_{vi}^{obs} = \sum_{j \neq i} X_{vj}$ for observed data and $R_{vi}^{rep(s)} = \sum_{j \neq i} Y_{vj}^{rep(s)}$ for replicated data.

3. Cross-tabulate item score $\times$ rest-score and compute the Goodman-Kruskal gamma for both observed and replicated data.
4. Compare the two gammas across draws.

Items with ppp close to 1 have observed item-restscore association that is consistently stronger than the model predicts. This typically indicates that the item discriminates more than assumed under the equal-discrimination Rasch model (i.e., a violation of the Rasch assumption). Items with ppp close to 0 discriminate less than expected.

Value

A [tibble](#) with the following columns:

item The item identifier.

gamma_obs Posterior mean of the observed Goodman-Kruskal gamma between this item and the rest-score.

gamma_rep Posterior mean of the replicated gamma.

gamma_diff Posterior mean of $\text{gamma_obs} - \text{gamma_rep}$. Positive values indicate the observed item-restscore association is stronger than the model expects.

ppp Posterior predictive p-value: $\text{mean}(\text{gamma_obs} > \text{gamma_rep})$ across draws. Values close to 1 indicate the item discriminates more than the model predicts (too high discrimination). Values close to 0 indicate the item discriminates less than expected (too low discrimination, e.g., noise or miskeyed item).

gamma_obs_q025, gamma_obs_q975 95\ the observed gamma.

gamma_obs_q005, gamma_obs_q995 99\ the observed gamma.

gamma_diff_q025, gamma_diff_q975 95\ the gamma difference.

gamma_diff_q005, gamma_diff_q995 99\ the gamma difference.

References

- Kreiner, S. (2011). A note on item-restscore association in Rasch models. *Applied Psychological Measurement*, 35(7), 557–561.
- Goodman, L. A. & Kruskal, W. H. (1954). Measures of association for cross classifications. *Journal of the American Statistical Association*, 49(268), 732–764.
- Bürkner, P.-C. (2021). Bayesian Item Response Modeling in R with brms and Stan. *Journal of Statistical Software*, 100, 1–54. [doi:10.18637/jss.v100.i05](https://doi.org/10.18637/jss.v100.i05)

See Also

[fit_statistic_pcm](#) for posterior predictive fit statistics, [fit_statistic_rm](#) for posterior predictive fit statistics, [infit_statistic](#) for Bayesian infit/outfit, [q3_statistic](#) for Bayesian Q3 residual correlations, [posterior_predict](#).

Examples

```

library(brms)
library(dplyr)
library(tidyr)
library(tibble)

# --- Partial Credit Model ---

df_pcm <- eRm::pcmdat2 %>%
  mutate(across(everything(), ~ .x + 1)) %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_pcm <- brm(
  response | thres(gr = item) ~ 1 + (1 | id),
  data = df_pcm,
  family = acat,
  chains = 4,
  cores = 1, # use more cores if you have
  iter = 500 # use at least 2000
)

# Item-restscore association
irs <- item_restscore_statistic(
  model = fit_pcm,
  ndraws_use = 100 # use at least 500
)

# Post-process draws
irs_results <- item_restscore_post(irs)
irs_results$summary
irs_results$plot

# --- Dichotomous Rasch Model ---

df_rm <- eRm::raschdat3 %>%
  as.data.frame() %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_rm <- brm(
  response ~ 1 + (1 | item) + (1 | id),
  data = df_rm,
  family = bernoulli(),
  chains = 4,
  cores = 1, # use more cores if you have
  iter = 500 # use at least 2000
)

irs_rm <- item_restscore_statistic(
  model = fit_rm,
  ndraws_use = 100 # use at least 500
)

```

```

)

# Post-process draws
irs_results <- item_restscore_post(irs_rm)
irs_results$summary
irs_results$plot

```

person_parameters	<i>Extract Person Parameters from a Bayesian Rasch Model</i>
-------------------	--

Description

Extracts person ability estimates from a fitted Bayesian Rasch model. Returns both Bayesian EAP (expected a posteriori) estimates with posterior SDs and frequentist WLE (Warm's weighted likelihood) estimates with standard errors, plus a lookup table mapping ordinal sum scores to both scales.

Usage

```

person_parameters(
  model,
  item_var = item,
  person_var = id,
  draws = FALSE,
  center = TRUE,
  theta_range = c(-7, 7)
)

```

Arguments

model	A fitted brmsfit object. Supported parameterisations: Polytomous ordinal (PCM) e.g., family = acat with thres(gr = item), producing item-specific thresholds. Dichotomous Rasch (random items) e.g., response ~ 1 + (1 item) + (1 id) with family = bernoulli(). Dichotomous 1PL (fixed items) e.g., response ~ 0 + item + (1 id) with family = bernoulli().
item_var	An unquoted variable name identifying the item grouping variable in the model data. Default is item.
person_var	An unquoted variable name identifying the person grouping variable in the model data. Default is id.
draws	Logical. If TRUE, a matrix of full posterior draws (persons x draws) is included in the output. Default is FALSE.

center	Logical. If TRUE (the default), person parameters and item difficulties are re-centered so that the mean item difficulty is zero, matching the convention in frequentist CML Rasch estimation. If FALSE, raw brms parameterisation is used.
theta_range	A numeric vector of length 2 specifying the range for the Newton-Raphson WLE search. Default is <code>c(-7, 7)</code> .

Details

EAP estimates are extracted as the posterior means of the person random effects (`r_id[j, Intercept]`) from `as_draws_df`, with the posterior SD serving as the standard error. These are the standard Bayesian point estimates and reflect shrinkage toward the population mean.

WLE estimates (Warm, 1989) are computed from the posterior mean item parameters using Newton-Raphson iteration with adaptive step damping on the Warm-corrected likelihood. WLE adds a bias correction term $J(\theta)/(2I(\theta))$ to the score equations, where $I(\theta)$ is the test information and $J(\theta) = \sum_i \sum_c (c - E_i)^3 P_{ic}$ is the sum of third central moments. This produces estimates with reduced finite-sample bias compared to MLE, especially at extreme scores (Warm, 1989).

The Newton-Raphson algorithm uses adaptive step damping following the approach in **iarm** (Mueller): the maximum allowed step size shrinks by a factor of 1.05 each iteration, preventing overshoot and ensuring convergence for near-extreme scores.

For extreme scores (sum score = 0 or maximum possible), the WLE is not well-defined. These cases are assigned the boundary values of `theta_range` with NA standard errors.

When `center = TRUE` (the default), item difficulty parameters are shifted so their mean is zero, and EAP person parameters are shifted by the same constant. WLE is computed from the centered item parameters. This matches the convention in frequentist CML Rasch estimation.

Row ordering: Both `person_estimates` and `draws_matrix` preserve the original person order from the model data (order of first appearance of each person ID). This allows direct row-binding with the source data without re-matching.

Value

A list with the following elements:

person_estimates A **tibble** with one row per person containing: the person ID, `sum_score`, `eap` (posterior mean of person random effect), `eap_se` (posterior SD), `wle` (Warm's weighted likelihood estimate), and `wle_se` (asymptotic SE of the WLE). Rows are ordered to match the original person order in the model data (i.e., the order of first appearance of each person ID).

score_table A **tibble** mapping each observed ordinal sum score to its mean EAP, mean EAP SE, WLE, and WLE SE. Extreme scores (0 and maximum) receive WLE estimates at the boundary of `theta_range` with NA standard errors.

draws_matrix (Only if `draws = TRUE`) A numeric matrix with rows = persons and columns = posterior draws. Row names are person IDs. Rows are ordered to match the original person order in the model data. Can be passed directly to `RMUreliability`.

References

Warm, T. A. (1989). Weighted likelihood estimation of ability in item response theory. *Psychometrika*, 54(3), 427–450. doi:10.1007/BF02294627

Bürkner, P.-C. (2021). Bayesian Item Response Modeling in R with brms and Stan. *Journal of Statistical Software*, 100, 1–54. doi:10.18637/jss.v100.i05

Christensen, K. B., Kreiner, S. & Mesbah, M. (Eds.) (2013). *Rasch Models in Health*. Iste and Wiley, pp. 63–70.

See Also

[RMUreliability](#), [plot_targeting](#), [ranef](#), [as_draws_df](#).

Examples

```
library(brms)
library(dplyr)
library(tidyr)
library(tibble)

# --- Dichotomous Rasch Model (random items) ---

df_rm <- eRm::raschdat3 %>%
  as.data.frame() %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_rm <- brm(
  response ~ 1 + (1 | item) + (1 | id),
  data = df_rm,
  family = bernoulli(),
  chains = 4, cores = 2, iter = 1000 # use more iter and cores
)

# Basic usage – rows match original person order
pp <- person_parameters(fit_rm)
pp$person_estimates
pp$score_table

# With full posterior draws (e.g., for RMUreliability)
pp_draws <- person_parameters(fit_rm, draws = TRUE)
RMUreliability(pp_draws$draws_matrix)

# --- Polytomous PCM (acat) ---

df_pcm <- eRm::pcmdat2 %>%
  mutate(across(everything(), ~ .x + 1)) %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_pcm <- brm(
  response | thres(gr = item) ~ 1 + (1 | id),
  data = df_pcm,
  family = acat,
  chains = 4, cores = 2, iter = 1000 # use more iter and cores
)
```

```
pp_pcm <- person_parameters(fit_pcm)
pp_pcm$score_table
```

```
person_parameters_hpcm
```

Extract Person Parameters from a Hurdle Partial Credit Model

Description

Extracts person trait estimates from a fitted hurdle Rasch partial credit model (family = `hurdle_acat()`), returning a per-submodel breakdown: a presence trait θ_{hurdle} (susceptibility) on the Bernoulli gate and a severity trait θ_{pcm} on the partial credit submodel. Each submodel's output mirrors the shape of `person_parameters` so `res$hurdle` and `res$pcm` can be passed to existing downstream functions (`RMUreliability`, `plot_targeting`, etc.).

Usage

```
person_parameters_hpcm(
  model,
  item_var = item,
  person_var = id,
  draws = FALSE,
  center = TRUE,
  theta_range = c(-7, 7)
)
```

Arguments

model	A fitted <code>brmsfit</code> object using the <code>hurdle_acat</code> custom family.
item_var	An unquoted variable name identifying the item grouping variable in the model data. Default is <code>item</code> .
person_var	An unquoted variable name identifying the person grouping variable in the model data. Default is <code>id</code> .
draws	Logical. If TRUE, a matrix of full posterior draws (persons x draws) is included in each submodel's output. Default is FALSE.
center	Logical. If TRUE (the default), item parameters and person parameters are shifted within each submodel so that the mean item parameter is zero. Person traits are shifted by the same constant as the items in their submodel.
theta_range	A numeric vector of length 2 giving the range for the Newton-Raphson WLE search on the hurdle submodel. Default is <code>c(-7, 7)</code> .

Details

Hurdle person trait. Defined as $\theta_{hurdle,v} = -r_{id}(v, \text{hu_Intercept})$, i.e., the brms random effect on hu with its sign flipped. Under this convention, the hurdle submodel reads $P(Y > 0) = \text{plogis}(\theta_{hurdle,v} - \delta_{hurdle,i})$, so higher θ_{hurdle} means greater presence / susceptibility. WLE is computed by treating the gate as a dichotomous Rasch model on $1[Y > 0]$ with item difficulties $\delta_{hurdle,i}$ from [item_parameters_hpcm](#).

Partial credit person trait. Defined as $\theta_{pcm,v} = r_{id}(v, \text{Intercept})$. Higher values mean greater severity given presence.

Why no WLE on the partial credit submodel. Conditional on the hurdle, the severity submodel is a PCM, but the set of items contributing to person v 's severity score depends on v 's own pattern of $1[Y_{vi} > 0]$: only items with $Y_{vi} > 0$ provide PCM information about $\theta_{pcm,v}$. The sum of partial credit scores is therefore not a sufficient statistic across the sample, and a standard score-table WLE is not well defined. The Bayesian EAP, which integrates over the (correlated) multivariate prior on $(\theta_{hurdle}, \theta_{pcm})$, remains well defined for all persons (including those with all-zero patterns) and is therefore the recommended point estimate; see Magnus and Garnier-Villarreal (2022, p. 272ff) for discussion in the closely-related MZI-GRM.

Centering. When `center = TRUE`, the same shifts that [item_parameters_hpcm](#) applies to the item parameters are applied here to the corresponding person traits. The likelihood is unchanged.

Row ordering. `person_estimates` and `draws_matrix` preserve the order of first appearance of each person ID in the model data, matching [person_parameters](#).

Value

A list with three elements:

hurdle A list with the same structure as [person_parameters](#) for a dichotomous Rasch model: `person_estimates` (one row per person; `sum_score` is the number of items with $Y > 0$; both EAP and Warm's WLE are provided), `score_table`, and optionally `draws_matrix`. The hurdle EAP is the sign-flipped brms random effect on hu, so higher values mean greater presence / susceptibility.

pcm A list with `person_estimates` (columns `id`, `sum_score`, `n_active`, `eap`, `eap_se`), `score_table`, and optionally `draws_matrix`. `sum_score` is the sum of Y across items with $Y_{vi} > 0$ for person v ; `n_active` is the count of such items. WLE columns are omitted intentionally (see Details).

correlation A [tibble](#) summarising the posterior of $\rho(\theta_{hurdle}, \theta_{pcm})$, identical to the correlation element returned by [item_parameters_hpcm](#).

References

- Magnus, B. E. & Garnier-Villarreal, M. (2022). A multidimensional zero-inflated graded response model for ordinal symptom data. *Psychological Methods*, 27(2), 261-279. doi:10.1037/met0000395
- Warm, T. A. (1989). Weighted likelihood estimation of ability in item response theory. *Psychometrika*, 54(3), 427-450. doi:10.1007/BF02294627

See Also

[personParameters](#) for the single-submodel version, [itemParameters_hpcm](#) for the item-side counterpart, [hurdle_acat](#) for the custom brms family.

plotBars

*Item Response Distribution Bar Chart***Description**

Creates a faceted bar chart showing the response distribution for each item, with counts and percentages displayed on each bar. Each item gets its own panel, with response categories on the x-axis and percentage of responses on the y-axis. This is a descriptive data visualization tool intended for use before model fitting.

Usage

```
plotBars(
  data,
  item_labels = NULL,
  category_labels = NULL,
  ncol = 1,
  label_wrap = 25,
  text_y = 6,
  viridis_option = "A",
  viridis_end = 0.9,
  font = "sans"
)
```

Arguments

<code>data</code>	A data frame in wide format containing only the item response columns. Each column is one item, each row is one person. All columns must be numeric (integer-valued). Response categories may be coded starting from 0 or 1. Do not include person IDs, grouping variables, or other non-item columns.
<code>item_labels</code>	An optional character vector of descriptive labels for the items (facet strips). Must be the same length as <code>ncol(data)</code> . If <code>NULL</code> (the default), column names are used. Labels are displayed as "column_name - label".
<code>category_labels</code>	An optional character vector of labels for the response categories (x-axis). Must be the same length as the number of response categories spanning from the minimum to the maximum observed value. If <code>NULL</code> (the default), numeric category values are used.
<code>ncol</code>	Integer. Number of columns in the faceted layout. Default is 1.
<code>label_wrap</code>	Integer. Number of characters per line in facet strip labels before wrapping. Default is 25.

text_y	Numeric. Vertical position (in percent units) for the count labels on each bar. Adjust upward if bars are tall. Default is 6.
viridis_option	Character. Viridis palette option for the count text color. One of "A" through "H". Default is "A".
viridis_end	Numeric in [0, 1]. End point of the viridis color scale for count text. Adjust if text is hard to read against the bar colors. Default is 0.9.
font	Character. Font family for all text. Default is "sans".

Details

Each item is displayed as a separate facet panel with the item label in the strip on the left side. Bars are colored by response category using the viridis palette. Each bar shows the count ($n = X$) as text.

Input requirements:

- All columns must be numeric (integer-valued).
- The data frame must contain at least 2 columns (items) and at least 1 row (person).

Value

A `ggplot` object.

Examples

```
library(ggplot2)
if (requireNamespace("eRm", quietly = TRUE))

# Basic response distribution plot
plot_bars(eRm::pcmdat2)

# With custom item labels
plot_bars(
  eRm::pcmdat2,
  item_labels = c("Mood", "Sleep", "Appetite", "Energy")
)

# Two-column layout with wrapped labels
plot_bars(
  eRm::pcmdat2,
  item_labels = c(
    "General mood and emotional wellbeing",
    "Quality of sleep at night",
    "Appetite and eating habits",
    "Overall energy level during the day"
  ),
  ncol = 2, label_wrap = 20
)

# With custom category labels
plot_bars(
  eRm::pcmdat2,
```

```
category_labels = c("Never", "Sometimes", "Often")
)
```

plot_icc

Item Characteristic Curves with Class Intervals

Description

Plots Item Characteristic Curves (ICCs) for Bayesian Rasch-family models fitted with **brms**. Each item panel shows the model-expected item score curve (with a credible interval ribbon) overlaid with observed average item scores computed within class intervals. Optionally, observed scores can be split by a grouping variable to visually assess differential item functioning (DIF).

Usage

```
plot_icc(
  model,
  item_var = item,
  person_var = id,
  items = NULL,
  n_intervals = 5,
  theta_range = c(-4, 4),
  n_points = 200,
  center = TRUE,
  prob = 0.95,
  ncol = NULL,
  line_size = 0.8,
  ribbon_alpha = 0.3,
  point_size = 2.5,
  dif_var = NULL,
  dif_data = NULL,
  dif_labels = NULL,
  dif_stats = TRUE,
  min_n = 5,
  palette = NULL
)
```

Arguments

model	A fitted <code>brmsfit</code> object from an ordinal IRT model (e.g., <code>family = acat</code>) or a dichotomous model (<code>family = bernoulli()</code>).
item_var	An unquoted variable name identifying the item grouping variable in the model data. Default is <code>item</code> .
person_var	An unquoted variable name identifying the person grouping variable in the model data. Default is <code>id</code> .

items	An optional character vector of item names to plot. If NULL (the default), all items are plotted.
n_intervals	Integer. The number of class intervals into which persons are binned along the sum score. Default is 5.
theta_range	A numeric vector of length 2 specifying the range of theta for the expected curve. Default is <code>c(-4, 4)</code> .
n_points	Integer. Number of evenly spaced theta values for computing the expected curve. Default is 200.
center	Logical. If TRUE (the default), the scale is recentered so the grand mean of item thresholds = 0, consistent with plot_targeting .
prob	Numeric in (0, 1). Width of the credible interval ribbon around the expected curve. Default is 0.95.
ncol	Integer. Number of columns in the faceted layout. If NULL, chosen automatically.
line_size	Numeric. Line width for the expected curve. Default is 0.8.
ribbon_alpha	Numeric in [0, 1]. Transparency of the credible interval ribbon. Default is 0.3.
point_size	Numeric. Size of observed score points. Default is 2.5.
dif_var	An optional unquoted variable name for a grouping variable to assess DIF visually. If supplied, observed scores are computed separately per group and coloured accordingly.
dif_data	An optional data frame containing the DIF variable. Required when <code>dif_var</code> is specified and the variable was not part of the model formula (since <code>brms</code> drops unused variables from <code>model\$data</code>). Must have the same rows and row order as the original model data.
dif_labels	An optional character vector of labels for the DIF groups. If NULL, factor levels are used.
dif_stats	Logical. If TRUE (the default when <code>dif_var</code> is specified), the partial gamma coefficient for each item is annotated in the plot panel. The partial gamma measures the strength of DIF, stratified by total score.
min_n	Integer. Minimum number of persons required in a class interval for the observed mean to be plotted. Intervals with fewer persons are dropped to avoid unstable estimates. Default is 5.
palette	An optional character vector of colors. If NULL, <code>viridis</code> colors are used.

Details

This is the Bayesian analogue of `ICCPlot()` from the **iarm** package, using the class interval method.

Expected curve: For each item, the expected item score $E_i(\theta)$ is computed for a dense grid of theta values using the posterior draws of item thresholds. For the adjacent category (PCM/acat) family:

$$E_i(\theta) = \sum_{c=0}^K c \cdot P_{ic}(\theta)$$

where P_{ic} are the category probabilities. For dichotomous items, $E_i(\theta) = P_i(\theta)$.

The posterior mean of $E_i(\theta)$ is plotted as a line, with a credible interval ribbon showing the prob interval across posterior draws.

Class intervals: Following the approach in `iarm::ICCplot()`, persons are binned into class intervals by their ordinal *sum score* (the sufficient statistic in Rasch models), not by their EAP theta estimate. Within each interval, the mean observed item response is computed and plotted at the theta value corresponding to the mean sum score in that interval, obtained by inverting the posterior-mean expected total score function. Persons with extreme sum scores (0 and maximum) are excluded from the class intervals, as their theta positions cannot be estimated.

Uncertainty around observed points: For each class interval, the standard error of the mean observed response is computed and displayed as error bars showing ± 1.96 SE. These reflect sampling variability of the observed mean within each bin.

DIF overlay: When `dif_var` is specified, observed scores are computed separately for each level of the DIF variable, producing group-specific points connected by lines. Deviations between groups that track alongside each other (but offset from the expected curve) suggest uniform DIF. Crossing group lines suggest non-uniform DIF.

Partial gamma: When `dif_stats = TRUE` (default when DIF is requested), the Goodman-Kruskal partial gamma coefficient is displayed in each item panel. This measures the ordinal association between item response and DIF group, stratified by total score. Values near 0 indicate no DIF; values near ± 1 indicate strong DIF. The computation reuses the concordant/discordant pair counting algorithm from [gk_gamma](#).

Value

A [ggplot](#) object.

See Also

[plot_ipf](#) for category probability curves, [plot_targeting](#) for person-item maps, [dif_statistic](#) for formal Bayesian DIF testing, [gk_gamma](#) for the underlying gamma algorithm.

Examples

```
library(brms)
library(dplyr)
library(tidyr)
library(tibble)
library(ggplot2)

# --- Partial Credit Model ---

df_pcm <- eRm::pcmdat2 %>%
  mutate(across(everything(), ~ .x + 1)) %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_pcm <- brm(
  response | thres(gr = item) ~ 1 + (1 | id),
  data = df_pcm,
  family = acat,
```

```

    chains = 4, cores = 2, iter = 1000 # use more iter and cores
  )

  # Basic ICC plot with 5 class intervals
  plot_icc(fit_pcm)

  # Select specific items, more intervals
  plot_icc(fit_pcm, items = c("I1", "I2"), n_intervals = 5)

  # With DIF overlay and partial gamma annotation

  # same dataset, adding a `gender` DIF variable
  df_pcm <- eRm::pcmdat2 %>%
    mutate(across(everything(), ~ .x + 1)) %>%
    rownames_to_column("id") %>%
    mutate(gender = sample(c("M", "F"), nrow(.), TRUE)) %>%
    pivot_longer(!c(id, gender), names_to = "item", values_to = "response")

  plot_icc(fit_pcm, dif_var = gender, dif_data = df_pcm,
    dif_labels = c("Female", "Male"), n_intervals = 5)

```

plot_icc_hpcm

Item Characteristic Curves with Class Intervals for Hurdle PCM

Description

Plots Item Characteristic Curves (ICCs) with class-interval overlays separately for each submodel of a hurdle partial credit model fitted with the [hurdle_acat](#) custom family. Returns two **ggplot2** plots — one for the Bernoulli hurdle on $P(Y > 0)$, one for the conditional partial credit $E(Y | Y > 0)$ — that can be inspected separately or combined with **patchwork**. Conceptually the Bayesian / hurdle-PCM analogue of [plot_icc](#), which only handles single-submodel ordinal / dichotomous IRT.

Usage

```

plot_icc_hpcm(
  model,
  item_var = item,
  person_var = id,
  items = NULL,
  n_intervals = 5,
  theta_range = c(-4, 4),
  n_points = 200,
  center = TRUE,
  prob = 0.95,
  ncol = NULL,
  line_size = 0.8,

```

```

    ribbon_alpha = 0.3,
    point_size = 2.5,
    min_n = 5
  )

```

Arguments

model	A fitted <code>brmsfit</code> object using the <code>hurdle_acat</code> custom family.
item_var	An unquoted variable name identifying the item grouping variable in the model data. Default is <code>item</code> .
person_var	An unquoted variable name identifying the person grouping variable in the model data. Default is <code>id</code> .
items	An optional character vector of item names to plot. If <code>NULL</code> (the default), all items are plotted.
n_intervals	Integer. The number of class intervals into which persons are binned within each submodel. Default is 5.
theta_range	A numeric vector of length 2 specifying the range of theta for the expected curves. Default is <code>c(-4, 4)</code> .
n_points	Integer. Number of evenly spaced theta values for computing the expected curves. Default is 200.
center	Logical. If <code>TRUE</code> (the default), each submodel is recentered so that the mean item parameter is zero, matching <code>item_parameters_hpcm</code> and <code>person_parameters_hpcm</code> .
prob	Numeric in $(0, 1)$. Width of the credible interval ribbon around each expected curve. Default is 0.95.
ncol	Integer. Number of columns in the faceted layout. If <code>NULL</code> , chosen automatically.
line_size	Numeric. Line width for the expected curves. Default is 0.8.
ribbon_alpha	Numeric in $[0, 1]$. Transparency of the credible interval ribbons. Default is 0.3.
point_size	Numeric. Size of observed score points. Default is 2.5.
min_n	Integer. Minimum number of observations required in a class interval (per item, per submodel) for the observed mean to be plotted. Default is 5.

Details

Hurdle submodel. For each item the expected curve is $P(Y_{vi} > 0 \mid \theta_{hurdle}) = \text{plogis}(\theta_{hurdle} - \delta_{hurdle,i})$, computed across posterior draws of the hurdle item difficulty. Persons are binned into class intervals by their hurdle sum score (count of items with $Y > 0$); each interval is positioned on the θ_{hurdle} axis by inverting the posterior-mean expected total $\sum_i P(Y_i > 0 \mid \theta_{hurdle})$, exactly analogous to the procedure in `plot_icc`. The y-axis ranges from 0 to 1.

Partial credit submodel. For each item the expected curve is the conditional expected severity $E(Y_{vi} \mid Y_{vi} > 0, \theta_{pcm}) = \sum_{c=1}^{K-1} c \cdot P(Y_{vi} = c \mid Y_{vi} > 0)$, with conditional PCM category probabilities computed from the posterior draws of τ_{ik} . The y-axis ranges from 1 to $K - 1$.

Persons are binned into class intervals by their **EAP** θ_{pcm} (taken from `ranef`), not by a sum score. This is intentional: under the hurdle PCM the sum of Y over items with $Y_{vi} > 0$ is not a sufficient statistic for θ_{pcm} (because the number of contributing items varies across persons;

see [person_parameters_hpcm](#)). EAP-based binning sidesteps the inverse-CDF construction and avoids privileging any particular summary score. Within each interval, the observed mean response for an item is computed *only* over persons with $Y > 0$ on that item — i.e., the cells the PCM submodel actually applies to.

Uncertainty. Each expected curve carries a credible interval ribbon at width prob. Observed points have $\pm 1.96 SE$ error bars where SE is the within-bin sampling standard error of the mean (binomial for the hurdle, ordinal for the partial credit submodel).

Why two separate plots and not patchwork-combined. The two submodels have qualitatively different y-axes (probability vs. expected severity) and different x-axis interpretations (presence trait vs. severity trait). Returning them as a list preserves the option to inspect each in isolation; combine them yourself with **patchwork** when a side-by-side or stacked layout is wanted:

```
plots <- plot_icc_hpcm(fit)
patchwork::wrap_plots(plots$hurdle, plots$pcm, ncol = 1)
```

Value

A list with two elements:

hurdle A [ggplot](#) object showing per-item Bernoulli ICCs on the $P(Y > 0)$ scale, with class-interval observed empirical hurdle-crossing rates overlaid.

pcm A [ggplot](#) object showing per-item conditional partial credit ICCs on the $E(Y | Y > 0)$ scale, with class-interval observed mean severities overlaid (restricted to persons with $Y > 0$ on that item).

References

Magnus, B. E. & Garnier-Villarreal, M. (2022). A multidimensional zero-inflated graded response model for ordinal symptom data. *Psychological Methods*, 27(2), 261-279. doi:10.1037/met0000395

See Also

[plot_icc](#) for the single-submodel ICC plot, [item_parameters_hpcm](#), [person_parameters_hpcm](#).

plot_ipf

Item Category Probability Function Curves for Polytomous IRT Models

Description

Plots item category probability functions (ICPFs) for polytomous Bayesian IRT models fitted with **brms**. For each item, the probability of endorsing each response category is plotted as a function of the latent variable (theta), with separate colored curves per category. All items are displayed in a combined faceted plot.

Usage

```
plot_ipf(
  model,
  item_var = item,
  person_var = id,
  items = NULL,
  theta_range = c(-4, 4),
  n_points = 100,
  ncol = NULL,
  line_size = 0.8,
  ribbon_alpha = 0.15,
  prob = 0.95,
  category_labels = NULL,
  palette = NULL
)
```

Arguments

model	A fitted brmsfit object from a polytomous IRT model (e.g., family = acat for a partial credit model or family = cumulative for a graded response model).
item_var	An unquoted variable name identifying the item grouping variable in the model data (e.g., item).
person_var	An unquoted variable name identifying the person grouping variable in the model data (e.g., id).
items	An optional character vector of item names to plot. If NULL (the default), all items in the model are plotted.
theta_range	A numeric vector of length 2 specifying the range of the latent variable (theta) for the x-axis. Default is c(-4, 4).
n_points	Integer. Number of evenly spaced theta values at which to evaluate the category probabilities. Default is 100.
ncol	Integer. Number of columns in the faceted plot layout. If NULL (the default), an appropriate number is chosen automatically.
line_size	Numeric. Line width for the probability curves. Default is 0.8.
ribbon_alpha	Numeric in [0, 1]. Transparency of the credible interval ribbons. Default is 0.15. Set to 0 to hide ribbons.
prob	Numeric in (0, 1). Width of the credible interval for the ribbons. Default is 0.95.
category_labels	An optional character vector of labels for the response categories. If NULL (the default), categories are labelled as integers starting from 1.
palette	An optional character vector of colors, one per response category. If NULL (the default), the viridis discrete scale from ggplot2 is used.

Details

The function computes category probabilities directly from the posterior draws of the item threshold parameters. For the brms `acat` (adjacent category / partial credit) family with logit link, the density is:

$$P(Y = y|\eta) = \frac{\exp(\sum_{k=1}^y(\eta - \tau_k))}{\sum_{k=0}^K \exp(\sum_{j=1}^k(\eta - \tau_j))}$$

where η is the linear predictor (i.e., theta for a Rasch model with no additional fixed effects) and τ_k are the item thresholds. Analogous formulas are used for the `cumulative`, `sratio`, and `cratio` families.

Posterior uncertainty in the thresholds propagates into credible interval ribbons around the category probability curves — a Bayesian advantage over point-estimate-based plots.

Value

A `ggplot` object. The plot can be further customised using standard **ggplot2** functions.

References

Bürkner, P.-C. (2021). Bayesian Item Response Modeling in R with brms and Stan. *Journal of Statistical Software*, 100, 1–54. doi:10.18637/jss.v100.i05

See Also

`posterior_epred`, `conditional_effects`,

Examples

```
library(brms)
library(dplyr)
library(tidyr)
library(tibble)
library(ggplot2)

# --- Partial Credit Model ---

df_pcm <- eRm::pcmdat2 %>%
  mutate(across(everything(), ~ .x + 1)) %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_pcm <- brm(
  response | thres(gr = item) ~ 1 + (1 | id),
  data = df_pcm,
  family = acat,
  chains = 4,
  cores = 1, # use more cores if you have
  iter = 500 # use at least 2000
)

# Plot all items
```

```

plot_ipf(fit_pcm, item_var = item, person_var = id)

# Plot a subset of items
plot_ipf(fit_pcm, item_var = item, person_var = id,
         items = c("I1", "I2", "I3"))

# Customise appearance
plot_ipf(fit_pcm, item_var = item, person_var = id,
         theta_range = c(-6, 6), ncol = 3, prob = 0.90) +
  theme_minimal() +
  labs(title = "Item Category Probability Functions")

```

plot_residual_pca

Residual PCA Contrast Plot for Bayesian IRT Models

Description

Assesses dimensionality of Bayesian IRT models by performing a principal component analysis (PCA) of standardized residuals for each posterior draw. The item loadings on the first residual contrast are plotted against item locations, with posterior uncertainty displayed as 2D density contours, crosshairs, or both. A posterior predictive p-value for the first-contrast eigenvalue tests whether the observed residual structure exceeds what the model predicts under unidimensionality.

Usage

```

plot_residual_pca(
  model,
  item_var = item,
  person_var = id,
  center = TRUE,
  prob = 0.95,
  ndraws_use = NULL,
  style = c("both", "density", "crosshair"),
  density_alpha = 0.3,
  density_bins = 6,
  density_palette = NULL,
  label_items = TRUE,
  point_size = 2.5,
  point_color = "#0072B2"
)

```

Arguments

model	A fitted brmsfit object from an ordinal IRT model (e.g., family = acat) or a dichotomous model (family = bernoulli()).
-------	--

item_var	An unquoted variable name identifying the item grouping variable in the model data. Default is item.
person_var	An unquoted variable name identifying the person grouping variable in the model data. Default is id.
center	Logical. If TRUE (the default), item locations are mean-centered to zero, matching the convention used in plot_targeting .
prob	Numeric in (0, 1). Width of the credible intervals for both loading and location whiskers. Default is 0.95.
ndraws_use	Optional positive integer. Number of posterior draws to use. If NULL (the default), up to 500 draws are used.
style	Character. Visual style for displaying uncertainty. "density" (the default) overlays filled 2D density contours per item computed from the draw-level location and loading values, showing the full joint posterior uncertainty. "crosshair" shows point estimates with horizontal and vertical credible interval bars. "both" displays density contours with crosshairs on top.
density_alpha	Numeric in [0, 1]. Maximum opacity of the density contours when style is "density" or "both". Default is 0.3.
density_bins	Integer. Number of contour bins for geom_density_2d_filled . Default is 6.
density_palette	An optional character vector of colors for the density contour fills (from low to high density). If NULL (the default), a blue sequential ramp is used. The length should match density_bins; the lowest (background) level is always transparent.
label_items	Logical. If TRUE (the default), item names are displayed next to points.
point_size	Numeric. Size of the item points. Default is 2.5.
point_color	Color for the item points and error bars. Default is "#0072B2".

Details

The procedure for each posterior draw s is:

1. Obtain category probabilities from [posterior_epred](#). Compute expected values $E_{vi}^{(s)}$ and variances $Var_{vi}^{(s)}$.
2. Compute standardized residuals for observed data:

$$z_{vi}^{obs(s)} = \frac{X_{vi} - E_{vi}^{(s)}}{\sqrt{Var_{vi}^{(s)}}}$$

3. Generate replicated data $Y^{rep(s)}$ from [posterior_predict](#) and compute standardized residuals:

$$z_{vi}^{rep(s)} = \frac{Y_{vi}^{rep(s)} - E_{vi}^{(s)}}{\sqrt{Var_{vi}^{(s)}}}$$

4. Reshape both sets of residuals into person $\times$ item matrices and perform SVD on each.

5. Extract the first-contrast eigenvalue and item loadings from both observed and replicated SVDs.
6. Compare eigenvalues across draws: the posterior predictive p-value $\text{ppp} = \text{mean}(\text{eigenvalue_obs} > \text{eigenvalue_rep})$ tests whether the observed residual structure is stronger than what the model produces under its own assumptions.

When `style = "density"` or `style = "both"`, the draw-level (location, loading) pairs for each item are used to construct filled 2D kernel density contours via `geom_density_2d_filled`. The lowest contour level (outside all contours) is set to transparent so the white panel background shows through. Higher density regions use progressively darker fills.

Item loadings are aligned across draws using majority-sign alignment to resolve the sign indeterminacy of eigenvectors.

Value

A list with three elements:

plot A `ggplot` object showing item loadings on the first residual contrast (y-axis) versus item locations (x-axis).

data A `tibble` with columns: `item`, `location` (posterior mean item location), `location_lower`, `location_upper` (location CI), `loading` (posterior mean loading on first contrast), `loading_lower`, `loading_upper` (loading CI).

eigenvalue A `tibble` with columns: `eigenvalue_obs` (posterior mean observed eigenvalue), `eigenvalue_rep` (posterior mean replicated eigenvalue), `eigenvalue_diff` (posterior mean difference), `ppp` (posterior predictive p-value), `var_explained_obs`, `var_explained_rep` (posterior mean proportions of residual variance explained).

References

Smith, E. V. (2002). Detecting and evaluating the impact of multidimensionality using item fit statistics and principal component analysis of residuals. *Journal of Applied Measurement*, 3, 205–231.

Bürkner, P.-C. (2021). Bayesian Item Response Modeling in R with brms and Stan. *Journal of Statistical Software*, 100, 1–54. doi:10.18637/jss.v100.i05

See Also

`plot_targeting` for person-item maps, `plot_ipf` for item category probability curves, `q3_statistic` for Q3 residual correlations (another local dependence / dimensionality diagnostic).

Examples

```
## Not run:
library(brms)
library(dplyr)
library(tidyr)
library(tibble)
library(ggplot2)
```

```

# --- Partial Credit Model ---

df_pcm <- eRm::pcmdat2 %>%
  mutate(across(everything(), ~ .x + 1)) %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_pcm <- brm(
  response | thres(gr = item) ~ 1 + (1 | id),
  data = df_pcm,
  family = acat,
  chains = 4,
  cores = 4,
  iter = 2000
)

# 2D density contours (default)
result <- plot_residual_pca(fit_pcm)
result$plot

# Crosshair style
result_c <- plot_residual_pca(fit_pcm, style = "crosshair")
result_c$plot

# Both combined
result_b <- plot_residual_pca(fit_pcm, style = "both")
result_b$plot

# Custom warm palette
result_w <- plot_residual_pca(
  fit_pcm,
  density_palette = c("#FEE8C8", "#FDBB84", "#E34A33",
    "#B30000", "#7F0000", "#4A0000"),
  point_color = "#B30000"
)
result_w$plot

## End(Not run)

```

plot_stackedbars

Stacked Bar Chart of Item Response Distributions

Description

Creates a horizontal stacked bar chart showing the response distribution for all items. Each bar represents one item, with segments colored by response category. Counts are displayed as text labels within each segment. This is a descriptive data visualization tool intended for use before model fitting.

Usage

```
plot_stackedbars(
  data,
  item_labels = NULL,
  category_labels = NULL,
  show_n = TRUE,
  show_percent = FALSE,
  text_color = "sienna1",
  text_size = 3,
  min_label_n = 0,
  viridis_option = "D",
  viridis_end = 0.99,
  title = "Item responses"
)
```

Arguments

<code>data</code>	A data frame in wide format containing only the item response columns. Each column is one item, each row is one person. All columns must be numeric (integer-valued). Response categories may be coded starting from 0 or 1. Do not include person IDs, grouping variables, or other non-item columns.
<code>item_labels</code>	An optional character vector of descriptive labels for the items (y-axis). Must be the same length as <code>ncol(data)</code> . If <code>NULL</code> (the default), column names are used.
<code>category_labels</code>	An optional character vector of labels for the response categories (legend). Must be the same length as the number of response categories spanning from the minimum to the maximum observed value, ordered from lowest to highest category. If <code>NULL</code> (the default), numeric category values are used.
<code>show_n</code>	Logical. If <code>TRUE</code> (the default), the count of responses is displayed as a text label inside each bar segment.
<code>show_percent</code>	Logical. If <code>TRUE</code> , the percentage of responses is displayed instead of (or in addition to) counts. Default is <code>FALSE</code> .
<code>text_color</code>	Character. Color for the count/percentage labels. Default is "sienna1".
<code>text_size</code>	Numeric. Size of the count/percentage labels. Default is 3.
<code>min_label_n</code>	Integer. Minimum count required for a label to be displayed within a bar segment. Segments with fewer responses are left unlabelled to avoid clutter. Default is 0 (all segments labelled).
<code>viridis_option</code>	Character. Viridis palette option. One of "A" through "H". Default is "D".
<code>viridis_end</code>	Numeric in [0, 1]. End point of the viridis color scale. Default is 0.99.
<code>title</code>	Character. Plot title. Default is "Item responses".

Details

Items are displayed on the y-axis in the same order as the columns in `data` (first column at the top). Each bar is divided into segments representing response categories, with the lowest category on the

left and the highest on the right. The total bar length equals the number of non-missing responses for that item.

Categories with zero responses still appear in the legend but produce no visible bar segment, which helps identify gaps in the response distribution.

Input requirements:

- All columns must be numeric (integer-valued).
- The data frame must contain at least 2 columns (items) and at least 1 row (person).

Value

A `ggplot` object.

Examples

```
library(ggplot2)
if (requireNamespace("eRm", quietly = TRUE))

# Basic stacked bar chart
plot_stackedbars(eRm::pcmdat2)

# With custom item and category labels
plot_stackedbars(
  eRm::pcmdat2,
  item_labels = c("Mood", "Sleep", "Appetite", "Energy"),
  category_labels = c("Never", "Sometimes", "Often")
)

# Show percentages, suppress small segments
plot_stackedbars(
  eRm::pcmdat2,
  show_percent = TRUE,
  show_n = FALSE,
  min_label_n = 5
)
```

plot_targeting

Person-Item Map (Targeting Plot) for Bayesian IRT Models

Description

Plots a person-item map (also known as a Wright map or targeting plot) for Bayesian IRT models fitted with **brms**. The plot consists of three vertically stacked panels sharing the same latent variable (theta / logit) x-axis:

Usage

```
plot_targeting(
  model,
  item_var = item,
  person_var = id,
  robust = FALSE,
  center = TRUE,
  sort_items = c("data", "location"),
  bins = 30,
  prob = 0.95,
  palette = NULL,
  person_fill = "#0072B2",
  threshold_fill = "#D55E00",
  height_ratios = c(3, 2, 5)
)
```

Arguments

model	A fitted <code>brmsfit</code> object from an ordinal IRT model (e.g., family = <code>acat</code>) or a dichotomous model (family = <code>bernoulli()</code>).
item_var	An unquoted variable name identifying the item grouping variable in the model data. Default is <code>item</code> .
person_var	An unquoted variable name identifying the person grouping variable in the model data. Default is <code>id</code> .
robust	Logical. If <code>FALSE</code> (the default), the histogram annotations use mean $\pm$ SD. If <code>TRUE</code> , median $\pm$ MAD is used instead.
center	Logical. If <code>TRUE</code> (the default), the scale is recentered so that the grand mean of all item threshold locations is zero, following the convention in frequentist Rasch analysis. Person estimates are shifted by the same constant. If <code>FALSE</code> , the raw <code>brms</code> parameterisation is used.
sort_items	Character. How to order items on the y-axis of the bottom panel. <code>"data"</code> (the default) preserves the order in which items first appear in the model data, with the first item at the top. <code>"location"</code> sorts items by their mean threshold location (easiest at top, hardest at bottom).
bins	Integer. Number of bins for both histograms. Default is 30.
prob	Numeric in (0, 1). Width of the credible intervals for the item threshold whiskers. Default is 0.95.
palette	An optional character vector of colors for the response categories. If <code>NULL</code> (the default), the <code>viridis</code> discrete scale is used.
person_fill	Fill color for the person histogram. Default is <code>"#0072B2"</code> (blue).
threshold_fill	Fill color for the threshold histogram. Default is <code>"#D55E00"</code> (vermillion).
height_ratios	Numeric vector of length 3 specifying the relative heights of the top (person), middle (threshold), and bottom (dot-whisker) panels. Default is <code>c(3, 2, 5)</code> .

Details

1. **Top:** A histogram of person ability estimates, with a reference line for the mean (or median) and shading for ± 1 SD (or ± 1 MAD).
2. **Middle:** An inverted histogram of item threshold locations, with a reference line for the mean (or median) and shading for ± 1 SD (or ± 1 MAD), mirroring the top panel to visualise the overlap between person abilities and item difficulties.
3. **Bottom:** A dot-and-whisker plot of item thresholds by item, with credible intervals and color-coded response categories.

Together, the top and middle panels form a half-moon (or back-to-back histogram) display that makes it easy to assess whether the test is well-targeted to the sample.

Person estimates are obtained as the posterior means of the person random effects from the fitted model via [ranef](#).

Item thresholds are extracted from the posterior draws. For models with grouped thresholds (`thres(gr = item)`), each item has its own set of threshold parameters. For models with a single set of thresholds (e.g., dichotomous Rasch with `(1 | item)`), the item random effects are subtracted from the global thresholds to obtain item-specific locations.

When `center = TRUE` (the default), the grand mean of all item threshold posterior means is computed and subtracted from every threshold estimate, its credible interval bounds, and every person estimate. This is a uniform translation of the entire scale that preserves all relative distances and matches the zero-centered item difficulty convention used in frequentist CML estimation.

Value

A patchwork object (combined `ggplot`).

References

- Wright, B. D. & Stone, M. H. (1979). *Best Test Design*. MESA Press.
- Bürkner, P.-C. (2021). Bayesian Item Response Modeling in R with `brms` and `Stan`. *Journal of Statistical Software*, 100, 1–54. doi:10.18637/jss.v100.i05

See Also

[plot_ipf](#) for item category probability curves, [ranef](#), [as_draws_df](#).

Examples

```
library(brms)
library(dplyr)
library(tidyr)
library(tibble)
library(ggplot2)
library(patchwork)

# --- Partial Credit Model ---

df_pcm <- eRm::pcmdat2 %>%
  mutate(across(everything(), ~ .x + 1)) %>%
```

```

rownames_to_column("id") %>%
pivot_longer(!id, names_to = "item", values_to = "response")

fit_pcm <- brm(
  response | thres(gr = item) ~ 1 + (1 | id),
  data = df_pcm,
  family = acat,
  chains = 4,
  cores = 1, # use more cores if you have
  iter = 500 # use at least 2000
)

# Default: centered, mean ± SD, items in data order
plot_targeting(fit_pcm)

# Uncentered (raw brms parameterisation)
plot_targeting(fit_pcm, center = FALSE)

# Robust: median ± MAD, items sorted by location
plot_targeting(fit_pcm, robust = TRUE, sort_items = "location")

# --- Dichotomous Rasch Model ---

df_rm <- eRm::raschdat3 %>%
  as.data.frame() %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_rm <- brm(
  response ~ 1 + (1 | item) + (1 | id),
  data = df_rm,
  family = bernoulli(),
  chains = 4,
  cores = 1, # use more cores if you have
  iter = 500 # use at least 2000
)

plot_targeting(fit_rm, sort_items = "location")

```

plot_targeting_hpcm *Person-Item Targeting Plot for the Hurdle Partial Credit Model*

Description

Builds two person-item targeting plots — one for each submodel of a hurdle partial credit model fitted with the [hurdle_acat](#) custom family. Each plot is a three-panel **patchwork** stack with the same anatomy as [plot_targeting](#): a person histogram, an inverted item-location histogram, and a dot-and-whisker by item. Returning two plots — one for the presence trait ($\theta_{hurdle}, \delta_{hurdle}$) and

one for the severity trait (θ_{pcm}, τ_{ik}) — reflects the fact that the two submodels live on distinct latent scales and should be inspected on their own axes.

Usage

```
plot_targeting_hpcm(
  model,
  item_var = item,
  person_var = id,
  robust = FALSE,
  center = TRUE,
  sort_items = c("data", "location"),
  bins = 30,
  prob = 0.95,
  palette = NULL,
  person_fill = "#0072B2",
  threshold_fill = "#D55E00",
  height_ratios = c(3, 2, 5)
)
```

Arguments

model	A fitted <code>brmsfit</code> object using the <code>hurdle_acat</code> custom family.
item_var	An unquoted variable name identifying the item grouping variable in the model data. Default is <code>item</code> .
person_var	An unquoted variable name identifying the person grouping variable in the model data. Default is <code>id</code> .
robust	Logical. If TRUE, the central tendency / spread markers in the histogram panels use median $\pm$ MAD instead of mean $\pm$ SD. Default is FALSE.
center	Logical. If TRUE (the default), each submodel is recentered so the mean item parameter is zero, with the corresponding person trait shifted by the same constant — matching <code>item_parameters_hpcm</code> and <code>person_parameters_hpcm</code> .
sort_items	One of "data" (default) or "location", controlling the ordering of items in the dot-and-whisker panel.
bins	Integer. Number of histogram bins. Default is 30.
prob	Numeric in (0, 1). Width of the credible interval shown as horizontal whiskers in the dot-and-whisker panel. Default is 0.95.
palette	An optional character vector of colours for the threshold-category dot-and-whisker scale. If NULL, <code>viridis</code> is used. Applied to both submodels.
person_fill	Fill colour for the person histograms. Default "#0072B2".
threshold_fill	Fill colour for the threshold histograms. Default "#D55E00".
height_ratios	A numeric vector of length 3 giving the relative heights of the (person, threshold, dot-and-whisker) panels. Default <code>c(3, 2, 5)</code> .

Details

Hurdle scale. The presence person trait is taken as $\theta_{hurdle,v} = -r_{id}(v, \text{hu_Intercept})$ (the brms random effect on hu with its sign flipped, so higher values mean greater presence). Hurdle item difficulties are $\delta_{hurdle,i} = b_{hu_factoritem,i}$ directly (higher values = more zeros = harder hurdle to cross). Under this convention, $P(Y_{vi} > 0) = \text{plogis}(\theta_{hurdle,v} - \delta_{hurdle,i})$, and the histograms are directly comparable on a single x-axis.

Partial credit scale. The severity person trait is $\theta_{pcm,v} = r_{id}(v, \text{Intercept})$, and per-item thresholds are $\tau_{ik} = b_{\text{Intercept}[i,k]}$. The middle histogram aggregates thresholds across items and threshold indices, exactly as in [plot_targeting](#).

Independent centering per submodel. When `center = TRUE`, the hurdle is shifted by $\overline{\delta_{hurdle}}$ and the PCM by $\bar{\tau}$ (a different constant per submodel). The two resulting x-axes therefore have a shared origin interpretation (mean item parameter = 0) but cannot be combined on a single axis — these are different latent traits.

Combining the two plots. Each list element is a valid **patchwork** object, so they can be combined with the usual operators:

```
plots <- plot_targeting_hpcm(fit)
patchwork::wrap_plots(plots$hurdle, plots$pcm, ncol = 2)
```

For a tall, single-column layout, prefer `wrap_plots(..., ncol = 1)` so each submodel keeps its own three-panel column.

Value

A list with two elements:

hurdle A **patchwork** object stacking the θ_{hurdle} histogram, the δ_{hurdle} histogram (inverted), and the per-item hurdle difficulty dot-and-whisker with the credible interval given by `prob`.

pcm A **patchwork** object with the same anatomy for the partial credit submodel: θ_{pcm} histogram, PCM threshold histogram (inverted), and a per-item dot-and-whisker with one row per item and one coloured marker per threshold within item.

References

Wright, B. D. & Stone, M. H. (1979). *Best Test Design*. MESA Press.

Magnus, B. E. & Garnier-Villareal, M. (2022). A multidimensional zero-inflated graded response model for ordinal symptom data. *Psychological Methods*, 27(2), 261-279. doi:10.1037/met0000395

See Also

[plot_targeting](#) for the single-submodel version, [item_parameters_hpcm](#), [person_parameters_hpcm](#).

plot_tile

*Tile Plot of Item Response Distributions***Description**

Creates a tile (heat map) plot showing the distribution of responses across all items and response categories. Each cell displays the count (or percentage) of responses, with optional conditional highlighting for cells with low counts. This is a descriptive data visualization tool intended for use before model fitting.

Usage

```
plot_tile(
  data,
  cutoff = 10,
  highlight = TRUE,
  percent = FALSE,
  text_color = "orange",
  item_labels = NULL,
  category_labels = NULL
)
```

Arguments

<code>data</code>	A data frame in wide format containing only the item response columns. Each column is one item, each row is one person. All columns must be numeric (integer-valued). Response categories may be coded starting from 0 or 1. Do not include person IDs, grouping variables, or other non-item columns.
<code>cutoff</code>	Integer. Cells with counts below this value are highlighted (when <code>highlight = TRUE</code>). Default is 10.
<code>highlight</code>	Logical. If <code>TRUE</code> (the default), cell labels with counts below <code>cutoff</code> are displayed in red. This includes cells with zero responses (empty categories), which is useful for identifying gaps in the response distribution.
<code>percent</code>	Logical. If <code>TRUE</code> , cell labels show percentages instead of raw counts. Default is <code>FALSE</code> .
<code>text_color</code>	Character. Color for cell label text (when not highlighted). Default is "orange".
<code>item_labels</code>	An optional character vector of descriptive labels for the items (y-axis). Must be the same length as <code>ncol(data)</code> . If <code>NULL</code> (the default), column names are used.
<code>category_labels</code>	An optional character vector of labels for the response categories (x-axis). Must be the same length as the number of response categories spanning from the minimum to the maximum observed value. If <code>NULL</code> (the default), numeric category values are used.

Details

The plot displays items on the y-axis (in the same order as the columns in data, from top to bottom) and response categories on the x-axis. Cell shading represents the count of responses (darker = more responses). Cell labels show either raw counts or percentages.

Categories with zero responses are explicitly shown (as cells with $n = 0$), which helps identify gaps in the response distribution — one of the primary purposes of this plot.

Input requirements:

- All columns must be numeric (integer-valued).
- The data frame must contain at least 2 columns (items) and at least 1 row (person).

Value

A `ggplot` object.

Examples

```
library(ggplot2)
if (requireNamespace("eRm", quietly = TRUE))

# Basic tile plot
plot_tile(eRm::pcmdat2)

# With custom item labels
plot_tile(
  eRm::pcmdat2,
  item_labels = c("Mood", "Sleep", "Appetite", "Energy")
)

# With custom category labels and percentages
plot_tile(
  eRm::pcmdat2,
  category_labels = c("Never", "Sometimes", "Often"),
  percent = TRUE
)

# Adjust cutoff for highlighting
plot_tile(eRm::pcmdat2, cutoff = 20, highlight = TRUE)
```

posterior_to_prior

Extract Informative Priors from a Fitted Bayesian IRT Model

Description

Takes a fitted `brmsfit` object and constructs a `brmsprior` object in which each item parameter receives a normal(mean, sd) prior derived from its posterior distribution. The person-level random effect SD prior is also updated. The returned prior can be passed directly to `update` (or `brm`) to refit the model with empirical Bayes / informative priors — useful for anchoring scales, warm-starting a model on new data, or regularising estimation with small samples.

Usage

```
posterior_to_prior(
  model,
  item_var = item,
  person_var = id,
  mult = 1,
  target_link = c("source", "logit", "probit")
)
```

Arguments

model	A fitted brmsfit object. Supported parameterisations: Polytomous ordinal e.g., family = acat with thres(gr = item), producing item-specific thresholds. Dichotomous Rasch (random items) e.g., response ~ 1 + (1 item) + (1 id) with family = bernoulli(). Dichotomous 1PL (fixed items) e.g., response ~ 0 + item + (1 id) with family = bernoulli().
item_var	An unquoted variable name identifying the item grouping variable in the model data. Default is item.
person_var	An unquoted variable name identifying the person grouping variable in the model data. Default is id.
mult	Numeric multiplier applied to each posterior SD before it is used as the prior SD. Values > 1 widen the priors (less informative); values < 1 tighten them. Default is 1 (use posterior SD directly).
target_link	Character string specifying the link function of the model the priors will be used with. One of "logit", "probit", or "source" (the default). When "source", the link function of the fitted model is used and no transformation is applied. When different from the source model's link, all location and scale parameters are rescaled using the approximation $\beta_{\text{probit}} \approx \beta_{\text{logit}}/1.7$. This is useful when transferring priors from a logit-fitted model to a probit model or vice versa.

Details

The function extracts all posterior draws via [as_draws_df](#), computes the mean and SD of each parameter's marginal posterior, and constructs normal(mean, sd * mult) priors.

Polytomous ordinal models with grouped thresholds (thres(gr = item)): each threshold receives its own prior via `brms::set_prior("normal(...)", class = "Intercept", group = item, coef = threshold_index)`.

Dichotomous Rasch models parameterised as response ~ 1 + (1 | item) + (1 | id): priors are set on the global intercept (class = "Intercept"), the item-level SD (class = "sd", group = item_var), and the person-level SD.

Dichotomous 1PL models parameterised as response ~ 0 + item + (1 | id): each item-specific fixed effect (e.g., b_itemI1) receives its own normal(mean, sd) prior via `brms::set_prior(..., class = "b", coef = "itemI1")`.

In all cases the person-level SD receives a normal(mean, sd * mult) prior (brms applies the lower bound of zero automatically for SD parameters).

Link function transformation: When target_link differs from the source model's link function, all parameters (means and SDs) are rescaled by a factor of approximately 1.7. This uses the well-known approximation that $\Phi(x) \approx \text{logistic}(1.7x)$, so logit-scale parameters can be converted to probit-scale by dividing by 1.7, and vice versa. The approximation is excellent for parameters in the range $|\beta| < 3$ and adequate beyond that range.

Value

A `brmsprior` object that can be supplied to the prior argument of `brm` or `update`.

Examples

```
library(brms)
library(dplyr)
library(tidyr)
library(tibble)

# --- Partial Credit Model ---

df_pcm <- eRm::pcmdat2 %>%
  mutate(across(everything(), ~ .x + 1)) %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_pcm <- brm(
  response | thres(gr = item) ~ 1 + (1 | id),
  data = df_pcm,
  family = acat,
  chains = 4, cores = 2, iter = 1000 # use more iter (and cores if you have)
)

# Extract posterior-informed priors (same link)
new_priors <- posterior_to_prior(fit_pcm)
new_priors

# Narrow the prior's sd by a factor of 0.5
wide_priors <- posterior_to_prior(fit_pcm, mult = 0.5)

# Extract priors for use with a probit model
probit_priors <- posterior_to_prior(fit_pcm, target_link = "probit")

# --- Dichotomous 1PL (fixed item effects) ---

df_rm <- eRm::raschdat3 %>%
  as.data.frame() %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_1pl <- brm(
  response ~ 0 + item + (1 | id),
```

```

    data    = df_rm,
    family  = bernoulli(),
    chains  = 4, cores = 2, iter = 1000 # use more iter (and cores if you have)
  )

  priors_1pl <- posterior_to_prior(fit_1pl)
  priors_1pl

  # Transfer logit priors to a probit refit
  priors_probit <- posterior_to_prior(fit_1pl, target_link = "probit")

```

q3_post

Summarize and Plot Posterior Predictive Q3 Residual Correlations

Description

Postprocesses the output of [q3_statistic](#) to produce summary tables of posterior predictive Q3 statistics and a combined slab + interval plot comparing observed Q3 residual correlations to the posterior predictive distribution for each item pair.

Usage

```

q3_post(
  q3_draws,
  ci = 0.84,
  n_pairs = NULL,
  sort_by = c("q3_diff", "q3_obs", "ppp")
)

```

Arguments

q3_draws	A data frame (or tibble) as returned by q3_statistic containing at minimum the columns <code>item_pair</code> , <code>q3</code> (observed Q3 per draw), and <code>q3_rep</code> (replicated Q3 per draw).
ci	Numeric in (0, 1). Width of the credible interval used for the posterior predictive HDI and the slab display. Default is 0.84.
n_pairs	Integer. Maximum number of item pairs to display in the plot, selected by largest absolute <code>q3_diff</code> . If <code>NULL</code> (the default), all pairs are shown. Useful when the number of item pairs is large.
sort_by	Character. How to order item pairs on the y-axis. <code>"q3_diff"</code> (the default) sorts by the posterior mean difference between observed and replicated Q3 (largest at top). <code>"q3_obs"</code> sorts by the posterior mean observed Q3. <code>"ppp"</code> sorts by the posterior predictive p-value.

Details

Two complementary summary tables are provided:

summary Reports posterior mean observed and replicated Q3 values alongside the posterior predictive p-value (ppp). The ppp is the proportion of draws where the observed Q3 exceeds the replicated Q3. Under good fit (no local dependence), the ppp should be near 0.5. A ppp close to 1 indicates the observed correlation is systematically higher than expected (local dependence); a ppp close to 0 indicates it is systematically lower (local repulsion, e.g., speed-accuracy tradeoffs).

hdi Reports the probability that the observed Q3 falls above (local dependence) or below (local repulsion) the HDI of the replicated distribution. This provides a more distributional assessment than the ppp alone.

The plot uses two layers from the **ggdist** package:

stat_slab Displays the posterior predictive (replicated) Q3 distribution as a filled density slab per item pair, shaded by credible interval level.

stat_slabinterval Displays the observed Q3 distribution per item pair as a semi-transparent slab with point and interval summaries.

Value

A list with three elements:

summary A **tibble** with one row per item pair containing: `item_pair`, `item_1`, `item_2`, `q3_obs` (posterior mean observed Q3), `q3_rep` (posterior mean replicated Q3), `q3_diff` (posterior mean difference), and `q3_ppp` (posterior predictive p-value: proportion of draws where the observed Q3 exceeds the replicated Q3).

hdi A **tibble** with one row per item pair containing: `item_pair`, `item_1`, `item_2`, `ld` (local dependence probability: proportion of draws where observed Q3 exceeds upper HDI bound of replicated distribution), and `lr` (local repulsion probability: proportion of draws where observed Q3 falls below lower HDI bound).

plot A **ggplot** object showing the posterior predictive distribution of replicated Q3 (grey filled slab) overlaid with the observed Q3 distribution (coloured slab + interval), with a dashed reference line at 0.

See Also

[q3_statistic](#), [infit_post](#), [item_restscore_post](#).

Examples

```
## Not run:
library(brms)
library(ggplot2)

# Assuming fit_pcm is a fitted brmsfit object
q3_draws <- q3_statistic(fit_pcm, ndraws_use = 500)
```

```

result <- q3_post(q3_draws)
result$summary
result$hdi
result$plot

# Show only top 10 pairs by Q3 difference
result_top <- q3_post(q3_draws, n_pairs = 10)
result_top$plot

## End(Not run)

```

q3_statistic	<i>Posterior Predictive Q3 Residual Correlations for Bayesian IRT Models</i>
--------------	--

Description

Computes a Bayesian analogue of Yen's Q3 statistic (Yen, 1984) for detecting local dependence between item pairs in Rasch-family models fitted with **brms**. For each posterior draw, residual correlations are computed for both observed and replicated data, yielding draw-level Q3 values that can be summarized and visualized via [q3_post](#).

Usage

```
q3_statistic(model, item_var = item, person_var = id, ndraws_use = NULL)
```

Arguments

model	A fitted brmsfit object from an ordinal IRT model (e.g., family = acat for a partial credit model) or a dichotomous model (family = bernoulli()).
item_var	An unquoted variable name identifying the item grouping variable in the model data. Default is item.
person_var	An unquoted variable name identifying the person grouping variable in the model data. Default is id.
ndraws_use	Optional positive integer. If specified, a random subset of posterior draws of this size is used. If NULL (the default), all draws are used.

Details

The procedure works as follows for each posterior draw s :

1. Compute expected values $E_{vi}^{(s)}$ from the category probabilities returned by [posterior_epred](#).
For ordinal models: $E_{vi}^{(s)} = \sum_c c \cdot P^{(s)}(X_{vi} = c)$. For binary models: $E_{vi}^{(s)} = P^{(s)}(X_{vi} = 1)$.
2. Compute observed residuals: $d_{vi}^{(s)} = X_{vi} - E_{vi}^{(s)}$.
3. Compute replicated residuals: $d_{vi}^{rep(s)} = Y_{vi}^{rep(s)} - E_{vi}^{(s)}$, where Y^{rep} is drawn via [posterior_predict](#).
4. For each item pair (i, j) , compute Q3 as the Pearson correlation of residuals across all persons who responded to both items.

Value

A [tibble](#) in long format with one row per draw per item pair, containing:

draw Integer draw index.

item_pair Character label of the item pair ("item1 : item2").

item_1 First item in the pair.

item_2 Second item in the pair.

q3 Observed Q3 residual correlation for this draw.

q3_rep Replicated Q3 residual correlation for this draw.

This long-format output parallels the structure of [infit_statistic](#) and can be passed directly to [q3_post](#) for summary tables and plots.

References

Yen, W. M. (1984). Effects of local item dependence on the fit and equating performance of the three-parameter logistic model. *Applied Psychological Measurement*, 8(2), 125–145. doi:10.1177/014662168400800201

Christensen, K. B., Makransky, G. & Horton, M. (2017). Critical values for Yen's Q3: Identification of local dependence in the Rasch model using residual correlations. *Applied Psychological Measurement*, 41(3), 178–194. doi:10.1177/0146621616677520

Bürkner, P.-C. (2021). Bayesian Item Response Modeling in R with brms and Stan. *Journal of Statistical Software*, 100, 1–54. doi:10.18637/jss.v100.i05

See Also

[q3_post](#) for postprocessing summaries and plots, [infit_statistic](#) for Bayesian infit/outfit, [posterior_epred](#), [posterior_predict](#).

Examples

```
library(brms)
library(dplyr)
library(tidyr)
library(tibble)

# --- Partial Credit Model ---

df_pcm <- eRm::pcmdat2 %>%
  mutate(across(everything(), ~ .x + 1)) %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_pcm <- brm(
  response | thres(gr = item) ~ 1 + (1 | id),
  data = df_pcm,
  family = acat,
  chains = 4,
  cores = 1, # use more cores if you have
```

```

    iter    = 500 # use at least 2000
  )

  # Q3 residual correlations
  q3_draws <- q3_statistic(fit_pcm, ndraws_use = 500)

  # Postprocess
  result <- q3_post(q3_draws)
  result$summary
  result$hdi
  result$plot

  # --- Dichotomous Rasch Model ---

  df_rm <- eRm::raschdat3 %>%
    as.data.frame() %>%
    rownames_to_column("id") %>%
    pivot_longer(!id, names_to = "item", values_to = "response")

  fit_rm <- brm(
    response ~ 1 + (1 | item) + (1 | id),
    data    = df_rm,
    family  = bernoulli(),
    chains  = 4,
    cores   = 1, # use more cores if you have
    iter    = 500 # use at least 2000
  )

  q3_draws <- q3_statistic(fit_rm, ndraws_use = 500)

  # Postprocess
  result <- q3_post(q3_draws)
  result$summary
  result$hdi
  result$plot

```

q3_statistic_hpcm

Posterior Predictive Q3 Residual Correlations for the Hurdle PCM

Description

Computes Yen's Q3 residual correlations separately for the two submodels of a hurdle partial credit model fitted with **brms** using the [hurdle_acat](#) custom family: (i) the *hurdle* submodel for $P(Y > 0)$ (Bernoulli) and (ii) the *partial credit* severity submodel for $P(Y = k \mid Y > 0)$ on the positive categories. For each posterior draw, residuals are computed from each submodel's expected values and Pearson correlations between item-pair residuals are returned for both observed and replicated data.

Usage

```
q3_statistic_hpcm(model, item_var = item, person_var = id, ndraws_use = NULL)
```

Arguments

model	A fitted <code>brmsfit</code> object using the <code>hurdle_acat</code> custom family.
item_var	An unquoted variable name identifying the item grouping variable in the model data. Default is <code>item</code> .
person_var	An unquoted variable name identifying the person grouping variable in the model data. Default is <code>id</code> .
ndraws_use	Optional positive integer. If specified, a random subset of posterior draws of this size is used. If <code>NULL</code> (the default), all draws are used.

Details

The hurdle PCM has a Bernoulli hurdle $hu = P(Y = 0)$ and a partial credit severity process on the positive categories $1, \dots, K - 1$. `posterior_epred` for `hurdle_acat` returns an $S \times N \times K_{\text{total}}$ array whose first category is hu and whose remaining categories are $(1 - hu) \cdot P_{sev}(k)$.

Hurdle residuals. For each (draw, observation):

$$d_{hurdle} = 1[Y > 0] - (1 - hu).$$

All observations contribute. Replicated residuals use $1[Y_{rep} > 0]$ from `posterior_predict`.

Partial credit residuals. For each (draw, observation) with $Y_{obs} > 0$:

$$d_{pcm} = Y - \sum_{k=1}^{K-1} k \cdot P(Y = k \mid Y > 0),$$

with conditional probabilities computed as $epred[, k + 1] / (1 - hu)$. Replicated partial credit values are drawn independently per (draw, observation) from these conditional probabilities via inverse-CDF sampling, so the partial credit PPC is not contaminated by hurdle misfit. Rows with $Y_{obs} = 0$ are set to NA in the wide residual matrix and excluded via `use = "pairwise.complete.obs"` when correlations are computed.

Value

A list with two elements, each a `tibble` in the same long format as `q3_statistic` (and directly compatible with `q3_post`):

hurdle Q3 correlations of hurdle residuals $1[Y > 0] - (1 - hu)$, using *all* observations.

pcm Q3 correlations of partial credit residuals $Y - E_{pcm}$, using only observations with $Y_{obs} > 0$. Rows with $Y_{obs} = 0$ are set to NA and excluded pairwise.

References

- Yen, W. M. (1984). Effects of local item dependence on the fit and equating performance of the three-parameter logistic model. *Applied Psychological Measurement*, 8(2), 125–145.
- Christensen, K. B., Makransky, G. & Horton, M. (2017). Critical values for Yen’s Q3: Identification of local dependence in the Rasch model using residual correlations. *Applied Psychological Measurement*, 41(3), 178–194.
- Magnus, B. E. & Garnier-Villarreal, M. (2022). A multidimensional zero-inflated graded response model for ordinal symptom data. *Psychological Methods*, 27(2), 261–279. doi:10.1037/met0000395

See Also

[q3_statistic](#) for the single-submodel version, [q3_post](#) for summarising and plotting the draws, [infit_statistic_hpcm](#) for hPCM infit.

RMUreliability	<i>Estimate reliability (Relative Measurement Uncertainty) from Bayesian measurement models</i>
----------------	---

Description

This function measures reliability using posterior draws from a fitted Bayesian model.

Usage

```
RMUreliability(input_draws, verbose = FALSE, level = 0.95)
```

Arguments

input_draws	A matrix or data frame of posterior draws. Rows represent subjects and columns represent draws.
verbose	Logical. Print detailed information about the input data. Default is TRUE.
level	Numeric. Credibility level for the highest density continuous interval. Default is 0.95.

Details

To use this function, you will need to provide a matrix (input_draws) that contains the posterior draws for the parameter you wish to calculate reliability. The function assumes that rows of input_draws represent subjects and columns represent posterior draws.

For an example of how to apply this function to calculate mean score reliability using brms, see [this tutorial](#).

For an example of how to apply this function to go/go-no task data using brms, see [this tutorial](#).

Value

A list containing:

- `hdc`: A data frame with a point-estimate (posterior mean) and highest density continuous interval for reliability, calculated using the `ggdist::mean_hdc` function
- `reliability_posterior_draws`: A numeric vector of posterior draws for reliability, of length $K/2$ (K = number of columns/draws in your `input_draws` matrix)

References

Bignardi, G., Kievit, R., & Bürkner, P. C. (2025). A general method for estimating reliability using Bayesian Measurement Uncertainty. PsyArXiv. doi:[10.31234/osf.io/h54k8](https://doi.org/10.31234/osf.io/h54k8)

Examples

```
library(brms)
library(dplyr)
library(tidyr)
library(tibble)

# --- Dichotomous Rasch Model ---

df_rm <- eRm::raschdat3 %>%
  as.data.frame() %>%
  rownames_to_column("id") %>%
  pivot_longer(!id, names_to = "item", values_to = "response")

fit_rm <- brm(
  response ~ 1 + (1 | item) + (1 | id),
  data = df_rm,
  family = bernoulli(),
  chains = 4,
  cores = 1, # use more cores if you have
  iter = 500 # use at least 2000
)

# Extract posterior draws from brms model

posterior_draws = fit_rm %>%
  as_draws_df() %>%
  as_tibble() %>%
  select(starts_with("r_id")) %>%
  t()

# Calculate RMU

RMUreliability(posterior_draws)
```

RMUreliability_hpcm *Relative Measurement Uncertainty Reliability for the Hurdle PCM*

Description

Computes posterior reliability via Relative Measurement Uncertainty (Bignardi, Kievit & Bürkner, 2025) separately for each of the two person traits in a hurdle partial credit model fitted with the `hurdle_acat` custom family: the presence / susceptibility trait θ_{hurdle} and the severity trait θ_{pcm} . Internally extracts per-submodel person draws via `person_parameters_hpcm` and applies `RMUreliability` to each.

Usage

```
RMUreliability_hpcm(
  x,
  item_var = item,
  person_var = id,
  level = 0.95,
  verbose = FALSE,
  center = TRUE
)
```

Arguments

<code>x</code>	Either: - a fitted <code>brmsfit</code> object using the <code>hurdle_acat</code> family, or - the result of <code>person_parameters_hpcm(model, draws = TRUE)</code>, i.e., a list with <code>\$hurdle\$draws_matrix</code> and <code>\$pcm\$draws_matrix</code>. Passing a pre-computed <code>person_parameters_hpcm()</code> result avoids re-extracting the posterior draws from <code>brms</code>.
<code>item_var</code>	An unquoted variable name identifying the item grouping variable in the model data. Used only when <code>x</code> is a <code>brmsfit</code> ; ignored otherwise. Default is <code>item</code> .
<code>person_var</code>	An unquoted variable name identifying the person grouping variable in the model data. Used only when <code>x</code> is a <code>brmsfit</code> ; ignored otherwise. Default is <code>id</code> .
<code>level</code>	Numeric in $(0, 1)$. Credibility level for the highest density continuous interval on each reliability. Default is <code>0.95</code> .
<code>verbose</code>	Logical. If <code>TRUE</code> , print the number of subjects and posterior draws used for each submodel. Default is <code>FALSE</code> .
<code>center</code>	Logical. If <code>TRUE</code> (the default), use centered person draws (mean item parameter = 0 in each submodel). The centering shift is a constant per draw and does not affect the correlation across draw halves, so reliability is invariant to this choice; the argument is exposed only for consistency with <code>person_parameters_hpcm</code> .

Details

Two reliabilities, two traits. The hurdle PCM gives every person a posterior on θ_{hurdle} (presence / susceptibility) and on θ_{pcm} (severity given presence). Each posterior has its own measurement uncertainty, and the RMU framework therefore yields one reliability per trait. The two values address different questions: θ_{hurdle} reliability is about how well the items rank persons by overall endorsement (a function of how many zeros appear and how discriminating the items are at the gate), while θ_{pcm} reliability is about how well the conditional severity is recovered (a function of how many positive responses each person provides and how spread out the PCM thresholds are). Reporting both is necessary, as emphasised in Magnus and Garnier-Villareal (2022, p. 277).

What about all-zero responders? For persons with $Y = 0$ on every item, no items provide information about θ_{pcm} directly. Their posterior on θ_{pcm} is shaped by the multivariate-normal prior linking it to θ_{hurdle} (and is therefore wide but well-defined). These persons still contribute to the RMU calculation; their contribution is appropriately downweighted through their large posterior variance.

Computational note. When x is a `brmsfit`, this function calls `person_parameters_hpcm(..., draws = TRUE)` internally, which extracts the full posterior draws of the person random effects. For large models you may prefer to call `person_parameters_hpcm` once and pass its result to `RMUreliability_hpcm`, `plot_targeting`, etc.

Value

A list with two elements:

`hurdle` A data frame with one row containing `rmu_estimate`, `hdci_lowerbound`, and `hdci_upperbound` for reliability of θ_{hurdle} , as returned by `RMUreliability`.

`pcm` Same format, for reliability of θ_{pcm} .

References

Bignardi, G., Kievit, R., & Bürkner, P.-C. (2025). A general method for estimating reliability using Bayesian Measurement Uncertainty. *PsyArXiv*. doi:10.31234/osf.io/h54k8_v1

Magnus, B. E. & Garnier-Villareal, M. (2022). A multidimensional zero-inflated graded response model for ordinal symptom data. *Psychological Methods*, 27(2), 261-279. doi:10.1037/met0000395

See Also

`RMUreliability` for the single-trait version, `person_parameters_hpcm` for the underlying person draws extraction.

Examples

```
## Not run:
# Fit a hurdle PCM with the hurdle_acat() family
fit <- brms::brm(
  brms::bf(
    response | brms::thres(gr = item) ~ 1 + (1 | id),
    hu ~ 0 + factor(item) + (1 | id)
  ),
  data = dat,
```

```
    family    = hurdle_acat(),
    stanvars = hurdle_acat_stanvars()
  )

# Option 1: pass the brmsfit directly
rel <- RMUreliability_hpcm(fit)
rel$hurdle
rel$pcm

# Option 2: reuse a cached person_parameters_hpcm() result
pp <- person_parameters_hpcm(fit, draws = TRUE)
rel <- RMUreliability_hpcm(pp)

## End(Not run)
```

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