

Package ‘missingHE’

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Type Package

Title Missing Outcome Data in Health Economic Evaluation

Version 1.3.2

Description Contains a suite of functions for health economic evaluations with missing outcome data.

The package can fit different types of statistical models under a fully Bayesian approach using the software 'JAGS' (which should be installed locally and which is loaded in 'missingHE' via the 'R' package 'R2jags').

Three classes of models can be fitted under a variety of missing data assumptions: selection models, pattern mixture models and hurdle models.

In addition to model fitting, 'missingHE' provides a set of specialised functions to assess model convergence and fit, and to summarise the statistical and economic results using different types of measures and graphs.

The methods implemented are described in Mason (2018) <doi:10.1002/hec.3793>, Molenberghs (2000) <doi:10.1007/978-1-4419-0300-6_18> and Gabrio (2019) <doi:10.1002/sim.8045>.

Depends R (>= 3.5.0)

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data_read_hurdle	<i>A function to read and re-arrange the data in different ways for the hurdle model</i>
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Description

This internal function imports the data and outputs only those variables that are needed to run the hurdle model according to the information provided by the user.

Usage

```
data_read_hurdle(data, model.eff, model.cost, model.se, model.sc, se, sc,
  type = type, center)
```

Arguments

data	A data frame in which to find variables supplied in model.eff, model.cost (model formulas for effects and costs) and model.se, model.sc (model formulas for the structural effect and cost models). Among these, effectiveness, cost and treatment indicator (only two arms) variables must always be provided and named 'e', 'c' and 't' respectively.
------	---

model.eff	A formula expression in conventional R linear modelling syntax. The response must be a health economics effectiveness outcome ('e') whose name must correspond to that used in data, and any covariates are given on the right-hand side. If there are no covariates, specify 1 on the right hand side. By default, covariates are placed on the "location" parameter of the distribution through a linear model.
model.cost	A formula expression in conventional R linear modelling syntax. The response must be a health economics cost outcome ('c') whose name must correspond to that used in data, and any covariates are given on the right-hand side. If there are no covariates, specify 1 on the right hand side. By default, covariates are placed on the "location" parameter of the distribution through a linear model.
model.se	A formula expression in conventional R linear modelling syntax. The response must be a health economics effectiveness outcome ('e') whose name must correspond to that used in data, and any covariates used to estimate the probability of structural effects are given on the right-hand side. If there are no covariates, specify 1 on the right hand side. By default, covariates are placed on the "probability" parameter for the structural effects through a logistic-linear model.
model.sc	A formula expression in conventional R linear modelling syntax. The response must be a health economics cost outcome ('c') whose name must correspond to that used in data, and any covariates used to estimate the probability of structural costs are given on the right-hand side. If there are no covariates, specify 1 on the right hand side. By default, covariates are placed on the "probability" parameter for the structural costs through a logistic-linear model.
se	Structural value to be found in the effect data defined in data. If set to NULL, no structural value is chosen and a standard model for the effects is run.
sc	Structural value to be found in the cost data defined in data. If set to NULL, no structural value is chosen and a standard model for the costs is run.
type	Type of structural value mechanism assumed, either 'SCAR' (Structural Completely At Random) or 'SAR' (Structural At Random).
center	Logical. If center is TRUE all the covariates in the model are centered.

Examples

```
#Internal function only
#no examples
#
#
```

data_read_pattern *A function to read and re-arrange the data in different ways*

Description

This internal function imports the data and outputs only those variables that are needed to run the model according to the information provided by the user.

Usage

```
data_read_pattern(data, model.eff, model.cost, type = type, center)
```

Arguments

data	A data frame in which to find variables supplied in <code>model.eff</code> and <code>model.cost</code> . Among these, effectiveness, cost and treatment indicator (only two arms) variables must always be provided and named 'e', 'c' and 't' respectively.
model.eff	A formula expression in conventional R linear modelling syntax. The response must be a health economics effectiveness outcome ('e') whose name must correspond to that used in <code>data</code> , and any covariates are given on the right-hand side. If there are no covariates, specify 1 on the right hand side. By default, covariates are placed on the "location" parameter of the distribution through a linear model.
model.cost	A formula expression in conventional R linear modelling syntax. The response must be a health economics cost outcome ('c') whose name must correspond to that used in <code>data</code> , and any covariates are given on the right-hand side. If there are no covariates, specify 1 on the right hand side. By default, covariates are placed on the "location" parameter of the distribution through a linear model.
type	Type of missingness mechanism assumed. Choices are Missing At Random (MAR) and Missing Not At Random (MNAR).
center	Logical. If center is TRUE all the covariates in the model are centered.

Examples

```
#Internal function only
#no examples
#
#
```

data_read_selection *A function to read and re-arrange the data in different ways*

Description

This internal function imports the data and outputs only those variables that are needed to run the model according to the information provided by the user.

Usage

```
data_read_selection(data, model.eff, model.cost, model.me, model.mc,
  type = type, center)
```

Arguments

<code>data</code>	A data frame in which to find variables supplied in <code>model.eff</code> and <code>model.cost</code> . Among these, effectiveness, cost and treatment indicator (only two arms) variables must always be provided and named 'e', 'c' and 't' respectively.
<code>model.eff</code>	A formula expression in conventional R linear modelling syntax. The response must be a health economics effectiveness outcome ('e') whose name must correspond to that used in <code>data</code> , and any covariates are given on the right-hand side. If there are no covariates, specify 1 on the right hand side. By default, covariates are placed on the "location" parameter of the distribution through a linear model.
<code>model.cost</code>	A formula expression in conventional R linear modelling syntax. The response must be a health economics cost outcome ('c') whose name must correspond to that used in <code>data</code> , and any covariates are given on the right-hand side. If there are no covariates, specify 1 on the right hand side. By default, covariates are placed on the "location" parameter of the distribution through a linear model.
<code>model.me</code>	A formula expression in conventional R linear modelling syntax. The response must be indicated with the term 'me'(missing effects) and any covariates used to estimate the probability of missing effects are given on the right-hand side. If there are no covariates, specify 1 on the right hand side. By default, covariates are placed on the "probability" parameter for the missing effects through a logistic-linear model.
<code>model.mc</code>	A formula expression in conventional R linear modelling syntax. The response must be indicated with the term 'mc'(missing costs) and any covariates used to estimate the probability of missing costs should be given on the right-hand side. If there are no covariates, specify 1 on the right hand side. By default, covariates are placed on the "probability" parameter for the missing costs through a logistic-linear model.
<code>type</code>	Type of missingness mechanism assumed. Choices are Missing At Random (MAR) and Missing Not At Random (MNAR).
<code>center</code>	Logical. If center is TRUE all the covariates in the model are centered.

Examples

```
#Internal function only
#no examples
#
#
```

<code>diagnostic</code>	<i>Diagnostic checks for assessing MCMC convergence of Bayesian models fitted in JAGS using the function selection, pattern or hurdle</i>
-------------------------	---

Description

The focus is restricted to full Bayesian models in cost-effectiveness analyses based on the function `selection`, `pattern` and `hurdle`, with convergence of the MCMC chains that is assessed through graphical checks of the posterior distribution of the parameters of interest, Examples are density plots, trace plots, autocorrelation plots, etc. Other types of posterior checks are related to some summary MCMC statistics that are able to detect possible issues in the convergence of the algorithm, such as the potential scale reduction factor or the effective sample size. Different types of diagnostic tools and statistics are used to assess model convergence using functions contained in the package `ggmcmc` and `mcmcplots`. Graphics and plots are managed using functions contained in the package `ggplot2` and `ggthemes`.

Usage

```
diagnostic(x, type = "histogram", param = "all", theme = NULL, ...)
```

Arguments

<code>x</code>	An object of class "missingHE" containing the posterior results of a full Bayesian model implemented using the function <code>selection</code> , <code>pattern</code> or <code>hurdle</code> .
<code>type</code>	Type of diagnostic check to be plotted for the model parameter selected. Available choices include: 'histogram' for histogram plots, 'denplot' for density plots, 'traceplot' for trace plots, 'acf' for autocorrelation plots, 'running' for running mean plots, 'compare' for comparing the distribution of the whole chain with only its last part, 'cross' for crosscorrelation plots, 'Rhat' for the potential scale reduction factor, 'geweke' for the geweke diagnostic, 'pairs' for posterior correlation among the parameters, 'caterpillar' for caterpillar plots. In addition the class 'summary' provides an overview of some of the most popular diagnostic checks for each parameter selected.
<code>param</code>	Name of the family of parameters to process, as given by a regular expression. For example the mean parameters for the effect and cost variables can be specified using 'mu.e' and 'mu.c', respectively. Different types of models may have different parameters depending on the assumed distributions and missing data assumptions. To see a complete list of all possible parameters by types of models assumed see details.
<code>theme</code>	Type of ggplot theme among some pre-defined themes. For a full list of available themes see details.
<code>...</code>	Additional parameters that can be provided to manage the graphical output of diagnostic.

Details

Depending on the types of plots specified in the argument `type`, the output of `diagnostic` can produce different combinations of MCMC visual posterior checks for the family of parameters indicated in the argument `param`. For a full list of the available plots see the description of the argument `type` or see the corresponding plots in the package `ggmcmc`.

The parameters that can be assessed through `diagnostic` are only those included in the object `x` (see Arguments). Specific character names must be specified in the argument `param` according to the specific model implemented. The available names and the parameters associated with them are:

- "mu.e" the mean parameters of the effect variables in the two treatment arms.
- "mu.c" the mean parameters of the cost variables in the two treatment arms.
- "mu.e.p" the pattern-specific mean parameters of the effect variables in the two treatment arms (only with the function `pattern`).
- "mu.c.p" the pattern-specific mean parameters of the cost variables in the two treatment arms (only with the function `pattern`).
- "sd.e" the standard deviation parameters of the effect variables in the two treatment arms.
- "sd.c" the standard deviation parameters of the cost variables in the two treatment arms.
- "alpha" the regression intercept and covariate coefficient parameters for the effect variables in the two treatment arms.
- "beta" the regression intercept and covariate coefficient parameters for the cost variables in the two treatment arms.
- "p.e" the probability parameters of the missingness or structural values mechanism for the effect variables in the two treatment arms (only with the function `selection` or `hurdle`).
- "p.c" the probability parameters of the missingness or structural values mechanism for the cost variables in the two treatment arms (only with the function `selection` or `hurdle`).
- "gamma.e" the regression intercept and covariate coefficient parameters of the missingness or structural values mechanism for the effect variables in the two treatment arms (only with the function `selection` or `hurdle`).
- "gamma.c" the regression intercept and covariate coefficient parameters of the missingness or structural values mechanism for the cost variables in the two treatment arms (only with the function `selection` or `hurdle`).
- "pattern" the probabilities associated with the missingness patterns in the data (only with the function `pattern`).
- "delta.e" the mnr parameters of the missingness mechanism for the effect variables in the two treatment arms (only with the function `selection` or `pattern`).
- "delta.c" the mnr parameters of the missingness mechanism for the cost variables in the two treatment arms (only with the function `selection` or `pattern`).
- "all" all available parameters stored in the object `x`.

When the object `x` is created using the function `pattern`, pattern-specific standard deviation ("sd.e", "sd.c") and regression coefficient parameters ("alpha", "beta") for both outcomes can be visualised. The parameters associated with a missingness mechanism can be accessed only when `x` is created using the function `selection` or `pattern`, while the parameters associated with the model for the structural values mechanism can be accessed only when `x` is created using the function `hurdle`.

The argument `theme` allows to customise the graphical output of the plots generated by `diagnostic` and allows to choose among a set of possible pre-defined themes taken from the package **ggthemes**. For a complete list of the available character names for each theme, see **ggthemes**.

Value

A **ggplot** object containing the plots specified in the argument `type`

Author(s)

Andrea Gabrio

References

Gelman, A. Carlin, JB., Stern, HS. Rubin, DB.(2003). *Bayesian Data Analysis, 2nd edition*, CRC Press.

Brooks, S. Gelman, A. Jones, JL. Meng, XL. (2011). *Handbook of Markov Chain Monte Carlo*, CRC/Chapman and Hall.

See Also

[ggs selection selection hurdle](#).

Examples

```
#For examples see the function selection, pattern or hurdle
#
#
```

hurdle

Full Bayesian Models to handle missingness in Economic Evaluations (Hurdle Models)

Description

Full Bayesian cost-effectiveness models to handle missing data in the outcomes using Hurdle models under a variety of alternative parametric distributions for the effect and cost variables. Alternative assumptions about the mechanisms of the structural values are implemented using a hurdle approach. The analysis is performed using the BUGS language, which is implemented in the software JAGS using the function [jags](#). The output is stored in an object of class 'missingHE'.

Usage

```
hurdle(data, model.eff, model.cost, model.se = se ~ 1, model.sc = sc ~
  1, se = 1, sc = 0, dist_e, dist_c, type, prob = c(0.05, 0.95),
  n.chains = 2, n.iter = 20000, n.burnin = floor(n.iter/2),
  inits = NULL, n.thin = 1, ppc = FALSE, save_model = FALSE,
  prior = "default", ...)
```

Arguments

data A data frame in which to find the variables supplied in `model.eff`, `model.cost` (model formulas for effects and costs) and `model.se`, `model.sc` (model formulas for the structural effect and cost models). Among these, effectiveness, cost and treatment indicator (only two arms) variables must always be provided and named 'e', 'c' and 't', respectively.

<code>model.eff</code>	A formula expression in conventional R linear modelling syntax. The response must be a health economic effectiveness outcome ('e') whose name must correspond to that used in data. Any covariates in the model must be provided on the right-hand side of the formula. If there are no covariates, 1 should be specified on the right hand side of the formula. By default, covariates are placed on the "location" parameter of the distribution through a linear model.
<code>model.cost</code>	A formula expression in conventional R linear modelling syntax. The response must be a health economic cost outcome ('c') whose name must correspond to that used in data. Any covariates in the model must be provided on the right-hand side of the formula. If there are no covariates, 1 should be specified on the right hand side of the formula. By default, covariates are placed on the "location" parameter of the distribution through a linear model. A joint bivariate distribution for effects and costs can be specified by including 'e' on the right-hand side of the formula for the costs model.
<code>model.se</code>	A formula expression in conventional R linear modelling syntax. The response must be indicated with the term 'se'(structural effects). Any covariates in the model must be provided on the right-hand side of the formula. If there are no covariates, 1 should be specified on the right hand side of the formula. By default, covariates are placed on the "probability" parameter for the structural effects through a logistic-linear model.
<code>model.sc</code>	A formula expression in conventional R linear modelling syntax. The response must be indicated with the term 'sc'(structural costs). Any covariates in the model must be provided on the right-hand side of the formula. If there are no covariates, 1 should be specified on the right hand side of the formula. By default, covariates are placed on the "probability" parameter for the structural costs through a logistic-linear model.
<code>se</code>	Structural value to be found in the effect variables defined in data. If set to NULL, no structural value is chosen and a standard model for the effects is run.
<code>sc</code>	Structural value to be found in the cost variables defined in data. If set to NULL, no structural value is chosen and a standard model for the costs is run.
<code>dist_e</code>	Distribution assumed for the effects. Current available choices are: Normal ('norm'), Beta ('beta'), Gamma ('gamma'), Exponential ('exp'), Weibull ('weibull'), Logistic ('logis'), Poisson ('pois'), Negative Binomial ('nbinom') or Bernoulli ('bern').
<code>dist_c</code>	Distribution assumed for the costs. Current available choices are: Normal ('norm'), Gamma ('gamma') or LogNormal ('lnorm').
<code>type</code>	Type of structural value mechanism assumed. Choices are Structural Completely At Random (SCAR), and Structural At Random (SAR).
<code>prob</code>	A numeric vector of probabilities within the range (0,1), representing the upper and lower CI sample quantiles to be calculated and returned for the imputed values.
<code>n.chains</code>	Number of chains.
<code>n.iter</code>	Number of iterations.
<code>n.burnin</code>	Number of warmup iterations.

<code>inits</code>	A list with elements equal to the number of chains selected; each element of the list is itself a list of starting values for the JAGS model, or a function creating (possibly random) initial values. If <code>inits</code> is <code>NULL</code> , JAGS will generate initial values for all the model parameters.
<code>n.thin</code>	Thinning interval.
<code>ppc</code>	Logical. If <code>ppc</code> is <code>TRUE</code> , the estimates of the parameters that can be used to generate replications from the model are saved.
<code>save_model</code>	Logical. If <code>save_model</code> is <code>TRUE</code> a <code>txt</code> file containing the model code is printed in the current working directory.
<code>prior</code>	A list containing the hyperprior values provided by the user. Each element of this list must be a vector of length two containing the user-provided hyperprior values and must be named with the name of the corresponding parameter. For example, the hyperprior values for the standard deviation parameter for the effects can be provided using the list <code>prior = list('sigma.prior.e' = c(0, 100))</code> . For more information about how to provide prior hypervalues for different types of parameters and models see details. If <code>prior</code> is set to <code>'default'</code> , the default values will be used.
<code>...</code>	Additional arguments that can be provided by the user. Examples are <code>d_e</code> and <code>d_c</code> , which should correspond to two binary indicator vectors with length equal to the number of rows of data. By default these variables are constructed within the function based on the observed data but it is possible for the user to directly provide them as a means to explore some Structural Not At Random (SNAR) mechanism assumptions about one or both outcomes. Individuals whose corresponding indicator value is set to 1 or 0 will be respectively associated with the structural or non-structural component in the model. Other optional arguments are <code>center = TRUE</code> , which centers all the covariates in the model or the additional arguments that can be provided to the function bcea to summarise the health economic evaluation results.

Details

Depending on the distributions specified for the outcome variables in the arguments `dist_e` and `dist_c` and the type of structural value mechanism specified in the argument `type`, different hurdle models are built and run in the background by the function `hurdle`. These are mixture models defined by two components: the first one is a mass distribution at the spike, while the second is a parametric model applied to the natural range of the relevant variable. Usually, a logistic regression is used to estimate the probability of incurring a "structural" value (e.g. 0 for the costs, or 1 for the effects); this is then used to weigh the mean of the "non-structural" values estimated in the second component. A simple example can be used to show how hurdle models are specified. Consider a data set comprising a response variable y and a set of centered covariate X_j . Specifically, for each subject in the trial $i = 1, \dots, n$ we define an indicator variable d_i taking value 1 if the i -th individual is associated with a structural value and 0 otherwise. This is modelled as:

$$d_i \text{ Bernoulli}(\pi_i)$$

$$\text{logit}(\pi_i) = \gamma_0 + \sum \gamma_j X_j$$

where

- π_i is the individual probability of a structural value in y .
- γ_0 represents the marginal probability of a structural value in y on the logit scale.
- γ_j represents the impact on the probability of a structural value in y of the centered covariates X_j .

When $\gamma_j = 0$, the model assumes a 'SCAR' mechanism, while when $\gamma_j \neq 0$ the mechanism is 'SAR'. For the parameters indexing the structural value model, the default prior distributions assumed are the following:

- γ_0 *Logistic*(0, 1)
- γ_j *Normal*(0, 0.01)

When user-defined hyperprior values are supplied via the argument `prior` in the function `hurdle`, the elements of this list (see Arguments) must be vectors of length 2 containing the user-provided hyperprior values and must take specific names according to the parameters they are associated with. Specifically, the names accepted by **missingHE** are the following:

- location parameters α_0, β_0 : "mean.prior.e"(effects) and/or "mean.prior.c"(costs)
- auxiliary parameters σ : "sigma.prior.e"(effects) and/or "sigma.prior.c"(costs)
- covariate parameters α_j, β_j : "alpha.prior"(effects) and/or "beta.prior"(costs)
- marginal probability of structural values γ_0 : "p.prior.e"(effects) and/or "p.prior.c"(costs)
- covariate parameters in the model of the structural values γ_j (if covariate data provided): "gamma.prior.e"(effects) and/or "gamma.prior.c"(costs)

For simplicity, here we have assumed that the set of covariates X_j used in the models for the effects/costs and in the model of the structural effect/cost values is the same. However, it is possible to specify different sets of covariates for each model using the arguments in the function `hurdle` (see Arguments).

Value

An object of the class 'missingHE' containing the following elements

data_set A list containing the original data set provided in `data` (see Arguments), the number of observed and missing individuals, the total number of individuals by treatment arm and the indicator vectors for the structural values

model_output A list containing the output of a JAGS model generated from the functions `jags`, and the posterior samples for the main parameters of the model and the imputed values

cea A list containing the output of the economic evaluation performed using the function `bcea`

type A character variable that indicate which type of structural value mechanism has been used to run the model, either SCAR or SAR (see details)

Author(s)

Andrea Gabrio

References

- Ntzoufras I. (2009). *Bayesian Modelling Using WinBUGS*, John Wiley and Sons.
- Daniels, MJ. Hogan, JW. (2008). *Missing Data in Longitudinal Studies: strategies for Bayesian modelling and sensitivity analysis*, CRC/Chapman Hall.
- Baio, G.(2012). *Bayesian Methods in Health Economics*. CRC/Chapman Hall, London.
- Gelman, A. Carlin, JB., Stern, HS. Rubin, DB.(2003). *Bayesian Data Analysis, 2nd edition*, CRC Press.
- Plummer, M. *JAGS: A program for analysis of Bayesian graphical models using Gibbs sampling*. (2003).

See Also

[jags](#), [bcea](#)

Examples

```
# Quick example to run using subset of MenSS dataset
MenSS.subset <- MenSS[50:100, ]

# Run the model using the hurdle function assuming a SCAR mechanism
# Use only 100 iterations to run a quick check
model.hurdle <- hurdle(data = MenSS.subset, model.eff = e ~ 1, model.cost = c ~ 1,
  model.se = se ~ 1, model.sc = sc ~ 1, se = 1, sc = 0, dist_e = "norm", dist_c = "norm",
  type = "SCAR", n.chains = 2, n.iter = 100, ppc = FALSE)

# Print the results of the JAGS model
print(model.hurdle)
#

# Use dic information criterion to assess model fit
pic.dic <- pic(model.hurdle, criterion = "dic", module = "total")
pic.dic
#

# Assess model convergence using graphical tools
# Produce histograms of the posterior samples for the mean effects
diag.hist <- diagnostic(model.hurdle, type = "histogram", param = "mu.e")
#

# Compare observed effect data with imputations from the model
# using plots (posterior means and credible intervals)
p1 <- plot(model.hurdle, class = "scatter", outcome = "effects")
#

# Summarise the CEA information from the model
summary(model.hurdle)
```

```

# Further examples which take longer to run
model.hurdle <- hurdle(data = MenSS, model.eff = e ~ u.0, model.cost = c ~ e,
  model.se = se ~ u.0, model.sc = sc ~ 1, se = 1, sc = 0, dist_e = "norm", dist_c = "norm",
  type = "SAR", n.chains = 2, n.iter = 500, ppc = FALSE)
#
# Print results for all imputed values
print(model.hurdle, value.mis = TRUE)

# Use looic to assess model fit
pic.looic <- pic(model.hurdle, criterion = "looic", module = "total")
pic.looic

# Show density plots for all parameters
diag.hist <- diagnostic(model.hurdle, type = "denplot", param = "all")

# Plots of imputations for all data
p1 <- plot(model.hurdle, class = "scatter", outcome = "all")

# Summarise the CEA results
summary(model.hurdle)

#
#

```

jagsresults

An internal function to summarise results from BUGS model

Description

This function hides missing data distribution from summary results of BUGS models

Usage

```

jagsresults(x, params, regex = FALSE, invert = FALSE,
  probs = c(0.025, 0.25, 0.5, 0.75, 0.975), signif, ...)

```

Arguments

x	The <code>rjags</code> , <code>rjags.parallel</code> , or <code>mcmc.list</code> object for which results will be printed.
params	Character vector or a regular expression pattern. The parameters for which results will be printed (unless <code>invert</code> is <code>FALSE</code> , in which case results for all parameters other than those given in <code>params</code> will be returned). If <code>regex</code> is <code>FALSE</code> , only those parameters that match <code>params</code> exactly will be returned. If <code>regex</code> is <code>TRUE</code> , <code>param</code> should be a character string giving the regular expression pattern to be matched.

regex	If regex is TRUE, then param is expected to be a single string giving a text pattern to be matched. Parameters with names matching the pattern will be returned (unless invert is TRUE, which results in all parameters that do not match the pattern being returned). Text pattern matching uses regular expressions (regex).
invert	Logical. If invert is TRUE, only those parameters that do not match elements of params will be returned.
probs	A numeric vector of probabilities within range [0, 1], representing the sample quantiles to be calculated and returned.
signif	If supplied, all columns other than n.eff will have their values rounded such that the most extreme value has the specified number of significant digits.
...	Additional arguments accepted by grep , e.g. perl=TRUE, to allow look-around pattern matching.

Examples

```
## Not run:
## Data
N <- 100
temp <- runif(N)
rain <- runif(N)
wind <- runif(N)
a <- 0.13
beta.temp <- 1.3
beta.rain <- 0.86
beta.wind <- -0.44
sd <- 0.16
y <- rnorm(N, a + beta.temp*temp + beta.rain*rain + beta.wind*wind, sd)
dat <- list(N=N, temp=temp, rain=rain, wind=wind, y=y)

### bugs example
library(R2jags)

## Model
M <- function() {
  for (i in 1:N) {
    y[i] ~ dnorm(y.hat[i], sd^-2)
    y.hat[i] <- a + beta.temp*temp[i] + beta.rain*rain[i] + beta.wind*wind[i]
    resid[i] <- y[i] - y.hat[i]
  }
  sd ~ dunif(0, 100)
  a ~ dnorm(0, 0.0001)
  beta.temp ~ dnorm(0, 0.0001)
  beta.rain ~ dnorm(0, 0.0001)
  beta.wind ~ dnorm(0, 0.0001)
}

## Fit model
jagsfit <- jags(dat, inits=NULL,
               parameters.to.save=c('a', 'beta.temp', 'beta.rain',
                                   'beta.wind', 'sd', 'resid'),
               model.file=M, n.iter=10000)
```

```

## Output
# model summary
jagsfit

# Results for beta.rain only
jagsresults(x=jagsfit, param='beta.rain')

# Results for 'a' and 'sd' only
jagsresults(x=jagsfit, param=c('a', 'sd'))
jagsresults(x=jagsfit, param=c('a', 'sd'),
            probs=c(0.01, 0.025, 0.1, 0.25, 0.5, 0.75, 0.9, 0.975))

# Results for all parameters including the string 'beta'
jagsresults(x=jagsfit, param='beta', regex=TRUE)

# Results for all parameters not including the string 'beta'
jagsresults(x=jagsfit, param='beta', regex=TRUE, invert=TRUE)

# Note that the above is NOT equivalent to the following, which returns all
# parameters that are not EXACTLY equal to 'beta'.
jagsresults(x=jagsfit, param='beta', invert=TRUE)

# Results for all parameters beginning with 'b' or including 'sd'.
jagsresults(x=jagsfit, param='^b|sd', regex=TRUE)

# Results for all parameters not beginning with 'beta'.
# This is equivalent to using param='^beta' with invert=TRUE and regex=TRUE
jagsresults(x=jagsfit, param='^(?!beta)', regex=TRUE, perl=TRUE)

## End(Not run)
#
#

```

Description

Data from a pilot RCT trial (The MenSS trial) on young men at risk of Sexually Transmitted Infections (STIs). A total of 159 individuals were enrolled in trial: 75 in the control (t=1) and 84 in the active intervention (t=2). Clinical and health economic outcome data were collected via self-reported questionnaires at four time points throughout the study: baseline, 3 months, 6 months and 12 months follow-up. Health economic data include utility scores related to quality of life and costs, from which QALYs and total costs were then computed using the area under the curve method and by summing up the cost components at each time point. Clinical data include the total number of instances of unprotected sex and whether the individual was associated with an STI diagnosis or not. Baseline data are available for the utilities (no baseline costs collected), instances of unprotected sex, sti diagnosis, age, ethnicity and employment variables.

Usage

```
data(MenSS)
```

Format

A data frame with 159 rows and 12 variables

Details

id id number

e Quality Adjusted Life Years (QALYs)

c Total costs in pounds

u.0 baseline utilities

age Age in years

ethnicity binary: white (1) and other (0)

employment binary: working (1) and other (0)

t Treatment arm indicator for the control (t=1) and the active intervention (t=2)

sex_inst.0 baseline number of instances of unprotected sex

sex_inst number of instances of unprotected sex at 12 months follow-up

sti.0 binary : baseline sti diagnosis (1) and no baseline sti diagnosis (0)

sti binary : sti diagnosis (1) and no sti diagnosis (0) at 12 months follow-up

References

Bailey et al. (2016) Health Technology Assessment 20 ([PubMed](#))

Examples

```
MenSS <- data(MenSS)
summary(MenSS)
str(MenSS)
```

pattern

*Full Bayesian Models to handle missingness in Economic Evaluations
(Pattern Mixture Models)*

Description

Full Bayesian cost-effectiveness models to handle missing data in the outcomes under different missingness mechanism assumptions, using alternative parametric distributions for the effect and cost variables and a pattern mixture approach to identify the model. The analysis is performed using the BUGS language, which is implemented in the software JAGS using the function [jags](#). The output is stored in an object of class 'missingHE'.

Usage

```
pattern(data, model.eff, model.cost, dist_e, dist_c, Delta_e, Delta_c,
        type, restriction = "CC", prob = c(0.05, 0.95), n.chains = 2,
        n.iter = 20000, n.burnin = floor(n.iter/2), inits = NULL,
        n.thin = 1, ppc = FALSE, save_model = FALSE, prior = "default",
        ...)
```

Arguments

<code>data</code>	A data frame in which to find the variables supplied in <code>model.eff</code> , <code>model.cost</code> (model formulas for effects and costs). Among these, effectiveness, cost and treatment indicator (only two arms) variables must always be provided and named 'e', 'c' and 't', respectively.
<code>model.eff</code>	A formula expression in conventional R linear modelling syntax. The response must be a health economic effectiveness outcome ('e') whose name must correspond to that used in data. Any covariates in the model must be provided on the right-hand side of the formula. If there are no covariates, 1 should be specified on the right hand side of the formula. By default, covariates are placed on the "location" parameter of the distribution through a linear model.
<code>model.cost</code>	A formula expression in conventional R linear modelling syntax. The response must be a health economic cost outcome ('c') whose name must correspond to that used in data. Any covariates in the model must be provided on the right-hand side of the formula. If there are no covariates, 1 should be specified on the right hand side of the formula. By default, covariates are placed on the "location" parameter of the distribution through a linear model. A joint bivariate distribution for effects and costs can be specified by including 'e' on the right-hand side of the formula for the costs model.
<code>dist_e</code>	Distribution assumed for the effects. Current available choices are: Normal ('norm'), Beta ('beta'), Gamma ('gamma'), Exponential ('exp'), Weibull ('weibull'), Logistic ('logis'), Poisson ('pois'), Negative Binomial ('nbinom') or Bernoulli ('bern').
<code>dist_c</code>	Distribution assumed for the costs. Current available choices are: Normal ('norm'), Gamma ('gamma') or LogNormal ('lnorm').
<code>Delta_e</code>	Range of values for the prior on the sensitivity parameters used to identify the mean of the effects under MNAR. The value must be set to 0 under MAR.
<code>Delta_c</code>	Range of values for the prior on the sensitivity parameters used to identify the mean of the costs under MNAR. The value must be set to 0 under MAR.
<code>type</code>	Type of missingness mechanism assumed. Choices are Missing At Random (MAR) and Missing Not At Random (MNAR).
<code>restriction</code>	type of identifying restriction to be imposed to identify the distributions of the missing data in each pattern. Available choices are: complete case restriction ('CC') - default - or available case restriction ('AC').
<code>prob</code>	A numeric vector of probabilities within the range (0,1), representing the upper and lower CI sample quantiles to be calculated and returned for the imputed values.

n.chains	Number of chains.
n.iter	Number of iterations.
n.burnin	Number of warmup iterations.
inits	A list with elements equal to the number of chains selected; each element of the list is itself a list of starting values for the JAGS model, or a function creating (possibly random) initial values. If inits is NULL, JAGS will generate initial values for all the model parameters.
n.thin	Thinning interval.
ppc	Logical. If ppc is TRUE, the estimates of the parameters that can be used to generate replications from the model are saved.
save_model	Logical. If save_model is TRUE a txt file containing the model code is printed in the current working directory.
prior	A list containing the hyperprior values provided by the user. Each element of this list must be a vector of length two containing the user-provided hyperprior values and must be named with the name of the corresponding parameter. For example, the hyperprior values for the standard deviation effect parameters can be provided using the list <code>prior = list('sigma.prior.e' = c(0, 100))</code> . For more information about how to provide prior hypervalues for different types of parameters and models see details. If prior is set to 'default', the default values will be used.
...	Additional arguments that can be provided by the user. Examples are <code>center = TRUE</code> to center all the covariates in the model or the additional arguments that can be provided to the function <code>bcea</code> to summarise the health economic evaluation results.

Details

Depending on the distributions specified for the outcome variables in the arguments `dist_e` and `dist_c` and the type of missingness mechanism specified in the argument `type`, different pattern mixture models are built and run in the background by the function `pattern`. The model for the outcomes is fitted in each missingness pattern and the parameters indexing the missing data distributions are identified using: the corresponding parameters identified from the observed data in other patterns (under 'MAR'); or a combination of the parameters identified by the observed data and some sensitivity parameters (under 'MNAR'). A simple example can be used to show how pattern mixture models are specified. Consider a data set comprising a response variable y and a set of centered covariate X_j . We denote with d_i the patterns' indicator variable for each subject in the trial $i = 1, \dots, n$ such that: $d_i = 1$ indicates the completers (both e and c observed), $d_i = 2$ and $d_i = 3$ indicate that only the costs or effects are observed, respectively, while $d_i = 4$ indicates that neither of the two outcomes is observed. In general, a different number of patterns can be observed between the treatment groups and `missingHE` accounts for this possibility by modelling a different patterns' indicator variables for each arm. For simplicity, in this example, we assume that the same number of patterns is observed in both groups. d_i is assigned a multinomial distribution, which probabilities are modelled using a Dirichlet prior (by default giving to each pattern the same weight). Next, the model specified in `dist_e` and `dist_c` is fitted in each pattern. The parameters that cannot be identified by the observed data in each pattern ($d = 2, 3, 4$), e.g. the means. $\mu_{e_c}[d]$ and $\mu_{c_c}[d]$, can be identified using the parameters estimated from other patterns. Two choices

are currently available: the complete cases ('CC') or available cases ('AC'). For example, using the 'CC' restriction, the parameters indexing the distributions of the missing data are identified as:

$$mu_e[2] = \mu_e[4] = \mu_e[1] + \Delta_e$$

$$mu_c[3] = \mu_c[4] = \mu_c[1] + \Delta_c$$

where

- $\mu_e[1]$ is the effects mean for the completers.
- $\mu_c[1]$ is the costs mean for the completers.
- Δ_e is the sensitivity parameters associated with the marginal effects mean.
- Δ_c is the sensitivity parameters associated with the marginal costs mean.

If the 'AC' restriction is chosen, only the parameters estimated from the observed data in pattern 2 (costs) and pattern 3 (effects) are used to identify those in the other patterns. When $\Delta_e = 0$ and $\Delta_c = 0$ the model assumes a 'MAR' mechanism. When $\Delta_e \neq 0$ and/or $\Delta_c \neq 0$ 'MNAR' departures for the effects and/or costs are explored assuming a Uniform prior distributions for the sensitivity parameters. The range of values for these priors is defined based on the boundaries specified in `Delta_e` and `Delta_c` (see Arguments), which must be provided by the user. When user-defined hyperprior values are supplied via the argument `prior` in the function `pattern`, the elements of this list (see Arguments) must be vectors of length two containing the user-provided hyperprior values and must take specific names according to the parameters they are associated with. Specifically, the names for the parameters indexing the model which are accepted by **missingHE** are the following:

- location parameters α_0 and β_0 : "mean.prior.e"(effects) and/or "mean.prior.c"(costs)
- auxiliary parameters σ : "sigma.prior.e"(effects) and/or "sigma.prior.c"(costs)
- covariate parameters α_j and β_j : "alpha.prior"(effects) and/or "beta.prior"(costs)

The only exception is the missingness patterns' probability π , denoted with "patterns.prior", whose hyperprior values must be provided as a list formed by two elements. These must be vectors of the same length equal to the number of patterns in the control (first element) and intervention (second element) group.

Value

An object of the class 'missingHE' containing the following elements

data_set A list containing the original data set provided in `data` (see Arguments), the number of observed and missing individuals, the total number of individuals by treatment arm and the indicator vectors for the missing values

model_output A list containing the output of a JAGS model generated from the functions `jags`, and the posterior samples for the main parameters of the model and the imputed values

cea A list containing the output of the economic evaluation performed using the function `bcea`

type A character variable that indicate which type of missingness assumption has been used to run the model, either MAR or MNAR (see details)

Author(s)

Andrea Gabrio

References

Daniels, MJ. Hogan, JW. *Missing Data in Longitudinal Studies: strategies for Bayesian modelling and sensitivity analysis*, CRC/Chapman Hall.

Baio, G.(2012). *Bayesian Methods in Health Economics*. CRC/Chapman Hall, London.

Gelman, A. Carlin, JB., Stern, HS. Rubin, DB.(2003). *Bayesian Data Analysis, 2nd edition*, CRC Press.

Plummer, M. *JAGS: A program for analysis of Bayesian graphical models using Gibbs sampling*. (2003).

See Also

[jags](#), [bcea](#)

Examples

```
# Quick example to run using subset of MenSS dataset
MenSS.subset <- MenSS[50:100, ]

# Run the model using the pattern function assuming a SCAR mechanism
# Use only 100 iterations to run a quick check
model.pattern <- pattern(data = MenSS.subset,model.eff = e~1,model.cost = c~1,
  dist_e = "norm", dist_c = "norm",type = "MAR", Delta_e = 0, Delta_c = 0,
  n.chains = 2, n.iter = 100, ppc = TRUE)

# Print the results of the JAGS model
print(model.pattern)
#

# Use dic information criterion and predictive checks to assess model fit
pic.dic <- pic(model.pattern, criterion = "dic", module = "total")
pic.dic
ppc(model.pattern, type = "histogram", ndisplay = 5)
#

# Assess model convergence using graphical tools
# Produce histograms of the posterior samples for the mean effects
diag.hist <- diagnostic(model.pattern, type = "histogram", param = "mu.e")
#

# Compare observed effect data with imputations from the model
# using plots (posterior means and credible intervals)
p1 <- plot(model.pattern, class = "scatter", outcome = "effects")
#
```

```

# Summarise the CEA information from the model
summary(model.pattern)

# Further examples which take longer to run
model.pattern <- pattern(data = MenSS, model.eff = e ~ u.0, model.cost = c ~ e,
  Delta_e = 0, Delta_c = 0, dist_e = "norm", dist_c = "norm",
  type = "MAR", n.chains = 2, n.iter = 500, ppc = FALSE)
#
# Print results for all imputed values
print(model.pattern, value.mis = TRUE)

# Use looic to assess model fit
pic.looic <- pic(model.pattern, criterion = "looic", module = "total")
pic.looic

# Show density plots for all parameters
diag.hist <- diagnostic(model.pattern, type = "denplot", param = "all")

# Plots of imputations for all data
p1 <- plot(model.pattern, class = "scatter", outcome = "all")

# Summarise the CEA results
summary(model.pattern)

#
#

```

pic *Predictive information criteria for Bayesian models fitted in JAGS using the function [selection](#), [pattern](#) or [hurdle](#)*

Description

Efficient approximate leave-one-out cross validation (LOO), deviance information criterion (DIC) and widely applicable information criterion (WAIC) for Bayesian models, calculated on the observed data.

Usage

```
pic(x, criterion = "dic", module = "total")
```

Arguments

x	A missingHE object containing the results of a Bayesian model fitted in cost-effectiveness analysis using the function selection , pattern or hurdle .
criterion	type of information criteria to be produced. Available choices are 'dic' for the Deviance Information Criterion, 'waic' for the Widely Applicable Information Criterion, and 'looic' for the Leave-One-Out Information Criterion.

module The modules with respect to which the information criteria should be computed. Available choices are 'total' for the whole model, 'e' for the effectiveness variables only, 'c' for the cost variables only, and 'both' for both outcome variables.

Details

The Deviance Information Criterion (DIC), Leave-One-Out Information Criterion (LOOIC) and the Widely Applicable Information Criterion (WAIC) are methods for estimating out-of-sample predictive accuracy from a Bayesian model using the log-likelihood evaluated at the posterior simulations of the parameters. DIC is computationally simple to calculate but it is known to have some problems, arising in part from it not being fully Bayesian in that it is based on a point estimate. LOOIC can be computationally expensive but can be easily approximated using importance weights that are smoothed by fitting a generalised Pareto distribution to the upper tail of the distribution of the importance weights. For more details about the methods used to compute LOOIC see the PSIS-LOO section in [loo-package](#). WAIC is fully Bayesian and closely approximates Bayesian cross-validation. Unlike DIC, WAIC is invariant to parameterisation and also works for singular models. In finite cases, WAIC and LOO give similar estimates, but for influential observations WAIC underestimates the effect of leaving out one observation.

Value

A named list containing different predictive information criteria results and quantities according to the value of `criterion`. In all cases, the measures are computed on the observed data for the specific modules of the model selected in `module`.

d_bar Posterior mean deviance (only if `criterion` is 'dic').

pD Effective number of parameters calculated with the formula used by JAGS (only if `criterion` is 'dic').

dic Deviance Information Criterion calculated with the formula used by JAGS (only if `criterion` is 'dic').

d_hat Deviance evaluated at the posterior mean of the parameters and calculated with the formula used by JAGS (only if `criterion` is 'dic')

elpd, elpd_se Expected log pointwise predictive density and standard error calculated on the observed data for the model nodes indicated in `module` (only if `criterion` is 'waic' or 'loo').

p, p_se Effective number of parameters and standard error calculated on the observed data for the model nodes indicated in `module` (only if `criterion` is 'waic' or 'loo').

looi, looi_se The leave-one-out information criterion and standard error calculated on the observed data for the model nodes indicated in `module` (only if `criterion` is 'loo').

waic, waic_se The widely applicable information criterion and standard error calculated on the observed data for the model nodes indicated in `module` (only if `criterion` is 'waic').

pointwise A matrix containing the pointwise contributions of each of the above measures calculated on the observed data for the model nodes indicated in `module` (only if `criterion` is 'waic' or 'loo').

pareto_k A vector containing the estimates of the shape parameter k for the generalised Pareto fit to the importance ratios for each leave-one-out distribution calculated on the observed data for the model nodes indicated in `module` (only if `criterion` is 'loo'). See [loo](#) for details about interpreting k .

Author(s)

Andrea Gabrio

References

Plummer, M. *JAGS: A program for analysis of Bayesian graphical models using Gibbs sampling*. (2003).

Vehtari, A. Gelman, A. Gabry, J. (2016a) Practical Bayesian model evaluation using leave-one-out cross-validation and WAIC. *Statistics and Computing*. Advance online publication.

Vehtari, A. Gelman, A. Gabry, J. (2016b) Pareto smoothed importance sampling. *ArXiv* preprint.

Gelman, A. Hwang, J. Vehtari, A. (2014) Understanding predictive information criteria for Bayesian models. *Statistics and Computing* 24, 997-1016.

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

See Also

[jags](#), [loo](#), [waic](#)

Examples

```
#For examples see the function selection, pattern or hurdle
#
#
```

plot.missingHE	<i>Plot method for the imputed data contained in the objects of class missingHE</i>
----------------	---

Description

Produces a plot of the observed and imputed values (with credible intervals) for the effect and cost outcomes from a Bayesian cost-effectiveness analysis model with two treatment arms, implemented using the function [selection](#), [pattern](#) or [hurdle](#). The graphical layout is obtained from the functions contained in the package **ggplot2** and **ggthemes**.

Usage

```
## S3 method for class 'missingHE'
plot(x, prob = c(0.05, 0.95), class = "scatter",
     outcome = "all", theme = NULL, ...)
```

Arguments

x	A missingHE object containing the results of the Bayesian model for cost-effectiveness analysis.
prob	A numeric vector of probabilities within range [0, 1], representing the upper and lower CI sample quantiles to be calculated and returned for the imputed values.
class	Type of the plot comparing the observed and imputed outcome data. Available choices are 'histogram' and 'scatter' for a histogram or a scatter plot of the observed and imputed outcome data, respectively.
outcome	The outcome variables that should be displayed. Options are: 'all' (default) which shows the plots for both treatment arms and types of outcome variables; 'effects' and 'costs' which show the plots for the corresponding outcome variables in both arms; 'arm1' and 'arm2' which show the plots by the selected treatment arm. To select the plots for a specific outcome in a specific treatment arm the options that can be used are 'effects_arm1', 'effects_arm2', 'costs_arm1' or 'costs_arm2'.
theme	Type of ggplot theme among some pre-defined themes, mostly taken from the package ggthemes . For a full list of available themes see details.
...	Additional parameters that can be provided to manage the output of plot.missingHE.

Details

The function produces a plot of the observed and imputed effect and cost data in a two-arm based cost-effectiveness model implemented using the function [selection](#), [pattern](#) or [hurdle](#). The purpose of this graph is to visually compare the outcome values for the fully-observed individuals with those imputed by the model for the missing individuals. For the scatter plot, imputed values are also associated with the credible intervals specified in the argument prob. The argument theme allows to customise the graphical aspect of the plots generated by plot.missingHE and allows to choose among a set of possible pre-defined themes taken from the package **ggtheme**. For a complete list of the available character names for each theme and scheme set, see **ggthemes** and **bayesplot**.

Value

A ggplot object containing the plots specified in the argument class.

Author(s)

Andrea Gabrio

References

- Daniels, MJ. Hogan, JW. (2008) *Missing Data in Longitudinal Studies: strategies for Bayesian modelling and sensitivity analysis*, CRC/Chapman Hall.
- Molenberghs, G. Fitzmaurice, G. Kenward, MG. Tsiatis, A. Verbeke, G. (2015) *Handbook of Missing Data Methodology*, CRC/Chapman Hall.

See Also

[selection](#) [pattern](#) [hurdle](#) [diagnostic](#)

Examples

```
#For examples see the function selection, pattern or hurdle
#
#
```

ppc	<i>Posterior predictive checks for assessing the fit to the observed data of Bayesian models implemented in JAGS using the function selection, pattern or hurdle</i>
-----	--

Description

The focus is restricted to full Bayesian models in cost-effectiveness analyses based on the function [selection](#), [pattern](#) and [hurdle](#), with the fit to the observed data being assessed through graphical checks based on the posterior replications generated from the model. Examples include the comparison of histograms, density plots, intervals, test statistics, evaluated using both the observed and replicated data. Different types of posterior predictive checks are implemented to assess model fit using functions contained in the package **bayesplot**. Graphics and plots are managed using functions contained in the package **ggplot2** and **ggthemes**.

Usage

```
ppc(x, type = "histogram", outcome = "effects_arm1", ndisplay = 15,
    theme = NULL, scheme_set = NULL, legend = TRUE, ...)
```

Arguments

x	An object of class "missingHE" containing the posterior results of a full Bayesian model implemented using the function selection , pattern or hurdle .
type	Type of posterior predictive check to be plotted for assessing model fit. Available choices include: 'histogram', 'boxplot', 'freqpoly', 'dens', 'dens_overlay' and 'ecdf_overlay', which compare the empirical and replicated distributions of the data; 'stat' and 'stat_2d', which compare the value of some statistics evaluated on the observed data with the replicated values for those statistics from the posterior predictions; 'error_hist', 'error_scatter', 'error_scatter_avg' and 'error_binned', which display the predictive errors of the model; 'intervals' and 'ribbon', which compare medians and central interval estimates of the replications with the observed data overlaid; 'scatter' and 'scatter_avg', which display scatterplots of the observed and replicated data.
outcome	The outcome variables that should be displayed. Only separate plots for a specific outcome in a specific treatment arm are available. These can be displayed using the names 'effects_arm1', 'effects_arm2', 'costs_arm1' or 'costs_arm2' for the effectiveness and cost outcomes in the control and treatment groups, respectively.
ndisplay	Number of posterior replications to be displayed in the plots.

theme	Type of ggplot theme among some pre-defined themes, mostly taken from the package ggthemes . For a full list of available themes see details.
scheme_set	Type of scheme sets among some pre-defined schemes, mostly taken from the package bayesplot . For a full list of available themes see details.
legend	Logical. If legend is TRUE, a legend is added to the plot.
...	Additional parameters that can be provided to manage the output of ppc. For more details see bayesplot .

Details

The function produces different types of graphical posterior predictive checks using the estimates from a Bayesian cost-effectiveness model implemented with the function [selection](#), [pattern](#) or [hurdle](#). The purpose of these checks is to visually compare the distribution (or some relevant quantity) of the observed data with respect to that from the replicated data for both effectiveness and cost outcomes in each treatment arm. Since predictive checks are meaningful only with respect to the observed data, only the observed outcome values are used to assess the fit of the model. The arguments `theme` and `scheme_set` allow to customise the graphical aspect of the plots generated by `ppc` and allow to choose among a set of possible pre-defined themes and scheme sets taken from the package **ggtheme** and **bayesplot**. For a complete list of the available character names for each theme and scheme set, see **ggthemes** and **bayesplot**.

Value

A ggplot object containing the plots specified in the argument `type`.

Author(s)

Andrea Gabrio

References

Gelman, A. Carlin, JB., Stern, HS. Rubin, DB.(2003). *Bayesian Data Analysis, 2nd edition*, CRC Press.

See Also

[selection](#) [pattern](#) [hurdle](#) [diagnostic](#)

Examples

```
#For examples see the function selection, pattern or hurdle
#
#
```

print.missingHE	<i>Print method for the posterior results contained in the objects of class missingHE</i>
-----------------	---

Description

Prints the summary table for the model fitted, with the estimate of the parameters and/or missing values.

Usage

```
## S3 method for class 'missingHE'
print(x, value.mis = FALSE, only.means = FALSE,
      ...)
```

Arguments

x	A missingHE object containing the results of the Bayesian model run using the function selection , pattern or hurdle .
value.mis	Logical. If value.mis is TRUE, the model results displayed contain also the imputed values, else if value.mis is FALSE the missing values are hidden.
only.means	Logical. If only.means is TRUE, then the print function only shows the summary statistics for the mean effectiveness and costs. Defaults at FALSE (in which case, shows the summary statistics for all parameters in the model).
...	additional arguments affecting the printed output produced. For example: digits= number of significant digits to be shown in the printed table (default=3). Not available if value.mis=TRUE.

Author(s)

Andrea Gabrio

Examples

```
# For examples see the function \link{selection},
# \link{pattern} or \link{hurdle}
#
#
```

prior_hurdle	<i>An internal function to change the hyperprior parameters in the hurdle model provided by the user depending on the type of structural value mechanism and outcome distributions assumed</i>
--------------	--

Description

This function modifies default hyper prior parameter values in the type of hurdle model selected according to the type of structural value mechanism and distributions for the outcomes assumed.

Usage

```
prior_hurdle(type, dist_e, dist_c, pe, pc, ze, zc, se, sc)
```

Arguments

type	Type of structural value mechanism assumed. Choices are Structural Completely At Random (SCAR), and Structural At Random (SAR). For a complete list of all available hyper parameters and types of models see the manual.
dist_e	distribution assumed for the effects. Current available choices are: Normal ('norm'), Beta ('beta'), Gamma ('gamma'), Exponential ('exp'), Weibull ('weibull'), Logistic ('logis'), Poisson ('pois'), Negative Binomial ('nbinom') or Bernoulli ('bern')
dist_c	distribution assumed for the costs. Current available choices are: Normal ('norm'), Gamma ('gamma') or LogNormal ('lnorm')
pe	Number of covariates for the effectiveness model
pc	Number of covariates for the cost model
ze	Number of covariates or the structural indicators model for the effectiveness
zc	Number of covariates or the structural indicators model for the costs
se	Structural value for the effectiveness
sc	Structural value for the costs

Examples

```
#Internal function only
#no examples
#
#
```

prior_pattern	<i>An internal function to change the hyperprior parameters in the selection model provided by the user depending on the type of missingness mechanism and outcome distributions assumed</i>
---------------	--

Description

This function modifies default hyper prior parameter values in the type of selection model selected according to the type of missingness mechanism and distributions for the outcomes assumed.

Usage

```
prior_pattern(type, dist_e, dist_c, pe, pc, d_list, restriction)
```

Arguments

type	Type of missingness mechanism assumed. Choices are Missing At Random (MAR), Missing Not At Random for the effects (MNAR_eff), Missing Not At Random for the costs (MNAR_cost), and Missing Not At Random for both (MNAR). For a complete list of all available hyper parameters and types of models see the manual.
dist_e	distribution assumed for the effects. Current available choices are: Normal ('norm'), Beta ('beta'), Gamma ('gamma'), Exponential ('exp'), Weibull ('weibull'), Logistic ('logis'), Poisson ('pois'), Negative Binomial ('nbinom') or Bernoulli ('bern')
dist_c	Distribution assumed for the costs. Current available choices are: Normal ('norm'), Gamma ('gamma') or LogNormal ('lnorm')
pe	Number of covariates for the effectiveness model
pc	Number of covariates for the cost model
d_list	a list of the number and types of patterns in the data
restriction	type of identifying restriction to be imposed

Examples

```
#Internal function only
#no examples
#
#
```

prior_selection	<i>An internal function to change the hyperprior parameters in the selection model provided by the user depending on the type of missingness mechanism and outcome distributions assumed</i>
-----------------	--

Description

This function modifies default hyper prior parameter values in the type of selection model selected according to the type of missingness mechanism and distributions for the outcomes assumed.

Usage

```
prior_selection(type, dist_e, dist_c, pe, pc, ze, zc)
```

Arguments

type	Type of missingness mechanism assumed. Choices are Missing At Random (MAR), Missing Not At Random for the effects (MNAR_eff), Missing Not At Random for the costs (MNAR_cost), and Missing Not At Random for both (MNAR). For a complete list of all available hyper parameters and types of models see the manual.
dist_e	distribution assumed for the effects. Current available choices are: Normal ('norm'), Beta ('beta'), Gamma ('gamma'), Exponential ('exp'), Weibull ('weibull'), Logistic ('logis'), Poisson ('pois'), Negative Binomial ('nbinom') or Bernoulli ('bern')
dist_c	Distribution assumed for the costs. Current available choices are: Normal ('norm'), Gamma ('gamma') or LogNormal ('lnorm')
pe	Number of covariates for the effectiveness model
pc	Number of covariates for the cost model
ze	Number of covariates or the missingness indicators model for the effectiveness
zc	Number of covariates or the missingness indicators model for the costs

Examples

```
#Internal function only
#no examples
#
#
```

run_hurdle	<i>An internal function to execute a JAGS hurdle model and get posterior results</i>
------------	--

Description

This function fits a JAGS using the `jags` function and obtain posterior inferences.

Usage

```
run_hurdle(type, dist_e, dist_c, inits, se, sc, sde, sdc, ppc)
```

Arguments

<code>type</code>	Type of structural value mechanism assumed. Choices are Structural Completely At Random (SCAR), and Structural At Random (SAR).
<code>dist_e</code>	distribution assumed for the effects. Current available choices are: Normal ('norm'), Beta ('beta'), Gamma ('gamma'), Exponential ('exp'), Weibull ('weibull'), Logistic ('logis'), Poisson ('pois'), Negative Binomial ('nbinom') or Bernoulli ('bern')
<code>dist_c</code>	distribution assumed for the costs. Current available choices are: Normal ('norm'), Gamma ('gamma') or LogNormal ('lnorm').
<code>inits</code>	a list with elements equal to the number of chains selected; each element of the list is itself a list of starting values for the BUGS model, or a function creating (possibly random) initial values. If <code>inits</code> is NULL, JAGS will generate initial values for parameters
<code>se</code>	Structural value to be found in the effect data. If set to NULL, no structural value is chosen and a standard model for the effects is run.
<code>sc</code>	Structural value to be found in the cost data. If set to NULL, no structural value is chosen and a standard model for the costs is run.
<code>sde</code>	hyper-prior value for the standard deviation of the distribution of the structural effects. The default value is $1.0E-6$ to approximate a point mass at the structural value provided by the user.
<code>sdc</code>	hyper-prior value for the standard deviation of the distribution of the structural costs. The default value is $1.0E-6$ to approximate a point mass at the structural value provided by the user.
<code>ppc</code>	Logical. If <code>ppc</code> is TRUE, the estimates of the parameters that can be used to generate replications from the model are saved.

Examples

```
#Internal function only
#No examples
#
#
```

run_pattern	<i>An internal function to execute a JAGS pattern mixture model and get posterior results</i>
-------------	---

Description

This function fits a JAGS using the `jags` function and obtain posterior inferences.

Usage

```
run_pattern(type, dist_e, dist_c, inits, d_list, d1, d2, restriction, ppc)
```

Arguments

type	Type of missingness mechanism assumed. Choices are Missing At Random (MAR), Missing Not At Random for the effects (MNAR_eff), Missing Not At Random for the costs (MNAR_cost), and Missing Not At Random for both (MNAR).
dist_e	distribution assumed for the effects. Current available choices are: Normal ('norm'), Beta ('beta'), Gamma ('gamma'), Exponential ('exp'), Weibull ('weibull'), Logistic ('logis'), Poisson ('pois'), Negative Binomial ('nbinom') or Bernoulli ('bern').
dist_c	Distribution assumed for the costs. Current available choices are: Normal ('norm'), Gamma ('gamma') or LogNormal ('lnorm').
inits	a list with elements equal to the number of chains selected; each element of the list is itself a list of starting values for the BUGS model, or a function creating (possibly random) initial values. If inits is NULL, JAGS will generate initial values for parameters.
d_list	a list of the number and types of patterns in the data.
d1	Patterns in the control.
d2	Patterns in the intervention.
restriction	type of identifying restriction to be imposed.
ppc	Logical. If ppc is TRUE, the estimates of the parameters that can be used to generate replications from the model are saved.

Examples

```
#Internal function only
#No examples
#
#
```

run_selection	<i>An internal function to execute a JAGS selection model and get posterior results</i>
---------------	---

Description

This function fits a JAGS using the `jags` function and obtain posterior inferences.

Usage

```
run_selection(type, dist_e, dist_c, inits, ppc)
```

Arguments

<code>type</code>	Type of missingness mechanism assumed. Choices are Missing At Random (MAR), Missing Not At Random for the effects (MNAR_eff), Missing Not At Random for the costs (MNAR_cost), and Missing Not At Random for both (MNAR).
<code>dist_e</code>	distribution assumed for the effects. Current available choices are: Normal ('norm'), Beta ('beta'), Gamma ('gamma'), Exponential ('exp'), Weibull ('weibull'), Logistic ('logis'), Poisson ('pois'), Negative Binomial ('nbinom') or Bernoulli ('bern').
<code>dist_c</code>	Distribution assumed for the costs. Current available choices are: Normal ('norm'), Gamma ('gamma') or LogNormal ('lnorm').
<code>inits</code>	a list with elements equal to the number of chains selected; each element of the list is itself a list of starting values for the BUGS model, or a function creating (possibly random) initial values. If inits is NULL, JAGS will generate initial values for parameters.
<code>ppc</code>	Logical. If ppc is TRUE, the estimates of the parameters that can be used to generate replications from the model are saved.

Examples

```
#Internal function only
#No examples
#
#
```

 selection

*Full Bayesian Models to handle missingness in Economic Evaluations
(Selection Models)*

Description

Full Bayesian cost-effectiveness models to handle missing data in the outcomes under different missing data mechanism assumptions, using alternative parametric distributions for the effect and cost variables and using a selection model approach to identify the model. The analysis is performed using the BUGS language, which is implemented in the software JAGS using the function `jags`. The output is stored in an object of class 'missingHE'.

Usage

```
selection(data, model.eff, model.cost, model.me = me ~ 1, model.mc = mc
  ~ 1, dist_e, dist_c, type, prob = c(0.05, 0.95), n.chains = 2,
  n.iter = 20000, n.burnin = floor(n.iter/2), inits = NULL,
  n.thin = 1, ppc = FALSE, save_model = FALSE, prior = "default",
  ...)
```

Arguments

<code>data</code>	A data frame in which to find the variables supplied in <code>model.eff</code> , <code>model.cost</code> (model formulas for effects and costs) and <code>model.me</code> , <code>model.mc</code> (model formulas for the missing effect and cost models). Among these, effectiveness, cost and treatment indicator (only two arms) variables must always be provided and named 'e', 'c' and 't', respectively.
<code>model.eff</code>	A formula expression in conventional R linear modelling syntax. The response must be a health economic effectiveness outcome ('e') whose name must correspond to that used in <code>data</code> . Any covariates in the model must be provided on the right-hand side of the formula. If there are no covariates, 1 should be specified on the right hand side of the formula. By default, covariates are placed on the "location" parameter of the distribution through a linear model.
<code>model.cost</code>	A formula expression in conventional R linear modelling syntax. The response must be a health economic cost outcome ('c') whose name must correspond to that used in <code>data</code> . and any covariates are given on the right-hand side. Any covariates in the model must be provided on the right-hand side of the formula. If there are no covariates, 1 should be specified on the right hand side of the formula. By default, covariates are placed on the "location" parameter of the distribution through a linear model. A joint bivariate distribution for effects and costs can be specified by including 'e' on the right-hand side of the formula for the costs model.
<code>model.me</code>	A formula expression in conventional R linear modelling syntax. The response must be indicated with the term 'me'(missing effects) and any covariates must be provided on the right-hand side of the formula. If there are no covariates, 1

should be specified on the right hand side of the formula. By default, covariates are placed on the "probability" parameter for the missing effects through a logistic-linear model.

model.mc	A formula expression in conventional R linear modelling syntax. The response must be indicated with the term 'mc'(missing costs) and any covariates must be provided on the right-hand side of the formula. If there are no covariates, 1 should be specified on the right hand side of the formula. By default, covariates are placed on the "probability" parameter for the missing costs through a logistic-linear model.
dist_e	Distribution assumed for the effects. Current available choices are: Normal ('norm'), Beta ('beta'), Gamma ('gamma'), Exponential ('exp'), Weibull ('weibull'), Logistic ('logis'), Poisson ('pois'), Negative Binomial ('nbinom') or Bernoulli ('bern').
dist_c	Distribution assumed for the costs. Current available choices are: Normal ('norm'), Gamma ('gamma') or LogNormal ('lnorm').
type	Type of missingness mechanism assumed. Choices are Missing At Random (MAR) and Missing Not At Random (MNAR).
prob	A numeric vector of probabilities within the range (0,1), representing the upper and lower CI sample quantiles to be calculated and returned for the imputed values.
n.chains	Number of chains.
n.iter	Number of iterations.
n.burnin	Number of warmup iterations.
inits	A list with elements equal to the number of chains selected; each element of the list is itself a list of starting values for the JAGS model, or a function creating (possibly random) initial values. If inits is NULL, JAGS will generate initial values for all the model parameters.
n.thin	Thinning interval.
ppc	Logical. If ppc is TRUE, the estimates of the parameters that can be used to generate replications from the model are saved.
save_model	Logical. If save_model is TRUE, a txt file containing the model code is printed in the current working directory.
prior	A list containing the hyperprior values provided by the user. Each element of this list must be a vector of length two containing the user-provided hyperprior values and must be named with the name of the corresponding parameter. For example, the hyperprior values for the standard deviation effect parameters can be provided using the list <code>prior = list('sigma.prior.e' = c(0, 100))</code> . For more information about how to provide prior hypervalues for different types of parameters and models see details. If prior is set to 'default', the default values will be used.
...	Additional arguments that can be provided by the user. Examples are <code>center = TRUE</code> to center all the covariates in the model or the additional arguments that can be provided to the function <code>bcea</code> to summarise the health economic evaluation results.

Details

Depending on the distributions specified for the outcome variables in the arguments `dist_e` and `dist_c` and the type of missingness mechanism specified in the argument `type`, different selection models are built and run in the background by the function `selection`. These models consist in logistic regressions that are used to estimate the probability of missingness in one or both the outcomes. A simple example can be used to show how selection models are specified. Consider a data set comprising a response variable y and a set of centered covariate X_j . For each subject in the trial $i = 1, \dots, n$ we define an indicator variable m_i taking value 1 if the i -th individual is associated with a missing value and 0 otherwise. This is modelled as:

$$m_i \sim \text{Bernoulli}(\pi_i)$$

$$\text{logit}(\pi_i) = \gamma_0 + \sum \gamma_j X_j + \delta(y)$$

where

- π_i is the individual probability of a missing value in y
- γ_0 represents the marginal probability of a missing value in y on the logit scale.
- γ_j represents the impact on the probability of a missing value in y of the centered covariates X_j .
- δ represents the impact on the probability of a missing value in y of the missing value itself.

When $\delta = 0$ the model assumes a 'MAR' mechanism, while when $\delta \neq 0$ the mechanism is 'MNAR'. For the parameters indexing the missingness model, the default prior distributions assumed are the following:

- $\gamma_0 \sim \text{Logistic}(0, 1)$
- $\gamma_j \sim \text{Normal}(0, 0.01)$
- $\delta \sim \text{Normal}(0, 1)$

When user-defined hyperprior values are supplied via the argument `prior` in the function `selection`, the elements of this list (see Arguments) must be vectors of length two containing the user-provided hyperprior values and must take specific names according to the parameters they are associated with. Specifically, the names for the parameters indexing the model which are accepted by **missingHE** are the following:

- location parameters α_0 and β_0 : "mean.prior.e"(effects) and/or "mean.prior.c"(costs)
- auxiliary parameters σ : "sigma.prior.e"(effects) and/or "sigma.prior.c"(costs)
- covariate parameters α_j and β_j : "alpha.prior"(effects) and/or "beta.prior"(costs)
- marginal probability of missing values γ_0 : "p.prior.e"(effects) and/or "p.prior.c"(costs)
- covariate parameters in the missingness model γ_j (if covariate data provided): "gamma.prior.e"(effects) and/or "gamma.prior.c"(costs)
- mnar parameter δ : "delta.prior.e"(effects) and/or "delta.prior.c"(costs)

For simplicity, here we have assumed that the set of covariates X_j used in the models for the effects/costs and in the model of the missing effect/cost values is the same. However, it is possible to specify different sets of covariates for each model using the arguments in the function `selection` (see Arguments).

Value

An object of the class 'missingHE' containing the following elements

data_set A list containing the original data set provided in data (see Arguments), the number of observed and missing individuals, the total number of individuals by treatment arm and the indicator vectors for the missing values

model_output A list containing the output of a JAGS model generated from the functions [jags](#), and the posterior samples for the main parameters of the model and the imputed values

cea A list containing the output of the economic evaluation performed using the function [bcea](#)

type A character variable that indicate which type of missingness mechanism has been used to run the model, either MAR or MNAR (see details)

Author(s)

Andrea Gabrio

References

Daniels, MJ. Hogan, JW. *Missing Data in Longitudinal Studies: strategies for Bayesian modelling and sensitivity analysis*, CRC/Chapman Hall.

Baio, G.(2012). *Bayesian Methods in Health Economics*. CRC/Chapman Hall, London.

Gelman, A. Carlin, JB., Stern, HS. Rubin, DB.(2003). *Bayesian Data Analysis, 2nd edition*, CRC Press.

Plummer, M. *JAGS: A program for analysis of Bayesian graphical models using Gibbs sampling*. (2003).

See Also

[jags](#), [bcea](#)

Examples

```
# Quick example to run using subset of MenSS dataset
MenSS.subset <- MenSS[50:100, ]

# Run the model using the selection function assuming a SCAR mechanism
# Use only 100 iterations to run a quick check
model.selection <- selection(data = MenSS.subset, model.eff = e ~ 1, model.cost = c ~ 1,
  model.me = me ~ 1, model.mc = mc ~ 1, dist_e = "norm", dist_c = "norm",
  type = "MAR", n.chains = 2, n.iter = 100, ppc = TRUE)

# Print the results of the JAGS model
print(model.selection)
#

# Use dic information criterion and predictive checks to assess model fit
pic.dic <- pic(model.selection, criterion = "dic", module = "total")
pic.dic
ppc(model.selection, type = "histogram", ndisplay = 5)
```

```

#

# Assess model convergence using graphical tools
# Produce histograms of the posterior samples for the mean effects
diag.hist <- diagnostic(model.selection, type = "histogram", param = "mu.e")
#

# Compare observed effect data with imputations from the model
# using plots (posterior means and credible intervals)
p1 <- plot(model.selection, class = "scatter", outcome = "effects")
#

# Summarise the CEA information from the model
summary(model.selection)

# Further examples which take longer to run
model.selection <- selection(data = MenSS, model.eff = e ~ u.0, model.cost = c ~ e,
  model.se = me ~ u.0, model.mc = mc ~ 1, dist_e = "norm", dist_c = "norm",
  type = "MAR", n.chains = 2, n.iter = 500, ppc = FALSE)
#
# Print results for all imputed values
print(model.selection, value.mis = TRUE)

# Use looic to assess model fit
pic.looic <- pic(model.selection, criterion = "looic", module = "total")
pic.looic

# Show density plots for all parameters
diag.hist <- diagnostic(model.selection, type = "denplot", param = "all")

# Plots of imputations for all data
p1 <- plot(model.selection, class = "scatter", outcome = "all")

# Summarise the CEA results
summary(model.selection)

#
#

```

summary.missingHE

Summary method for objects in the class missingHE

Description

Produces a table printout with some summary results of the health economic evaluation probabilistic model run using the function [selection](#), [pattern](#) or [hurdle](#).

Usage

```
## S3 method for class 'missingHE'
summary(object, ...)
```

Arguments

object A missingHE object containing the results of the Bayesian modelling and the economic evaluation

... Additional arguments affecting the summary produced.

Value

Prints a table with some information on the health economic model based on the assumption selected for the missingness using the function selection, pattern or hurdle. Summary information on the main parameters of interests is provided.

Author(s)

Andrea Gabrio

References

Baio, G.(2012). *Bayesian Methods in Health Economics*. CRC/Chapman Hall, London.

See Also

[selection pattern hurdle diagnostic plot.missingHE](#)

Examples

```
#For examples see the function selection, pattern or hurdle
#
#
```

write_hurdle	<i>An internal function to select which type of hurdle model to execute for both effectiveness and costs. Alternatives vary depending on the type of distribution assumed for the effect and cost variables, type of structural value mechanism assumed and independence or joint modelling This function selects which type of model to execute.</i>
--------------	---

Description

An internal function to select which type of hurdle model to execute for both effectiveness and costs. Alternatives vary depending on the type of distribution assumed for the effect and cost variables, type of structural value mechanism assumed and independence or joint modelling This function selects which type of model to execute.

Usage

```
write_hurdle(dist_e, dist_c, ind, type, pe, pc, ze, zc, se, sc)
```

Arguments

dist_e	Distribution assumed for the effects. Current available choices are: Normal ('norm'), Beta ('beta'), Gamma ('gamma'), Exponential ('exp'), Weibull ('weibull'), Logistic ('logis'), Poisson ('pois'), Negative Binomial ('nbinom') or Bernoulli ('bern')
dist_c	Distribution assumed for the costs. Current available choices are: Normal ('norm'), Gamma ('gamma') or LogNormal ('lnorm')
ind	Logical; if TRUE independence between effectiveness and costs is assumed, else correlation is accounted for
type	Type of structural value mechanism assumed. Choices are Structural Completely At Random (SCAR) and Structural At Random (SAR)
pe	Number of covariates for the effectiveness model
pc	Number of covariates for the cost model
ze	Number of covariates or the structural indicators model for the effectiveness
zc	Number of covariates or the structural indicators model for the costs
se	Structural value for the effectiveness
sc	Structural value for the costs

Examples

```
#Internal function only
#No examples
#
#
```

write_pattern	<i>An internal function to select which type of pattern mixture model to execute. Alternatives vary depending on the type of distribution assumed for the effect and cost variables, type of missingness mechanism assumed and independence or joint modelling This function selects which type of model to execute.</i>
---------------	--

Description

An internal function to select which type of pattern mixture model to execute. Alternatives vary depending on the type of distribution assumed for the effect and cost variables, type of missingness mechanism assumed and independence or joint modelling This function selects which type of model to execute.

Usage

```
write_pattern(dist_e, dist_c, ind, type, pe, pc, d_list, d1, d2,
             restriction)
```

Arguments

dist_e	Distribution assumed for the effects. Current available choices are: Normal ('norm'), Beta ('beta'), Gamma ('gamma'), Exponential ('exp'), Weibull ('weibull'), Logistic ('logis'), Poisson ('pois'), Negative Binomial ('nbinom') or Bernoulli ('bern')
dist_c	Distribution assumed for the costs. Current available choices are: Normal ('norm'), Gamma ('gamma') or LogNormal ('lnorm')
ind	Logical; if TRUE independence between effectiveness and costs is assumed, else correlation is accounted for
type	Type of missingness mechanism assumed. Choices are Missing At Random (MAR), Missing Not At Random for the effects (MNAR_eff), Missing Not At Random for the costs (MNAR_cost), and Missing Not At Random for both (MNAR)
pe	Number of covariates for the effectiveness model
pc	Number of covariates for the cost model
d_list	Number and type of patterns
d1	Pattern indicator in the control
d2	Pattern indicator in the intervention
restriction	type of identifying restriction to be imposed

Examples

```
# Internal function only
# No examples
#
#
```

write_selection	<i>An internal function to select which type of selection model to execute. Alternatives vary depending on the type of distribution assumed for the effect and cost variables, type of missingness mechanism assumed and independence or joint modelling This function selects which type of model to execute.</i>
-----------------	--

Description

An internal function to select which type of selection model to execute. Alternatives vary depending on the type of distribution assumed for the effect and cost variables, type of missingness mechanism assumed and independence or joint modelling This function selects which type of model to execute.

Usage

```
write_selection(dist_e, dist_c, ind, type, pe, pc, ze, zc)
```

Arguments

dist_e	Distribution assumed for the effects. Current available choices are: Normal ('norm'), Beta ('beta'), Gamma ('gamma'), Exponential ('exp'), Weibull ('weibull'), Logistic ('logis'), Poisson ('pois'), Negative Binomial ('nbinom') or Bernoulli ('bern')
dist_c	Distribution assumed for the costs. Current available choices are: Normal ('norm'), Gamma ('gamma') or LogNormal ('lnorm')
ind	Logical; if TRUE independence between effectiveness and costs is assumed, else correlation is accounted for
type	Type of missingness mechanism assumed. Choices are Missing At Random (MAR), Missing Not At Random for the effects (MNAR_eff), Missing Not At Random for the costs (MNAR_cost), and Missing Not At Random for both (MNAR)
pe	Number of covariates for the effectiveness model
pc	Number of covariates for the cost model
ze	Number of covariates or the missingness indicators model for the effectiveness
zc	Number of covariates or the missingness indicators model for the costs

Examples

```
#Internal function only  
#No examples  
#  
#
```

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