Package 'rtmpt'

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```

Title Fitting RT-MPT Models

Author Raphael Hartmann [aut, cre], Karl C. Klauer [cph, aut, ctb, ths], Henrik Singmann [aut, ctb, cph], Jean Marie Linhart [cph]

Maintainer Raphael Hartmann <raphael.hartmann@protonmail.com>

Depends R (>= 3.0.0)

Imports coda, data.table, LaplacesDemon, loo, methods, stats, stringr, truncnorm, utils

Suggests knitr, rmarkdown

VignetteBuilder knitr

NeedsCompilation yes

SystemRequirements GSL (>=2.3)

Description Fit response-time extended multinomial processing tree (RT-MPT) models by Klauer and Kellen (2018) <doi:10.1016/j.jmp.2017.12.003>. The RT-MPT class not only incorporate frequencies like traditional multinomial processing tree (MPT) models, but also latencies. This enables it

to estimate process completion times and encoding plus motor execution times next to the process probabilities

of traditional MPTs. 'rtmpt' is a Bayesian framework and posterior samples are sampled using a Metropolis-Gibbs

sampler like the one described in the Klauer and Kellen (2018), but with some modifications. Other than in

the original C++ program we use the free and open source GNU Scientific Library (GSL). There is also the

possibility to suppress single process completion times.

License GPL (>= 2)

Encoding UTF-8

LazyData true

RoxygenNote 6.1.1

Archs i386, x64

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Description

Given model and data, this function calls an altered version of the C++ program by Klauer and Kellen (2018) to sample from the posterior distribution via a Metropolis-Gibbs sampler and storing it in an mcmc.list called samples. Posterior predictive checks developed by Klauer (2010), deviance information criterion (DIC; Spiegelhalter et al., 2002), 99% and 95% highest density intervals (HDI) together with the median will be provided for the main parameters in a list called diags. Optionally, the indices widely applicable information criterion (WAIC; Watanabe, 2010; Vehtari et al., 2017) and leave-one-out cross-validation (LOO; Vehtari et al., 2017) can be saved. Additionally the log-likelihood (LogLik) can also be stored. Some specifications of the function call are also saved in specs.

Usage

```
fit_rtmpt(model, data, n.chains = 4, n.iter = 5000, n.burnin = 200,
    n.thin = 1, Rhat_max = 1.05, Irep = 1000, prior_params = NULL,
    indices = FALSE, save_log_lik = FALSE, old_label = FALSE)
```

Arguments

model	A list of the class rtmpt_model.
data	Optimally, a list of class rtmpt_data. Also possible is a data. frame or a path to the text file. Both, data. frame and the text file must contain the column names "subj", "group", "tree", "cat", and "rt" preferably but not necessarily in this order. The values of the latter must be in milliseconds. It is always advised to use to_rtmpt_data first, which gives back an rtmpt_data list with informations about the changes in the data, that were needed.
n.chains	Number of chains to use. Default is 4. Must be larger than 1 and smaller or equal to 16.
n.iter	Number of samples per chain. Default is 5000.
n.burnin	Number of warm-up samples. Default is 200.
n.thin	Thinning factor. Default is 1.
Rhat_max	Maximal Potential scale reduction factor: A lower threshold that needs to be reached before the actual sampling starts. Default is 1.05

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Irep

Every Irep samples an interim state with the current maximal potential scale reduction factor is shown. Default is 1000. The following statements must hold true for Irep:

- n.burnin is smaller than or equal to Irep,
- Irep is a multiple of n. thin and
- n.iter is a multiple of Irep / n.thin.

prior_params

Named list with prior parameters. All parameters have default values, that lead to uninformative priors. Vectors are not allowed. Allowed parameters are:

- mean_of_exp_mu_beta: This is the a priori expected exponential rate (E(exp(beta)) = E(lambda)) and 1/mean_of_exp_mu_beta is the a priori expected process time (1/E(exp(beta)) = E(tau)). The default mean is set to 10, such that the expected a priori process time is 0.1 seconds.
- var_of_exp_mu_beta: The a priori group-specific variance of the exponential rates. Since exp(mu_beta) is Gamma distributed, the rate of the distribution is just mean divided by variance and the shape is the mean times the rate. The default is set to 100.
- mean_of_mu_gamma: This is the a priori expected *mean parameter* of the encoding and response execution times, which follow a normal distribution truncated from below at zero, so E(mu_gamma) < E(gamma). The default is 0.
- var_of_mu_gamma: The a priori group-specific variance of the mean parameter. Its default is 10.
- mean_of_omega_sqr: This is the a priori expected residual variance (E(omega^2)). Its distribution differs from the one used in the paper. Here it is a Gamma distribution instead of an improper one. The default is 0.005.
- var_of_omega_sqr: The a priori variance of the residual variance (Var(omega^2)).
 The default is 0.01. The default of the mean and variance is equivalent to
 a shape and rate of 0.0025 and 0.5, respectivly.
- df_of_sigma_sqr: A priori degrees of freedom for the individual variance
 of the response executions. The individual variance has a scaled inverse
 chi-squared prior with df_of_sigma_sqr degrees of freedom and omega^2
 as scale. 2 is the default and it should be an integer.
- sf_of_scale_matrix_SIGMA: The original scaling matrix (S) of the (scaled) inverse Wishart distribution for the process related parameters is an identity matrix S=I. sf_of_scale_matrix_SIGMA is a scaling factor, that scales this matrix (S=sf_of_scale_matrix_SIGMA*I). Its default is 1.
- sf_of_scale_matrix_GAMMA: The original scaling matrix (S) of the (scaled) inverse Wishart distribution for the encoding and motor execution parameters is an identity matrix S=I. sf_of_scale_matrix_GAMMA is a scaling factor, that scales this matrix (S=sf_of_scale_matrix_GAMMA*I). Its default is 1.
- prec_epsilon: This is epsilon in the paper. It is the precision of mu_alpha and all xi (scaling parameter in the scaled inverse Wishart distribution). Its default is also 1.
- add_df_to_invWish: If P is the number of parameters or rather the size of the scale matrix used in the (scaled) inverse Wishart distribution then add_df_to_invWish is the number of degrees of freedom that can be added to it. So DF = P + add_df_to_invWish. The default for add_df_to_invWish is 1, such that the correlations are uniformly distributed within [-1,1].

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Model selection indices. If set to TRUE the log-likelihood for each iteration and trial will be stored temporarily and with that the WAIC and LOO will be calculated via the loo package. If you want to have this log-likelihood matrix stored in the output of this function, you can set save_log_likelihood. TRUE. The default for indices is FALSE.

save_log_lik If set to TRUE and indices = TRUE the log-likelihood matrix for each iteration and trial will be saved in the output as a matrix. Its default is FALSE.

elements of the output list. Default is FALSE.

Value

A list of the class rtmpt_fit containing

- samples: the posterior samples as an mcmc.list object,
- diags: some diagnostics like deviance information criterion, posterior predictive checks for the frequencies and latencies, potential scale reduction factors, and also the 99% and 95% HDIs and medians for the group-level parameters,
- specs: some model specifications like the model, arguments of the model call, and information about the data transformation,
- indices (optional): if enabled, WAIC and LOO,
- LogLik (optional): if enabled, the log-likelihood matrix used for WAIC and LOO.
- summary includes posterior mean and median of the main parameters.

Author(s)

Raphael Hartmann

References

Klauer, K. C. (2010). Hierarchical multinomial processing tree models: A latent-trait approach. *Psychometrika*, 75(1), 70-98.

Klauer, K. C., & Kellen, D. (2018). RT-MPTs: Process models for response-time distributions based on multinomial processing trees with applications to recognition memory. *Journal of Mathematical Psychology*, 82, 111-130.

Spiegelhalter, D. J., Best, N. G., Carlin, B. P., & Van Der Linde, A. (2002). Bayesian measures of model complexity and fit. *Journal of the royal statistical society: Series b (statistical methodology)*, 64(4), 583-639.

Vehtari, A., Gelman, A., & Gabry, J. (2017). Practical Bayesian model evaluation using leave-one-out cross-validation and WAIC. *Statistics and Computing*, 27(5), 1413-1432.

Watanabe, S. (2010). Asymptotic equivalence of Bayes cross validation and widely applicable information criterion in singular learning theory. *Journal of Machine Learning Research*, 11(Dec), 3571-3594.

Examples

Detect-Guess variant of the Two-High Threshold model.

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```
mdl_2HTM <- "
# targets
do+(1-do)*g
(1-do)*(1-g)
# lures
(1-dn)*g
dn+(1-dn)*(1-g)
# do: detect old; dn: detect new; g: guess
model <- to_rtmpt_model(mdl_file = mdl_2HTM)</pre>
data_file <- system.file("extdata/data.txt", package="rtmpt")</pre>
data <- read.table(file = data_file, header = TRUE)</pre>
data_list <- to_rtmpt_data(raw_data = data, model = model)</pre>
# This might take some time
rtmpt_out <- fit_rtmpt(model = model, data = data_list)</pre>
rtmpt_out
# Type ?SimData for another working example.
```

fit_rtmpt_SBC

Simulate data from RT-MPT models

Description

Simulate data from RT-MPT models using rtmpt_model objects. The difference to sim_rtmpt_data is that here only scalars are allowed. This makes it usable for simulation-based calibration (SBC; Talts et al., 2018). You can specify the random seed, number of subjects, number of trials, and some parameters (same as prior_params from fit_rtmpt).

Usage

```
fit_rtmpt_SBC(model, seed, n.eff_samples = 99, n.chains = 4,
    n.iter = 5000, n.burnin = 200, n.thin = 1, Rhat_max = 1.05,
    Irep = 1000, n.subj = 40, n.trials = 30, prior_params = NULL)
```

Arguments

model	A list of the class rtmpt_model.
seed	Random seed number.
n.eff_samples	Number of effective samples. Default is 99, leading to 100 possible ranks (from 0 to 99).
n.chains	Number of chains to use. Default is 4. Must be larger than 1 and smaller or equal to 16.
n.iter	Number of samples per chain. Default is 5000. Must be larger or equal to $n.eff_samples$.
n.burnin	Number of warm-up samples. Default is 200.

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n.thin Thinning factor. Default is 1.

Maximal Potential scale reduction factor: A lower threshold that needs to be Rhat_max

reached before the actual sampling starts. Default is 1.05

Every Irep samples an interim state with the current maximal potential scale reduction factor is shown. Default is 1000. The following statements must hold true for Irep:

• n.burnin is smaller than or equal to Irep,

• Irep is a multiple of n. thin and

• n.iter is a multiple of Irep / n.thin.

Number of subjects. Default is 40.

n.trials Number of trials per tree. Default is 30.

> Named list of parameters from which the data will be generated. This must be the same named list as prior_params from fit_rtmpt and has the same defaults. It is not recommended to use the defaults since they lead to many probabilities close or equal to 0 and/or 1 and to RTs close or equal to 0. Allowed parameters are:

- mean_of_exp_mu_beta: This is the expected exponential rate (E(exp(beta))) = E(lambda)) and 1/mean_of_exp_mu_beta is the expected process time (1/E(exp(beta)) = E(tau)). The default mean is set to 10, such that the expected process time is 0.1 seconds.
- var_of_exp_mu_beta: The group-specific variance of the exponential rates. Since exp(mu_beta) is Gamma distributed, the rate of the distribution is just mean divided by variance and the shape is the mean times the rate. The default is set to 100.
- mean_of_mu_gamma: This is the expected *mean parameter* of the encoding and response execution times, which follow a normal distribution truncated from below at zero, so $E(mu_gamma) < E(gamma)$. The default is 0.
- var_of_mu_gamma: The group-specific variance of the *mean parameter*. Its default is 10.
- mean_of_omega_sqr: This is the expected residual variance (E(omega^2)). The default is 0.005.
- var_of_omega_sqr: The variance of the residual variance (Var(omega^2)). The default is 0.01. The default of the mean and variance is equivalent to a shape and rate of 0.0025 and 0.5, respectivly.
- df_of_sigma_sqr: degrees of freedom for the individual variance of the response executions. The individual variance follows a scaled inverse chisquared distribution with df_of_sigma_sqr degrees of freedom and omega^2 as scale. 2 is the default and it should be an integer.
- sf_of_scale_matrix_SIGMA: The original scaling matrix (S) of the (scaled) inverse Wishart distribution for the process related parameters is an identity matrix S=I. sf_of_scale_matrix_SIGMA is a scaling factor, that scales this matrix (S=sf_of_scale_matrix_SIGMA*I). Its default is 1.
- sf_of_scale_matrix_GAMMA: The original scaling matrix (S) of the (scaled) inverse Wishart distribution for the encoding and motor execution parameters is an identity matrix S=I. sf_of_scale_matrix_GAMMA is a scaling factor that scales this matrix (S=sf_of_scale_matrix_GAMMA*I). Its de-
- prec_epsilon: This is epsilon in the paper. It is the precision of mu_alpha and all xi (scaling parameter in the scaled inverse Wishart distribution). Its default is also 1.

n.subi

Irep

prior_params

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add_df_to_invWish: If P is the number of parameters or rather the size
 of the scale matrix used in the (scaled) inverse Wishart distribution then
 add_df_to_invWish is the number of degrees of freedom that can be added
 to it. So DF = P + add_df_to_invWish. The default for add_df_to_invWish
 is 1, such that the correlations are uniformly distributed within [-1,1].

Value

A list of the class rtmpt_sbc containing

- ranks: the rank statistic for all parameters,
- sim_list: an object of the class rtmpt_sim,
- fit_list: an object of the class rtmpt_fit,
- specs: some specifications like the model, seed number, etc.,

Author(s)

Raphael Hartmann

References

Talts, S., Betancourt, M., Simpson, D., Vehtari, A., & Gelman, A. (2018). Validating Bayesian inference algorithms with simulation-based calibration. *arXiv preprint arXiv:1804.06788*.

```
# Detect-Guess variant of the Two-High Threshold model.
# The encoding and motor execution times are assumed to be different for each response.
mdl_2HTM <- "
# targets
d+(1-d)*g
(1-d)*(1-g)
# lures
(1-d)*g
           ; 0
d+(1-d)*(1-g); 1
# d: detect; g: guess
model <- to_rtmpt_model(mdl_file = mdl_2HTM)</pre>
params <- list(mean_of_exp_mu_beta = 10,</pre>
           var_of_exp_mu_beta = 10,
           mean_of_mu_gamma = 0.5,
           var_of_mu_gamma = 0.0025,
           mean_of_omega_sqr = 0.005,
           var_of_omega_sqr = 0.000025,
           df_of_sigma_sqr = 10,
           sf_of_scale_matrix_SIGMA = 0.1,
           sf_of_scale_matrix_GAMMA = 0.01,
           prec_epsilon = 10,
```

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```
add_df_to_invWish = 5)
SBC_out <- fit_rtmpt_SBC(model, seed = 123, prior_params = params)</pre>
SBC_out$ranks
# For 2000 replications
## This takes too long to run and in addition Rhat should always be
## checked as well as the effective sample size.
R = 2000
rank_mat <- data.frame()</pre>
for (r in 1:R) {
 SBC_out <- fit_rtmpt_SBC(model, seed = r*123, prior_params = params, n.eff_samples = 99)</pre>
  rank_mat <- rbind(rank_mat, SBC_out$ranks)</pre>
}
## pearson chi square for testing uniformity
x <- apply(rank_mat[1:R,], 2, table)</pre>
expect < R/100 # 100 = number of bins/cells (0:99)
pearson <- apply(X = x, MARGIN = 2, FUN = function(x) \{sum((x-expect)^2/expect)\})
z95 \leftarrow gchisq(0.95, 99) # 99 = degrees of freedom
sum(pearson>z95) / length(pearson)
```

set_params

Set constants for probability parameters and suppress process times in a rtmpt_model list

Description

By using parameter = "probs" you can specify which of the probability parameters should be set to a constant by using values between zero and one. If you use NA the probability will be estimated. By using parameter = "tau_minus" or parameter = "tau_plus" you can suppress process times/rates. Here 0 will suppress the named process and NA allows the process time/rate to be estimated.

Usage

```
set_params(model, parameter, names, values = NA)
```

Arguments

model

A list of the class rtmpt_model.

parameter

Character of length one indicating the parameter to change. Allowed characters:

- "probs": probability parameters
- "tau_minus": rate parameters of the exponential distribution of the process times that lead to a negative outcome
- "tau_plus": rate parameters of the exponential distribution of the process times that lead to a positive outcome

names

Character vector with process names.

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values

Numerical vector of length length(names). By using parameter = "probs" you have the following options

- NA: estimate the named probability
- 0 < values < 1: set the named probability to a constant value between zero and one

Example: set_params(model = model, parameter = "probs", names = c("do", "dn", "g"), value = c(NA,NA,.5)) will set the guessing "old" (g) to the constant 0.5 in the 2HT model. By using parameter = "tau_minus" or parameter = "tau_plus" you have two options:

- NA: estimate the process time/rate
- 0: suppress the process time/rate

Example: set_params(model = model, parameter = "tau_minus", names = c("do", "dn", "g"), v = c(NA, NA, 0)) will suppress the process-completion time for guessing "new" in the 2HT model. This of course does not make sense here, but for some models it might be useful if you assume that a time-consuming process is not associated with certain process-outcome pairs (e.g., for technical parameters not corresponding to a psychological process).

Value

A list of the class rtmpt_model.

Author(s)

Raphael Hartmann

See Also

```
set_resps
```

```
# Detect-Guess variant of the Two-High Threshold model.
# The encoding and motor execution times are assumed to be equal for each category.
# The process completion times for both failed detections will be suppressed.
mdl_2HTM <- "
# targets
do+(1-do)*g
(1-do)*(1-g)
# lures
(1-dn)*g
dn+(1-dn)*(1-g)
# do: detect old; dn: detect new; g: guess
model <- to_rtmpt_model(mdl_file = mdl_2HTM)</pre>
## removing the process times for the failed detection ("tau_minus")
## of the detection parameters ("dn", "do")
```

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```
model <- set_params(model = model, parameter = "tau_minus", names = c("dn", "do"), values = c(0,0)
```

set_resps

Set responses in a rtmpt_model

Description

Change the responses for a tree and the categories within that tree.

Usage

```
set_resps(model, tree, categories, values = 0)
```

Arguments

model A list of the class rtmpt_model.

tree Character or numerical value of the tree for which the responses should be

changed.

categories Character or numerical vector identifying category/ies within the specified tree

for which the responses should be changed.

values Numerical vector of length length(categories) providing the responses. De-

fault is 0.

Value

A list of the class rtmpt_model.

Author(s)

Raphael Hartmann

See Also

```
set_params
```

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SimData

Data simulated from the restricted 2HTM

Description

Data set generated from a restricted Two-High Threshold model.

Usage

SimData

Format

A data frame with five variables: subj subjects number group group label of the subjects tree condition of the current trial cat observed response category rt observed response time in ms

Details

Fourty subjects with thirty trials per condition (Studied items, new Items) were simulated.

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```
d+(1-d)*(1-g) ; 1

# d: detect; g: guess
"

model <- to_rtmpt_model(mdl_file = mdl_2HTM)

data <- to_rtmpt_data(raw_data = SimData, model = model)

# this might take some time to run

rtmpt_out <- fit_rtmpt(model = model, data = data)

# convergence
## traceplot and summary of the first six parameters
plot(rtmpt_out$samples[,1:6])
summary(rtmpt_out)</pre>
```

sim_rtmpt_data

Simulate data from RT-MPT models

Description

Simulate data from RT-MPT models using rtmpt_model objects. You can specify the random seed, number of subjects, number of trials per tree, and some parameters (mainly the same as prior_params from fit_rtmpt).

Usage

```
sim_rtmpt_data(model, seed, n.subj, n.trials, params = NULL)
```

Arguments

params

model A list of the class rtmpt_model.

seed Random seed number.n.subj Number of subjects.n.trials Number of trials per tree.

Named list of parameters from which the data will be generated. This must be the same named list as prior_params from fit_rtmpt, except for "mean_of_mu_alpha" and "var_of_mu_alpha", and has the same defaults. The difference to prior_params is, that vectors are allowed, but must match the length of the parameters in the model. It is not recommended to use the defaults since they lead to many probabilities close or equal to 0 and/or 1 and to RTs close or equal to 0. Allowed parameters are:

- mean_of_mu_alpha: Probit transformed mean probability. If you want to have a group-level mean probability of 0.6, use mean_of_mu_alpha = qnorm(0.6) in the params list. Default is 0 or qnorm(.5).
- var_of_mu_alpha: Variance of the probit transformed group-level mean probability. If specified, mu_alpha will be sampled from N(mean_of_mu_alpha, var_of_mu_alpha. If not, mu_alpha = mean_of_mu_alpha.

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• mean_of_exp_mu_beta: This is the expected exponential rate (E(exp(beta)) = E(lambda)) and 1/mean_of_exp_mu_beta is the expected process time (1/E(exp(beta)) = E(tau)). The default mean is set to 10, such that the expected process time is 0.1 seconds. For a mean process time of 200 ms, wirte mean_of_exp_mu_beta = 1000/200.

- var_of_exp_mu_beta: The group-specific variance of the exponential rates.
 Since exp(mu_beta) is Gamma distributed, the rate of the distribution is
 just mean divided by variance and the shape is the mean times the rate. If
 specified, exp(mu_beta) is sampled from Gammashape = mean_of_exp_mu_beta^2/var_of_ex
 = mean_of_exp_mu_beta/var_of_exp_mu_beta). If not, mu_alpha = mean_of_exp_mu_beta.
- mean_of_mu_gamma: This is the expected *mean parameter* of the encoding and response execution times, which follow a normal distribution truncated from below at zero, so E(mu_gamma) < E(gamma). The default is 0. For a mean motor time of 550 ms write mean_of_mu_gamma = 550/1000.
- var_of_mu_gamma: The group-specific variance of the *mean parameter*. If specified, mu_gamma is sampled from N(mean_of_mu_gamma, var_of_mu_gamma). If not, mu_gamma = mean_of_mu_gamma.
- mean_of_omega_sqr: This is the expected residual variance (E(omega^2)).
 The default is 0.005.
- var_of_omega_sqr: The variance of the residual variance (Var(omega^2)).
 If specified, omega_sqr is sampled from GAMMA(shape = mean_of_omega_sqr^2/var_of_omega_sqr/omega_sqr/var_of_omega_sqr). If not, omega_sqr = mean_of_omega_sqr.
 0.01. The default of the mean and variance is equivalent to a shape and rate of 0.0025 and 0.5, respectively.
- df_of_sigma_sqr: Degrees of freedom for the individual variance of the response executions. The individual variance follows a scaled inverse chisquared distribution with df_of_sigma_sqr degrees of freedom and omega^2 as scale. 2 is the default and it should be an integer.
- sf_of_scale_matrix_SIGMA: The original scaling matrix (S) of the (scaled) inverse Wishart distribution for the process related parameters is an identity matrix S=I. sf_of_scale_matrix_SIGMA is a scaling factor, that scales this matrix (S=sf_of_scale_matrix_SIGMA*I). Its default is 1.
- sf_of_scale_matrix_GAMMA: The original scaling matrix (S) of the (scaled) inverse Wishart distribution for the encoding and motor execution parameters is an identity matrix S=I. sf_of_scale_matrix_GAMMA is a scaling factor that scales this matrix (S=sf_of_scale_matrix_GAMMA*I). Its defoult is 1
- prec_epsilon: This is epsilon in the paper. It is the precision of xi (scaling parameter in the scaled inverse Wishart distribution). Its default is also 1.
- add_df_to_invWish: If P is the number of parameters or rather the size of the scale matrix used in the (scaled) inverse Wishart distribution then add_df_to_invWish is the number of degrees of freedom that can be added to it. So DF = P + add_df_to_invWish. The default for add_df_to_invWish is 1, such that the correlations are uniformly distributed within [-1,1].
- SIGMA: Variance-covariance matrix of the process-related parameters. It must match the number of process-related parameters to be estimated. If scalars or vectors are given, they will be transformed into diagonal matrices using diag(SIGMA). If not specified it will be randomly generated using diag(xi)%*%LaplacesDemon::rinvwishart(nu,S)%*%diag(xi), where nu is the number of process-related group-level parameters to be estimated plus add_df_to_invWish, S is the identity matrix multiplied by sf_of_scale_matrix_SIGMA,

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and xi (randomly generated from N(1,1/prec_epsilon)) are the scaling factors for the scaled inverse wishart distribution. If SIGMA is used, sf_of_scale_matrix_SIGMA and add_df_to_invWish will be ignored for the process-related parameters.

• GAMMA: Variance-covariance matrix of the motor time parameters. It must match the number of motor time parameters to be estimated. If scalars or vectors are given, they will be transformed into diagonal matrices using diag(SIGMA). If not specified it will be randomly generated using diag(xi)%*%LaplacesDemon where nu is the number of motor time group-level parameters to be estimated plus add_df_to_invWish, S is the identity matrix multiplied by sf_of_scale_matrix_GAMMA, and xi (randomly generated from N(1,1/prec_epsilon)) are the scaling factors for the scaled inverse wishart distribution. If GAMMA is used, sf_of_scale_matrix_GAMMA and add_df_to_invWish will be ignored for the motor time parameters.

Value

A list of the class rtmpt_sim containing

- data: the data.frame with the simulated data,
- gen_list: a list containing lists of the group-level and subject-specific parameters for the
 process-related parameters and the motor-related parameters, and the trial-specific probabilities, process-times, and motor-times,
- specs: some specifications like the model, seed number, etc.,

Author(s)

Raphael Hartmann

```
# Detect-Guess variant of the Two-High Threshold model.
# The encoding and motor execution times are assumed to be different for each response.
mdl_2HTM <- "
# targets
do+(1-do)*g
           ; 0
(1-do)*(1-g)
           ; 1
# lures
(1-dn)*g
dn+(1-dn)*(1-g); 1
# do: detect old; dn: detect new; g: guess
model <- to_rtmpt_model(mdl_file = mdl_2HTM)</pre>
# random group-level parameters
params <- list(mean_of_mu_alpha = 0,</pre>
           #var_of_mu_alpha = 1
           mean_of_exp_mu_beta = 10,
           var_of_exp_mu_beta = 10,
```

sim_rtmpt_data_SBC

```
mean_of_mu_gamma = 0.5,
               var_of_mu_gamma = 0.0025,
               mean\_of\_omega\_sqr = 0.005
               var_of_omega_sqr = 0.000025,
               df_of_sigma_sqr = 10,
               sf_of_scale_matrix_SIGMA = 0.1,
               sf_of_scale_matrix_GAMMA = 0.01,
               prec_epsilon = 10,
               add_df_to_invWish = 5)
sim_dat <- sim_rtmpt_data(model, seed = 123, n.subj = 40, n.trials = 30, params = params)</pre>
# fixed group-level parameters
params <- list(mean_of_mu_alpha = 0,</pre>
               mean_of_exp_mu_beta = 10,
               mean_of_mu_gamma = 0.5,
               mean_of_omega_sqr = 0.005,
               df_of_sigma_sqr = 10,
               sf_of_scale_matrix_SIGMA = 0.1,
               sf_of_scale_matrix_GAMMA = 0.01,
               prec_epsilon = 10,
               add_df_to_invWish = 5,
               SIGMA = diag(9), # independent process-related params
               GAMMA = diag(2)) # independent motor time params
sim_dat <- sim_rtmpt_data(model, seed = 123, n.subj = 40, n.trials = 30, params = params)</pre>
```

sim_rtmpt_data_SBC

Simulate data from RT-MPT models

Description

Simulate data from RT-MPT models using rtmpt_model objects. The difference to sim_rtmpt_data is that here only scalars are allowed. This makes it usable for simulation-based calibration (SBC; Talts et al., 2018). You can specify the random seed, number of subjects, number of trials, and some parameters (same as prior_params from fit_rtmpt).

Usage

```
sim_rtmpt_data_SBC(model, seed, n.subj, n.trials, params = NULL)
```

Arguments

model	A list of the class rtmpt_model.
seed	Random seed number.
n.subj	<- Number of subjects.
n.trials	<- Number of trials per tree.
params	Named list of parameters from

Named list of parameters from which the data will be generated. This must be the same named list as prior_params from fit_rtmpt and has the same defaults. It is not recommended to use the defaults since they lead to many probabilities close or equal to 0 and/or 1 and to RTs close or equal to 0. Allowed

parameters are:

- mean_of_exp_mu_beta: This is the expected exponential rate (E(exp(beta)) = E(lambda)) and 1/mean_of_exp_mu_beta is the expected process time (1/E(exp(beta)) = E(tau)). The default mean is set to 10, such that the expected process time is 0.1 seconds.
- var_of_exp_mu_beta: The group-specific variance of the exponential rates. Since exp(mu_beta) is Gamma distributed, the rate of the distribution is just mean divided by variance and the shape is the mean times the rate. The default is set to 100.
- mean_of_mu_gamma: This is the expected *mean parameter* of the encoding and response execution times, which follow a normal distribution truncated from below at zero, so E(mu_gamma) < E(gamma). The default is 0.
- var_of_mu_gamma: The group-specific variance of the mean parameter. Its
 default is 10.
- mean_of_omega_sqr: This is the expected residual variance (E(omega^2)).
 The default is 0.005.
- var_of_omega_sqr: The variance of the residual variance (Var(omega^2)). The default is 0.01. The default of the mean and variance is equivalent to a shape and rate of 0.0025 and 0.5, respectivly.
- df_of_sigma_sqr: degrees of freedom for the individual variance of the response executions. The individual variance follows a scaled inverse chi-squared distribution with df_of_sigma_sqr degrees of freedom and omega^2 as scale. 2 is the default and it should be an integer.
- sf_of_scale_matrix_SIGMA: The original scaling matrix (S) of the (scaled) inverse Wishart distribution for the process related parameters is an identity matrix S=I. sf_of_scale_matrix_SIGMA is a scaling factor, that scales this matrix (S=sf_of_scale_matrix_SIGMA*I). Its default is 1.
- sf_of_scale_matrix_GAMMA: The original scaling matrix (S) of the (scaled) inverse Wishart distribution for the encoding and motor execution parameters is an identity matrix S=I. sf_of_scale_matrix_GAMMA is a scaling factor that scales this matrix (S=sf_of_scale_matrix_GAMMA*I). Its default is 1.
- prec_epsilon: This is epsilon in the paper. It is the precision of mu_alpha and all xi (scaling parameter in the scaled inverse Wishart distribution). Its default is also 1.
- add_df_to_invWish: If P is the number of parameters or rather the size of the scale matrix used in the (scaled) inverse Wishart distribution then add_df_to_invWish is the number of degrees of freedom that can be added to it. So DF = P + add_df_to_invWish. The default for add_df_to_invWish is 1, such that the correlations are uniformly distributed within [-1,1].

Value

A list of the class rtmpt_sim containing

- data: the data.frame with the simulated data,
- gen_list: a list containing lists of the group-level and subject-specific parameters for the
 process-related parameters and the motor-related parameters, and the trial-specific probabilities, process-times, and motor-times,
- specs: some specifications like the model, seed number, etc.,

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Author(s)

Raphael Hartmann

References

Talts, S., Betancourt, M., Simpson, D., Vehtari, A., & Gelman, A. (2018). Validating Bayesian inference algorithms with simulation-based calibration. *arXiv preprint arXiv:1804.06788*.

Examples

```
# Detect-Guess variant of the Two-High Threshold model.
\mbox{\tt\#} The encoding and motor execution times are assumed to be different for each response.
mdl_2HTM <- "
# targets
do+(1-do)*g
             ; 0
(1-do)*(1-g)
             ; 1
# lures
(1-dn)*g
             ; 0
dn+(1-dn)*(1-g); 1
# do: detect old; dn: detect new; g: guess
model <- to_rtmpt_model(mdl_file = mdl_2HTM)</pre>
params <- list(mean_of_exp_mu_beta = 10,</pre>
            var_of_exp_mu_beta = 10,
            mean_of_mu_gamma = 0.5,
            var_of_mu_gamma = 0.0025,
            mean\_of\_omega\_sqr = 0.005,
            var_of_omega_sqr = 0.000025,
            df_of_sigma_sqr = 10,
            sf_of_scale_matrix_SIGMA = 0.1,
            sf_of_scale_matrix_GAMMA = 0.01,
            prec_epsilon = 10,
            add_df_to_invWish = 5)
sim_dat <- rtmpt:::sim_rtmpt_data_SBC(model, seed = 123, n.subj = 40,</pre>
                                n.trials = 30, params = params)
```

 to_rtmpt_data

Transform data for use in fit_rtmpt

Description

Transform data, such that it can be used in fit_rtmpt. This implies changing each value/label in "subj", "group", "tree", and "cat" to numbers such that it starts from zero (e.g. data\$tree = c(1,1,3,3,2,2,...) will be changed to data\$tree = c(0,0,2,2,1,1,...)) and the columns will be ordered

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in the right way. "rt" must be provided in milliseconds. If it has decimal places it will be rounded to a whole number. fit_rtmpt will automatically call this function if its input is not already an rtmpt_data list, but it is advised to use it anyway because it provides information about the transformations of the data.

Usage

```
to_rtmpt_data(raw_data, model)
```

Arguments

raw_data data.frame or path to data containing columns "subj", "group", "tree", "cat", and "rt". If not provided in this order it will be reordered and unused variables

will be moved to the end of the new data frame.

model A list of the class rtmpt_model.

Value

A list of the class rtmpt_data containing transformed data and information about the transformation that has been done.

Author(s)

Raphael Hartmann

```
# Detect-Guess variant of the Two-High Threshold model.
# The encoding and motor execution times are assumed to be equal for each response.
eqn_2HTM <- "
# CORE MPT EQN
# tree ; cat ; mpt
target ; hit ; do
target; hit; (1-do)*g
target; miss; (1-do)*(1-g)
 lure ; f_a ; (1-dn)*g
 lure; c_r; dn
 lure; c_r; (1-dn)*(1-g)
model <- to_rtmpt_model(eqn_file = eqn_2HTM)</pre>
data_file <- system.file("extdata/labeled_data.txt", package="rtmpt")</pre>
data <- read.table(file = data_file, header = TRUE)</pre>
data_list <- to_rtmpt_data(raw_data = data, model = model)</pre>
data_list
```

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to_rtmpt_model Cred	<pre>ate a model list for fit_rtmpt</pre>
---------------------	---

Description

Create a model list of the class rtmpt_model by providing either eqn_file or mdl_file. If both are provided mdl_file will be used.

Usage

```
to_rtmpt_model(eqn_file = NULL, mdl_file = NULL)
```

Arguments

eqn_file	Character string as shown in example 2 or path to the text file that specifies the (RT-)MPT model with standard .eqn syntax (Heck et al., 2018; Hu, 1999). E.g. studied; hit; (1-do)*g for a correct guess in the detect-guess 2HT model.
mdl_file	Character string as shown in example 1 or path to the text file that specifies the (RT-)MPT model and gives on each line the equation of one category using + to separate branches and * to separate processes (Singmann and Kellen, 2013). E.g. do+(1-do)*g for the category "hit" in the detect-guess 2HT model.

Value

A list of the class rtmpt_model.

Note

Within a branch of a (RT-)MPT model it is not allowed to have the same process two or more times.

Author(s)

Raphael Hartmann

References

Heck, D. W., Arnold, N. R., & Arnold, D. (2018). TreeBUGS: An R package for hierarchical multinomial-processing-tree modeling. *Behavior Research Methods*, 50(1), 264-284.

Hu, X. (1999). Multinomial processing tree models: An implementation. *Behavior Research Methods, Instruments, & Computers, 31(4),* 689-695.

Singmann, H., & Kellen, D. (2013). MPTinR: Analysis of multinomial processing tree models in R. *Behavior Research Methods*, 45(2), 560-575.

See Also

- set_params
- set_resps

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```
# Detect-Guess variant of the Two-High Threshold model
# with constant guessing and
# suppressed process completion times for both failed detections.
# The encoding and motor execution times are assumed to be different for each response.
\#\# 1. using the mdl syntax
mdl_2HTM <- "
# targets
do+(1-do)*g
           ; 0
(1-do)*(1-g); 1
# lures
(1-dn)*g
           ; 0
dn+(1-dn)*(1-g); 1
# do: detect old; dn: detect new; g: guess
# OPTIONAL MPT CONSTRAINTS
# set probabilities to constants:
const_prob: g=0.5
# suppress process times:
suppress_process: dn-, do-
model <- to_rtmpt_model(mdl_file = mdl_2HTM)</pre>
model
## 2. using the eqn syntax
eqn_2HTM <- "
# CORE MPT EQN
# tree ; cat ; mpt
   0; 0; do
    0; 0; (1-do)*g
    0; 1; (1-do)*(1-g)
   1; 2; (1-dn)*g
    1 ; 3 ; dn
    1; 3; (1-dn)*(1-g)
# OPTIONAL MPT CONSTRAINTS
# set probabilities to constants:
const_prob: g=0.5
# suppress process times:
suppress_process: dn-, do-
    tree ; cat ; RESP
resp: 0; 0; 0
resp:
       0; 1;
                 1
resp:
     1; 2;
      1; 3;
# different motor execution times for old and new responses.
```

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"

```
model <- to_rtmpt_model(eqn_file = eqn_2HTM)
model</pre>
```

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