Package 'gemtc'

June 24, 2020

```
Version 0.8-6
Date 2020-06-23
Title Network Meta-Analysis Using Bayesian Methods
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Description Network meta-analyses (mixed treatment comparisons) in the Bayesian
      framework using JAGS. Includes methods to assess heterogeneity and
      inconsistency, and a number of standard visualizations.
      van Valkenhoef et al. (2012) <doi:10.1002/jrsm.1054>;
      van Valkenhoef et al. (2015) <doi:10.1002/jrsm.1167>.
Depends coda (>= 0.13)
Imports igraph (>= 1.0), meta (>= 2.1), plyr (>= 1.8), graphics,
      grDevices, stats, utils, grid, rjags (>= 4-0), truncnorm,
      Rglpk, forcats (\geq 0.5.0)
Suggests testthat (>= 0.8), Matrix, XML (>= 3.6)
URL http://github.com/gertvv/gemtc
License GPL-3
LazvData true
Collate 'anohe.R' 'arrayize.R' 'blobbogram.R' 'template.R' 'code.R'
      'data.R' 'deviance.R' 'forest.R' 'solveLP.R' 'inits.R'
      'likelihoods.R' 'll-helper.counts.R' 'll.binom.cloglog.R'
      'll.binom.log.R' 'll.binom.logit.R' 'll.call.R'
      'll.normal.identity.R' 'll.poisson.log.R'
      'minimum.diameter.spanning.tree.R' 'mtc.data.studyrow.R'
      'mtc.hy.prior.R' 'mtc.model.R' 'mtc.model.consistency.R'
      'mtc.model.nodesplit.R' 'mtc.model.regression.R'
      'mtc.model.ume.R' 'mtc.model.use.R' 'mtc.network.R'
      'mtc.network.xml.R' 'stopIfNotConsistent.R' 'mtc.result.R'
      'mtc.run.R' 'nodesplit.R' 'plotCovariateEffect.R' 'priors.R'
      'rank.probability.R' 'regression.R' 'relative.effect.R'
      'relative.effect.table.R'
```

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NeedsCompilation yes

Repository CRAN

Date/Publication 2020-06-24 08:40:20 UTC

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gemt	c-package	GeMTC:	Network	k meta-a	ınalysis ir	n R	

Description

An R package for performing network meta-analyses (mixed treatment comparisons).

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Details

Network meta-analysis, or mixed treatment comparison (MTC) is a technique to meta-analyze networks of trials comparing two or more treatments at the same time [Dias et al. 2013a]. Using a Bayesian hierarchical model, all direct and indirect comparisons are taken into account to arrive at a single consistent estimate of the effect of all included treatments based on all included studies.

This package allows the automated generation of network meta-analysis models [van Valkenhoef et al. 2012], inclusing both fixed effect and random effects network meta-analysis, node-splitting models to identify inconsistency, and network meta-regression models. Models are estimated using JAGS (through the rjags package).

The package can read GeMTC files that were created by the GeMTC GUI (now obsolete) or exported from ADDIS 1.x, if the optional package XML is installed. The source for GeMTC is available under the GPL-3 on Github.

Author(s)

Gert van Valkenhoef

References

- S. Dias, N.J. Welton, D.M. Caldwell, and A.E. Ades (2010), *Checking consistency in mixed treat-ment comparison meta-analysis*, Statistics in Medicine 29(7-8, Sp. Iss. SI):932-944. [doi:10.1002/sim.3767]
- S. Dias, A.J. Sutton, A.E. Ades, and N.J. Welton (2013a), *A Generalized Linear Modeling Framework for Pairwise and Network Meta-analysis of Randomized Controlled Trials*, Medical Decision Making 33(5):607-617. [doi:10.1177/0272989X12458724]
- S. Dias, A.J. Sutton, N.J. Welton, and A.E. Ades (2013b), *Heterogeneity Subgroups, Meta-Regression, Bias, and Bias-Adjustment*, Medical Decision Making 33(5):618-640. [doi:10.1177/0272989X13485157]
- S. Dias, N.J. Welton, A.J. Sutton, D.M. Caldwell, G. Lu, and A.E. Ades (2013c), *Inconsistency in Networks of Evidence Based on Randomized Controlled Trials*, Medical Decision Making 33(5):641-656. [doi:10.1177/0272989X12455847]
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- R.M. Turner, J. Davey, M.J. Clarke, S.G. Thompson, J.P.T. Higgins (2012), *Predicting the extent of heterogeneity in meta-analysis, using empirical data from the Cochrane Database of Systematic Reviews*, International Journal of Epidemiology 41(3):818-827. [doi:10.1093/ije/dys041]
- G. van Valkenhoef, G. Lu, B. de Brock, H. Hillege, A.E. Ades, and N.J. Welton (2012), *Automating network meta-analysis*, Research Synthesis Methods 3(4):285-299. [doi:10.1002/jrsm.1054]
- G. van Valkenhoef, S. Dias, A.E. Ades, and N.J. Welton (2015), *Automated generation of node-splitting models for assessment of inconsistency in network meta-analysis*, Research Synthesis Methods, accepted manuscript. [doi:10.1002/jrsm.1167]
- G. van Valkenhoef et al. (draft), *Modeling inconsistency as heterogeneity in network meta-analysis*, draft manuscript.

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D.E. Warn, S.G. Thompson, and D.J. Spiegelhalter (2002), *Bayesian random effects meta-analysis of trials with binary outcomes: methods for the absolute risk difference and relative risk scales*, Statistics in Medicine 21(11):1601-1623. [doi:10.1002/sim.1189]

See Also

```
mtc.network, mtc.model, mtc.run
```

Examples

```
# Load the example network and generate a consistency model:
model <- mtc.model(smoking, type="consistency")</pre>
# Load pre-generated samples instead of runing the model:
## Not run: results <- mtc.run(model, thin=10)</pre>
results <- dget(system.file("extdata/luades-smoking.samples.gz", package="gemtc"))
# Print a basic statistical summary of the results:
summary(results)
## Iterations = 5010:25000
## Thinning interval = 10
## Number of chains = 4
## Sample size per chain = 2000
##
## 1. Empirical mean and standard deviation for each variable,
      plus standard error of the mean:
##
##
##
           Mean
                    SD Naive SE Time-series SE
## d.A.B 0.4965 0.4081 0.004563
                                       0.004989
## d.A.C 0.8359 0.2433 0.002720
                                       0.003147
## d.A.D 1.1088 0.4355 0.004869
                                       0.005280
## sd.d 0.8465 0.1913 0.002139
                                       0.002965
## 2. Quantiles for each variable:
##
            2.5%
                    25%
                           50%
                                  75% 97.5%
## d.A.B -0.2985 0.2312 0.4910 0.7530 1.341
## d.A.C 0.3878 0.6720 0.8273 0.9867 1.353
## d.A.D 0.2692 0.8197 1.0983 1.3824 2.006
## sd.d 0.5509 0.7119 0.8180 0.9542 1.283
```

atrialFibrillation

Prevention of stroke in atrial fibrillation patients

Description

A dataset of 25 trials investigating 17 treatments for stroke prevention in atrial fibrillation patients. The main outcome is the number of patients with a stroke, and a covariate captures the proportion of patients with a prior stroke.

blobbogram 5

Data are take from Table 1 of Cooper et al. (2009), with the following corrections applied: SPAF 3 and AFASAK 2 do not have a treatment 13 arm, and SPAF 1 does not contain treatment 5, but treatment 6. Thanks to prof. Cooper for providing the original analysis dataset.

Format

A network meta-regression dataset containing 60 rows of arm-based data (responders and sample size).

Source

Cooper et al. (2009), Adressing between-study heterogeneity and inconsistency in mixed treatment comparisons: Application to stroke prevention treatments in individuals with non-rheumatic atrial fibrillation, Statistics in Medicine 28:1861-1881. [doi:10.1002/sim.3594]

Examples

blobbogram

Plot a blobbogram (AKA forest plot)

Description

blobbogram is a flexible function for creating blobbograms (forest plots), making no specific assumptions about the data being plotted. It supports column and row grouping as well as pagination.

Usage

```
blobbogram(data, id.label="Study", ci.label="Mean (95% CI)",
    left.label=NULL, right.label=NULL, center.label=NULL,
    log.scale=FALSE, xlim=NULL, styles=NULL,
```

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```
grouped=TRUE, group.labels=NULL,
  columns=NULL, column.labels=NULL,
  column.groups=NULL, column.group.labels=NULL,
  digits=2,
  ask=dev.interactive(orNone=TRUE),
  draw.no.effect=TRUE)
forest(x, ...)
```

Arguments

Х

8	
data	A data frame containing one row for each confidence interval to be visualized. The data format is described below.
id.label	Label to show above the row-id column.
ci.label	Label to show above the confidence intervals.
left.label	Label to show on the left-hand side of the no-difference line.
right.label	Label to show on the right-hand side of the no-difference line.
center.label	Label to show center-aligned with the no-difference line.
log.scale	If TRUE, the confidence intervals are given on a log scale, and axis labels will be exp() transformed.
xlim	The scale limits of the plot, if the confidence interval exceeds these limits an arrow will be shown at the limit. If unspecified, limits will be chosen that encompass all confidence intervals.
styles	A data frame describing the different row styles. By default, the styles "normal", "pooled" and "group" are defined.
grouped	If TRUE, and group.labels are specified, rows will be grouped according to the "group" column given in the data argument.
group.labels	Vector of group labels.
columns	Additional user-defined columns to be shown (names of columns given in the data argument).
column.labels	A vector of labels for the user-defined columns.
column.groups	Column groups, a numeric vector specifying the column group for each column.
column.group.l	abels
	A vector of labels for the column groups.
digits	The number of (significant) digits to print.
ask	If TRUE, a prompt will be displayed before generating the next page of a multipage plot.
draw.no.effect	If TRUE, draw the no-effect line.

An object to create a forest plot of.

Additional arguments.

blobbogram 7

Details

The forest function is a generic S3 method (definition compatible with the meta package). This package defines methods for mtc.result and mtc.relative.effect.table.

The blobbogram function creates a blobbogram (forest plot) from the given data (point estimates and confidence intervals) and meta-data (labels, column specifications, column groups, row groups, styles) using the grid package. If the plot would not fit the device's graphics region, the content is broken up into multiple plots generated in sequence (pagination).

The data argument is given as a data frame containing the following columns:

• id: identifier (label) for this row.

• group (optional): row group this row belongs to (indexes into the group.labels argument).

• pe: point estimate.

• ci.l: lower confidence interval limit.

• ci.u: upper confidence interval limit.

• style: the style to apply to this row (defined in the styles argument).

Additional user-defined columns can be specified using the columns and column.labels arguments.

The styles argument is given as a data frame containing the following columns:

• style: name of the style.

• weight: font weight.

• pe.style: symbol to draw for the point estimate ("circle" or "square", currently).

Value

None.

Note

This method should not be considered stable. We intend to generalize it further and possibly provide it in a separate package. The interface may change at any time.

Author(s)

Gert van Valkenhoef, Joël Kuiper

See Also

meta::forest, grid::Grid

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Examples

```
data <- read.table(textConnection('</pre>
"Study 1" 1 0.35 0.08 0.92 "normal" "2/46"
"Study 2" 1 0.43 0.15 1.14 "normal" "4/50"
"Study 3" 2 0.31 0.07 0.74 "normal" "2/97"
"Study 4" 2 0.86 0.34 2.90 "normal" "9/104"
"Study 5" 2 0.33 0.10 0.72 "normal" "4/74"
"Study 6" 2 0.47 0.23 0 91 "-
                                                                        value.A
                                                                                       value.B
                                                                           "7/46"
                                                                            "8/49"
                                                                            "10/100"
                            0.86 0.34 2.90 "normal" "9/104"
                                                                            "6/105"
                                                                            "14/74"
                            0.47 0.23 0.91 "normal" "11/120" "22/129"
 "Pooled"
               NA
                             0.42 0.15 1.04 "pooled" NA
 '), header=TRUE)
 data$pe <- log(data$pe)</pre>
 data$ci.l <- log(data$ci.l)</pre>
 data$ci.u <- log(data$ci.u)</pre>
 blobbogram(data, group.labels=c('GROUP 1', 'GROUP 2'),
      columns=c('value.A', 'value.B'), column.labels=c('r/n', 'r/n'),
      column.groups=c(1, 2), grouped=TRUE,
      column.group.labels=c('Intervention', 'Control'),
      id.label="Trial", ci.label="Odds Ratio (95% CrI)", log.scale=TRUE)
```

blocker

Beta blockers to prevent mortality after myocardial infarction

Description

A dataset of 22 trials investigating beta blockers versus control to prevent mortality after myocardial infarction. Number of events and sample size.

Format

A meta-analysis dataset containing 44 rows of arm-based data (responders and sample size).

Source

- S. Dias, A.J. Sutton, A.E. Ades, and N.J. Welton (2013a), *A Generalized Linear Modeling Framework for Pairwise and Network Meta-analysis of Randomized Controlled Trials*, Medical Decision Making 33(5):607-617. [doi:10.1177/0272989X12458724]
- J.B. Carlin (1992), *Meta-analysis for 2* \times 2 *tables: a Bayesian approach*, Statistics in Medicine 11(2):141-158. [doi:10.1002/sim.4780110202]

certolizumab 9

certolizumab

Certolizumab Pegol (CZP) for Rheumatoid Arthritis

Description

A dataset of 12 trials investigating 6 treatments and placebo for rheumatoid arthritis. The main outcome is the number of patients who improved by at least 50% on the American College of Rheumatology scale (ACR50) at 6 Months. A covariate is present for the mean disease duration at baseline (years).

Format

A network meta-regression dataset containing 24 rows of arm-based data (responders and sample size).

Source

S. Dias, A.J. Sutton, N.J. Welton, and A.E. Ades (2013b), *Heterogeneity - Subgroups, Meta-Regression, Bias, and Bias-Adjustment*, Medical Decision Making 33(5):618-640. [doi:10.1177/0272989X13485157]

Examples

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depression

Treatment response in major depression

Description

A dataset of 111 trials investigating 12 treatments for major depression on treatment response. Treatment response was defined as a reduction of at least 50% from the baseline score on the HAMD or MADRS at week 8 (or, if not available, another time between week 6 and 12).

Format

A network meta-analysis dataset containing 224 rows of arm-based data (responders and sample size).

Source

Cipriani et al. (2009), Comparative efficacy and acceptability of 12 new-generation antidepressants: a multiple-treatments meta-analysis, Lancet 373(9665):746-758. [doi:10.1016/S0140-6736(09)60046-5]

dietfat

Effects of low-fat diets on mortality

Description

A dataset of 10 trials investigating low-fat diet versus control diet for mortality. Number of events and exposure in person-years.

Format

A meta-analysis dataset containing 20 rows of arm-based data (responders, exposure, and sample size).

Source

S. Dias, A.J. Sutton, A.E. Ades, and N.J. Welton (2013a), *A Generalized Linear Modeling Framework for Pairwise and Network Meta-analysis of Randomized Controlled Trials*, Medical Decision Making 33(5):607-617. [doi:10.1177/0272989X12458724]

Hooper et al. (2000), *Reduced or modified dietary fat for preventing cardiovascular disease*, Cochrane Database of Systematic Reviews 2:CD002137. [doi:10.1002/14651858.CD002137]

hfPrevention 11

hfPrevention Statins versus placebo in primary and secondary preventi failure	ion of heart
----------------------------------------------------------------------------------	--------------

Description

A dataset of 19 trials comparing statins versus placebo or usual care for cholesterol lowering. The main outcome is the number of deaths. Trials are either primary prevention (no previous heart disease; secondary = 0) or secondary prevention (previous heart disease; secondary = 1).

Format

A meta-regression dataset containing 38 rows of arm-based data (responders and sample size).

Source

S. Dias, A.J. Sutton, N.J. Welton, and A.E. Ades (2013b), *Heterogeneity - Subgroups, Meta-Regression, Bias, and Bias-Adjustment*, Medical Decision Making 33(5):618-640. [doi:10.1177/0272989X13485157]

Examples

11.call

Call a likelihood/link-specific function

Description

GeMTC implements various likelihood/link combinations. Functionality specific to the likelihood/link is handled by methods with names ending in .likelihood>.hood>.hood>.hood>.

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Usage

```
11.call(fnName, model, ...)
```

Arguments

fnName The name of the function to call. See details for available functions.

Model An object of S3 class mtc.model describing a network meta-analysis model, or a list containing elements named 'likelihood' and 'link'.

Additional arguments to be passed to the function.

Details

The following methods currently need to be implemented to implement a likelihood/link:

- mtc.arm.mle: calculates a (corrected) maximum likelihood estimate for an arm-level effect.
 Used to generate starting values.
- mtc.rel.mle: calculates a (corrected) maximum likelihood estimate for a relative effect. Used to generate starting values.
- mtc.code.likelihood: generates JAGS code implementing the likelihood.
- scale.log: returns TRUE if plots should use the log scale.
- scale.name: returns the user-facing name of the outcome metric.
- scale.limit.inits: returns an upper and lower bound for the initial values, because some initial values might trigger boundary conditions such as probability 0 or 1 for the binomial.
- required.columns.ab: returns the required columns for arm-based data.

The first two methods can now also be used to selectively apply continuity corrections in case the maximum likelihood estimates are used for other purposes. mtc.arm.mle has an additional k=0.5 argument to specify the correction factor. mtc.rel.mle has arguments correction.force=TRUE to force application of the continuity correction even if unnecessary, correction.type="constant" to specify the type of correction (specify "reciprocal") for a correction proportional to the reciprocal of the size of the other arm, and correction.magnitude=1 to specify the (total) magnitude of the correction. These corrections apply only for count data, and will be ignored for continuous likelihood/links.

Value

The return value of the called function.

Author(s)

Gert van Valkenhoef

See Also

mtc.model

mtc.anohe 13

Examples

```
# The "model" may be a stub.
model <- list(likelihood="poisson", link="log")

ll.call("scale.name", model)
# "Hazard Ratio"

ll.call("mtc.arm.mle", model, c('responders'=12, 'exposure'=80))
# mean sd
#-1.8562980 0.1118034</pre>
```

mtc.anohe

Analysis of heterogeneity (ANOHE)

Description

(EXPERIMENTAL) Generate an analysis of heterogeneity for the given network. Three types of model are estimated: unrelated study effects, unrelated mean effects, and consistency. Output of the summary function can passed to plot for a visual representation.

Usage

```
mtc.anohe(network, ...)
```

Arguments

network An object of S3 class mtc.network.

Arguments to be passed to mtc.run or mtc.model. This can be used to set the likelihood/link or the number of iterations, for example.

Details

Analysis of heterogeneity is intended to be a unified set of statistics and a visual display that allows the simultaneous assessment of both heterogeneity and inconsistency in network meta-analysis [van Valkenhoef et al. 2014b (draft)].

mtc.anohe returns the MCMC results for all three types of model. To get appropriate summary statistics, call summary() on the results object. The summary can be plotted.

To control parameters of the MCMC estimation, see mtc.run. To specify the likelihood/link or to control other model parameters, see mtc.model. The ... arguments are first matched against mtc.run, and those that do not match are passed to mtc.model.

Value

For mtc. anohe: an object of class mtc. anohe. This is a list with the following elements:

```
result.use The result for the USE model (see mtc.run).

result.ume The result for the UME model (see mtc.run).
```

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```
result.cons The result for the consistency model (see mtc.run).
```

For summary: an object of class mtc.anohe.summary. This is a list with the following elements:

```
cons.model Generated consistency model.

studyEffects Study-level effect summaries (multi-arm trials downweighted).

pairEffects Pair-wise pooled effect summaries (from the UME model).

consEffects Consistency effect summaries.

indEffects Indirect effect summaries (back-calculated).

isquared.comp Per-comparison I-squared statistics.

isquared.glob Global I-squared statistics.
```

Note

This method should not be considered stable. It is an experimental feature and heavily work in progress. The interface may change at any time.

Author(s)

Gert van Valkenhoef, Joël Kuiper

See Also

```
mtc.model mtc.run
```

mtc.data.studyrow Convert one-study-per-row datasets

Description

Converts datasets in the one-study-per-row format to one-arm-per-row format used by GeMTC

Usage

```
mtc.data.studyrow(data,
    armVars=c('treatment'='t', 'responders'='r', 'sampleSize'='n'),
    nArmsVar='na',
    studyVars=c(),
    studyNames=1:nrow(data),
    treatmentNames=NA,
    patterns=c('%s..', '%s..%d.'))
```

mtc.data.studyrow 15

Arguments

data Data in one-study-per-row format.

armVars Vector of per-arm variables. The name of each component will be the column

name in the resulting dataset. The column name in the source dataset is derived

from the value of each component.

nArmsVar Variable holding the number of arms for each study.

studyVars Vector of per-study variables. The name of each component will be the column

name in the resulting dataset. The column name in the source dataset is derived

from the value of each component.

studyNames Vector of study names.

treatmentNames Vector of treatment names.

patterns Patterns to generate column names in the source dataset. The first is for per-

study variables, the second for per-arm variables.

Details

Maps the one-study-per-row format that is widely used and convenient for BUGS models to the one-arm-per-row format used by GeMTC. As the primary purpose is to input datasets from BUGS models, the defaults work for the standard BUGS data table format. In most cases, it should be possible to just copy/paste the BUGS data table (without the final 'END') and read.table it into R, then apply mtc.data.studyrow. In many cases, the resulting table can be processed directly by mtc.network.

Value

A data table with the requested columns.

Author(s)

Gert van Valkenhoef

See Also

```
mtc.network
```

Examples

```
## Example taken from the NICE DSU TSD series in Evidence Synthesis, #2
## Dopamine agonists for the treatment of Parkinson's

# Read the bugs-formatted data
data.src <- read.table(textConnection('
t[,1] t[,2] t[,3] y[,1] y[,2] y[,3] se[,1] se[,2] se[,3] na[]
1 3 NA -1.22 -1.53 NA 0.504 0.439 NA 2
1 2 NA -0.7 -2.4 NA 0.282 0.258 NA 2
1 2 A -0.3 -2.6 -1.2 0.505 0.510 0.478 3
3 4 NA -0.24 -0.59 NA 0.265 0.354 NA 2
3 4 NA -0.73 -0.18 NA 0.335 0.442 NA 2</pre>
```

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```
4 5 NA -2.2 -2.5 NA 0.197 0.190 NA 2
4 5 NA -1.8 -2.1 NA 0.200 0.250 NA 2'), header=TRUE)

# Convert the data, setting treatment names
data <- mtc.data.studyrow(data.src,
    armVars=c('treatment'='t', 'mean'='y', 'std.err'='se'),
    treatmentNames=c('Placebo', 'DA1', 'DA2', 'DA3', 'DA4'))

# Check that the data are correct
print(data)

# Create a network
network <- mtc.network(data)
```

mtc.deviance

Inspect residual deviance

Description

Inspect the posterior residual deviance and summarize it using plots

Usage

```
mtc.deviance(result)
mtc.devplot(x, ...)
mtc.levplot(x, ...)
## S3 method for class 'mtc.deviance'
plot(x, auto.layout=TRUE, ...)
```

Arguments

result An object of class mtc.result.

x An object of class mtc.deviance.

auto.layout If TRUE, the separate plots will be shown as panels on a single page.

Graphical parameters.

Details

mtc.devplot will generate a stem plot of the posterior deviance per arm (if there are only arm-based data) or the mean per data point deviance per study (if there are contrast-based data).

mtc.levplot will plot the leverage versus the square root of the residual deviance (mean per data point for each study).

The generic plot function will display both on a single page (unless auto.layout=FALSE).

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Value

mtc.deviance returns the deviance statistics of a mtc.result.

Author(s)

Gert van Valkenhoef

See Also

mtc.run

mtc.hy.prior

Set priors for the heterogeneity parameter

Description

These functions generate priors for the heterogeneity parameter in mtc.model. Priors can be set explicitly or, for outcomes on the log odds-ratio scale, based on empirical research.

Usage

```
mtc.hy.prior(type, distr, ...)
mtc.hy.empirical.lor(outcome.type, comparison.type)
```

Arguments

type	Type of he	terogeneity p	rior:	'std.dev',	'var', or	'prec'	for standard of	deviation,

variance, or precision respectively.

distr Prior distribution name (JAGS syntax). Typical ones would be 'dunif' (uni-

form), 'dgamma' (Gamma), or 'dlnorm' (log-normal). Use 'dhnorm' for the half-normal. Note that, as in JAGS, the precision (and not the variance or stan-

dard deviation) is used for the normal distribution and its derivatives.

. . Arguments to the distr. Can be numerical values or "om.scale" for the esti-

mated outcome measure scale (see mtc.model)

outcome.type The type of outcome to get an empirical prior for. Can be one of 'mortality'

(all-cause mortality), 'semi-objective' (e.g. cause-specific mortality, major morbidity event, drop-outs), or 'subjective' (e.g. pain, mental health, dichotomous

biomarkers).

comparison.type

The type of comparison to get an empirical prior for. Can be one of 'pharma-control' (pharmacological interventions versus control), 'pharma-pharma' (pharmacological versus pharmacological interventions) and 'non-pharma' (any other comparisons).

Details

The generated prior is a list, the structure of which may change without notice. It can be converted to JAGS compatible code using as . character.

Empirical priors for the log odds-ratio (LOR) are taken from [Turner et al. 2012].

Value

A value to be passed to mtc.model.

Author(s)

Gert van Valkenhoef

See Also

mtc.model

Examples

```
# NOTE: the mtc.run commands below are for illustrative purposes, such a small
# number of iterations should obviously not be used in practice.

# set a uniform prior standard deviation
model1 <- mtc.model(smoking, hy.prior=mtc.hy.prior("std.dev", "dunif", 0, 2))
result <- mtc.run(model1, n.adapt=10, n.iter=10)

# set an empirical (log-normal) prior on the variance
model2 <- mtc.model(smoking, hy.prior=mtc.hy.empirical.lor("subjective", "non-pharma"))
result <- mtc.run(model2, n.adapt=10, n.iter=10)

# set a gamma prior on the precision
model3 <- mtc.model(smoking, hy.prior=mtc.hy.prior("prec", "dgamma", 0.01, 0.01))
result <- mtc.run(model3, n.adapt=10, n.iter=10)</pre>
```

mtc.model

Generate network meta-analysis models

Description

The mtc.model function generates network meta-analysis models from an mtc.network object.

Usage

```
mtc.model(network, type = "consistency", factor = 2.5, n.chain = 4,
    likelihood=NULL, link=NULL, linearModel="random",
    om.scale=NULL, hy.prior=mtc.hy.prior("std.dev", "dunif", 0, "om.scale"),
    dic=TRUE, powerAdjust=NA, ...)
```

Arguments

network An object of S3 class mtc.network

type A string literal indicating the type of model (allowed values are "consistency",

"regression", "nodesplit", "ume", or "use").

factor Variance scaling factor for the starting values

n. chain Number of chains in the model

likelihood The likelihood to be used. If unspecified, a suitable likelihood will be inferred

for the given data.

link The link function to be used. If unspecified, a suitable link function will be

inferred for the given data.

linearModel The type of linear model to be generated. Can be "random" for a random effects

model, or "fixed" for a fixed effect model.

om. scale Outcome measure scale. Represents a "very large" difference on the analysis'

outcome scale. This is used to set vague priors. For the log odds-ratio, values between 2 and 5 are considered reasonable. For continuous outcomes, this depends heavily on the specific outcome. If left unspecified, it is determined from

the data.

hy.prior Heterogeneity prior. See mtc.hy.prior.

dic When set to TRUE, deviance and fitted values will be monitored to allow com-

putation of the Deviance Information Criterion (DIC) at residual.

powerAdjust Optional: the name of a column in the studies data frame of the mtc.network.

This column must contain values between 0 and 1. The likelihood for each study will be adjusted by inflating the variance, where 0 means the study is excluded

and 1 means it receives full weight. See details for more.

... Additional arguments to be passed to the type-specific model generation func-

tion.

Details

The mtc.model function generates an object of S3 class mtc.model, which can be visualized by the generic plot function or summarized by the generic summary function.

These likelihood/links are supported:

• normal/identity: for continuous (mean difference) data.

Required columns: [mean, std.err] or [mean, std.dev, sampleSize].

Result: relative mean difference.

• binom/logit: for dichotomous data.

Required columns [responders, sampleSize].

Result: (log) odds ratio.

• binom/log: for dichotomous data.

Required columns [responders, sampleSize].

Result: (log) risk ratio.

• binom/cloglog: for rate (survival) data - equal follow-up in each arm. Required columns [responders, sampleSize].

Result: (log) hazard ratio.

• poisson/log: for rate (survival) data.

Required columns [responders, exposure].

Result: (log) hazard ratio.

Most likelihood/links follow [Dias et al. 2013a], and the binom/log model follows [Warn et al. 2002].

The following model types are supported:

- consistency: ordinary consistency model. No additional parameters. [Dias et al. 2013a, van Valkenhoef et al. 2012]
- nodesplit: node-splitting model. Removes both arms used to estimate the direct evidence from the network of indirect evidence, rather than just one of those arms. This means that three-arm trials do not contribute any evidence in the network of indirect evidence. When relative effect data are present, these are transformed appropriately (using an assumption of normality) to enable this direct/indirect evidence split. Additional parameters: t1 and t2, which indicate the comparison to be split. [Dias et al. 2010, van Valkenhoef et al. 2015]
- regression: meta-regression model. Additional parameters: regressor, which indicates how to structure the treatment-interaction model for the regression. See below for details.
- use: unrelated study effects. Models the effects within each study as if the studies are independent. No additional parameters. [van Valkenhoef et al. (draft)]
- ume: unrelated mean effects. Models the effects within each comparison as if they are independent. Does not properly handle multi-arm trials, and warns when they are present in the network. No additional parameters. [Dias et al. 2013b, van Valkenhoef et al. (draft)]

Regressor specification: a list with elements: variable, coefficient, and either control or classes. The variable is the name of the covariate to include in the regression analysis, and must be a column of the studies data frame in the network. The regressor variable is automatically centered and standardized using the method recommended by Gelman et al. (2008). The coefficient indicates the type of treatment-interaction model: "shared", "unrelated", or "exchangeable". control, if specified, must be the ID of a treatment in the network. All other treatments have a coefficient relative to the control, which can be the same for all treatments ("shared"), different for all treatments ("unrelated") or exchangeable between treatments ("exchangeable"). classes is a named list of treatment classes, the first of which will act as the control class. Each class is a vector of treatment IDs from the network. Only "shared" coefficients can currently be used, meaning a single coefficient per class (except the control class). See also atrialFibrillation, certolizumab, and hfPrevention for examples of meta-regression analyses.

Studies can be downweighted by using the powerAdjust argument, which applies a variance inflation (also known as "power prior") to the likelihood. This allows a weight $\alpha_i \in [0,1]$ to be specified for each study i. The log-likelihood will be multiplied by a factor α_i , or equivalently for normal distributions the variance will be multiplied by $1/\alpha_i$. Setting $\alpha_i = 0$ will completely exclude that study, whereas setting $\alpha_i = 1$ will weight it fully. Essentially, down-weighted models modify the data and hence model fit statistics such as DIC can not be compared between models with different weightings.

Value

An object of class mtc.model. The following elements are descriptive:

type The type of model

network Network the model was generated from

tree Spanning tree formed by the basic parameters

var.scale The scaling factor used to over-disperse starting values

likelihood The likelihood used

link The link function used

om. scale The scale for the variance parameters

regressor Regressor specification (regression models only): includes additional elements

"center" and "scale" describing how the regressor input was standardized

These elements determine the model run by JAGS:

n.chain The number of chains

code Model code in JAGS syntax. Use cat() for proper formatting.

data Data in JAGS compatible format

inits Initial values in JAGS compatible format

monitors The nodes of the JAGS model to monitor

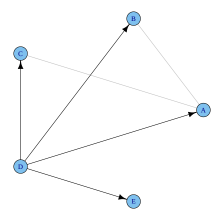
dic Whether to compute the DIC

These latter fields can be modified to alter the statistical model, but such changes may break the model or assumptions made elsewhere in the package.

Visualization

Calling the generic plot method on an S3 mtc.model object will show a graph with the treatments as vertices and the comparisons as edges. The lines with solid arrows represent basic parameters, and the other lines represent comparisons that are not associated with any parameter but do have direct evidence from trials.

The example code will generate the following graph:



The default layout algorithm is igraph::layout.circle, other layout algorithms can be used by passing them as an optional argument called layout to plot. The igraph::layout.fruchterman.reingold algorithm also seems to produce nice results and may be better for large graphs.

Author(s)

Gert van Valkenhoef, Joël Kuiper

See Also

```
mtc.network, mtc.run
```

Examples

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mtc.network	Create an mtc.network

Description

Creates an object of class mtc.network

Usage

Arguments

data.ab Arm-level data. A data frame defining the arms of each study, containing the

columns 'study' and 'treatment', where 'treatment' must refer to an existing treatment ID if treatments were specified. Further columns define the data per arm, and depend on the likelihood/link to be used. See mtc.model for supported

likelihood/links and their data requirements.

data.re Relative effect data. A data frame defining the arms of each study, containing

the columns 'study' and 'treatment', where 'treatment' must refer to an existing treatment ID if treatments were specified. The column 'diff' specifies the mean difference between the current arm and the baseline arm; set 'diff=NA' for the baseline arm. The column 'std.err' specifies the standard error of the mean difference (for non-baseline arms). For trials with more than two arms, specify the standard error of the mean of the baseline arm in 'std.err', as this determines

the covariance of the differences.

treatments Optional. A data frame with columns 'id' and 'description' defining the treat-

ments or a vector giving the treatment IDs.

studies Optional. A data frame with a 'study' column naming the studies and additional

columns containing covariate values.

description Optional. Short description of the network.

data Deprecated. Arm-level data; automatically assigned to data.ab if it is not speci-

fied. Present for compatibility with older versions.

x An mtc.network object.

layout An igraph-compatible layout.

dynamic.edge.width

If set to TRUE, dynamically set the edge width based on the number of studies.

use.description

Display treatment descriptions instead of treatment IDs.

.. Additional arguments passed to plot.igraph.

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Details

One-arm trials are automatically removed, which results in a warning.

Also see mtc.data.studyrow for a convenient way to import data from the one-study-per-row format, which is very popular for BUGS code.

Value

For mtc.network, an object of the class mtc.network which is a list containing:

```
description A short description of the network
treatments A data frame describing the treatments
data.ab A data frame containing the network data (arm-level)
data.re A data frame containing the network data (relative effects)
studies A data frame containing study-level information (covariates)
```

These are cleaned up and standardized versions of the arguments provided, or generated defaults for 'treatments' if the argument was omitted.

Author(s)

Gert van Valkenhoef, Joël Kuiper

See Also

```
mtc.data.studyrow, mtc.model
```

Examples

```
# Create a new network by specifying all information.
treatments <- read.table(textConnection('</pre>
 id description
      "Treatment A"
      "Treatment B"
     "Treatment C"'), header=TRUE)
data <- read.table(textConnection('</pre>
 study treatment responders sampleSize
 01
        Α
                    2
                                100
 01
        В
                    5
                                100
 02
        В
                    6
                                110
        С
                                110
 02
                    1
 03
                    3
                                60
        Α
 03
        С
                    4
                                80
                    7
 03
        В
                                80'), header=TRUE)
network <- mtc.network(data, description="Example", treatments=treatments)</pre>
# Create a new network by specifying only the data.
data <- read.table(textConnection('</pre>
 study treatment mean std.dev sampleSize
               -1.12 0.6
 01
        Α
                                    15
```

mtc.nodesplit 25

```
-1.55 0.5
                                   16
 02
        Α
                   -0.8 0.7
                                   33
 02
        В
                   -1.1 0.5
                                   31'), header=TRUE)
network <- mtc.network(data)</pre>
# Print the network
print(network)
## MTC dataset: Network
##
    study treatment mean std.dev sampleSize
## 1
                 A -1.12
                           0.6
        1
                 B -1.55
## 2
        1
                              0.5
                                          16
## 3
        2
                  A -0.80
                             0.7
                                          33
## 4
        2
                  B -1.10
                             0.5
                                          31
```

mtc.nodesplit

Node-splitting analysis of inconsistency

Description

Generate and run an ensemble of node-splitting models, results of which can be jointly summarized and plotted.

Usage

```
mtc.nodesplit(network, comparisons=mtc.nodesplit.comparisons(network), ...)
mtc.nodesplit.comparisons(network)
```

Arguments

network An object of S3 class mtc.network.

comparisons Data frame specifying the comparisons to be split. The frame has two columns: 't1' and 't2'.

Arguments to be passed to mtc.run or mtc.model. This can be used to set the likelihood/link or the number of iterations, for example.

Details

mtc.nodesplit returns the MCMC results for all relevant node-splitting models [van Valkenhoef et al. 2015]. To get appropriate summary statistics, call summary() on the results object. The summary can be plotted. See mtc.model for details on how the node-splitting models are generated.

To control parameters of the MCMC estimation, see mtc.run. To specify the likelihood/link or to control other model parameters, see mtc.model. The ... arguments are first matched against mtc.run, and those that do not match are passed to mtc.model.

mtc.nodesplit.comparisons returns a data frame enumerating all comparisons that can reasonably be split (i.e. have independent indirect evidence).

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Value

For mtc.nodesplit: an object of class mtc.nodesplit. This is a list with the following elements:

d. X. Y For each comparison (t1=X, t2=Y), the MCMC results

consistency The consistency model results

For summary: an object of class mtc.nodesplit.summary. This is a list with the following elements:

dir.effect Summary of direct effects for each split comparison ind.effect Summary of indirect effects for each split comparison

cons.effect Summary of consistency model effects for each split comparison

p.value Inconsistency p-values for each split comparison

cons.model The generated consistency model

Author(s)

Gert van Valkenhoef, Joël Kuiper

See Also

```
mtc.model mtc.run
```

Examples

```
# Run all relevant node-splitting models
## Not run: result.ns <- mtc.nodesplit(parkinson, thin=50)
# (read results from file instead of running:)
result.ns <- readRDS(system.file('extdata/parkinson.ns.rds', package='gemtc'))
# List the individual models
names(result.ns)
# Time series plots and convergence diagnostics for d.A.C model
plot(result.ns$d.A.C)
gelman.diag(result.ns$d.A.C, multivariate=FALSE)
# Overall summary and plot
summary.ns <- summary(result.ns)
print(summary.ns)
plot(summary.ns)</pre>
```

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mtc.run

Running an mtc.model using an MCMC sampler

Description

The function mtc.run is used to generate samples from a object of type mtc.model using a MCMC sampler. The resulting mtc.result object can be coerced to an mcmc.list for further analysis of the dataset using the coda package.

Usage

```
mtc.run(model, sampler = NA, n.adapt = 5000, n.iter = 20000, thin = 1)
## S3 method for class 'mtc.result'
summary(object, ...)
## S3 method for class 'mtc.result'
plot(x, ...)
## S3 method for class 'mtc.result'
forest(x, use.description=FALSE, ...)
## S3 method for class 'mtc.result'
print(x, ...)
## S3 method for class 'mtc.result'
as.mcmc.list(x, ...)
```

Arguments

model	An object of S3 class mtc.model describing a network meta-analysis model.	
sampler	Deprecated: gemtc now only supports the JAGS sampler. Specifying a sampler will result in a warning or error. This argument will be removed in future versions.	
n.adapt	Amount of adaptation (or tuning) iterations.	
n.iter	Amount of simulation iterations.	
thin	Thinning factor.	
object	Object of S3 class mtc.result.	
х	Object of S3 class mtc.result.	
use.description		
	Display treatment descriptions instead of treatment IDs.	
	Additional arguments.	

Value

An object of class mtc.result. This is a list with the following elements:

samples The samples resulting from running the MCMC model, in mcmc.list format.

model The mtc.model used to produce the samples.

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deviance

Residual deviance statistics, a list with the following elements. DIC: deviance information criterion at residual (Dbar + pD). Dbar: mean sum of residual deviance. pD: sum of leverage, also known as the effective number of parameters. dev.ab: mean posterior residual deviance of each arm (for arm-based data). fit.ab: deviance at the posterior mean of te fitted values (for arm-based data). dev.re: mean posterior residual deviance of each study (for relative-effect data). fit.re: deviance at the posterior mean of the fitted values (for relative-effect data).

The object can be coerced to an mcmc.list from the coda package by the generic S3 method as.mcmc.list.

Analysis of the results

Convergence of the model can be assessed using methods from the coda package. For example the Brooks-Gelman-Rubin method (coda::gelman.diag, coda::gelman.plot). The summary also provides useful information, such as the MCMC error and the time series and densities given by plot should also be inspected.

The forest function can provide forest plots for mtc.result objects. This is especially useful in combination with the relative.effect function that can be used to calculate relative effects compared to any baseline for consistency models. The rank.probability function calculates rank probabilities for consistency models.

Author(s)

Gert van Valkenhoef, Joël Kuiper

See Also

```
mtc.model
relative.effect.table, relative.effect, rank.probability
coda::gelman.diag, coda::gelman.plot
```

Examples

```
model <- mtc.model(smoking)

## Not run: results <- mtc.run(model, thin=10)
results <- dget(system.file("extdata/luades-smoking.samples.gz", package="gemtc"))

# Convergence diagnostics
gelman.plot(results)

# Posterior summaries
summary(results)

## Iterations = 5010:25000

## Thinning interval = 10

## Number of chains = 4

## Sample size per chain = 2000

##</pre>
```

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```
## 1. Empirical mean and standard deviation for each variable,
     plus standard error of the mean:
##
##
                    SD Naive SE Time-series SE
           Mean
## d.A.B 0.4965 0.4081 0.004563
                                      0.004989
## d.A.C 0.8359 0.2433 0.002720
                                      0.003147
## d.A.D 1.1088 0.4355 0.004869
                                      0.005280
## sd.d 0.8465 0.1913 0.002139
                                      0.002965
## 2. Quantiles for each variable:
##
##
            2.5%
                    25%
                           50%
                                  75% 97.5%
## d.A.B -0.2985 0.2312 0.4910 0.7530 1.341
## d.A.C 0.3878 0.6720 0.8273 0.9867 1.353
## d.A.D 0.2692 0.8197 1.0983 1.3824 2.006
## sd.d 0.5509 0.7119 0.8180 0.9542 1.283
plot(results) # Shows time-series and density plots of the samples
forest(results) # Shows a forest plot
```

parkinson

Mean off-time reduction in Parkinson's disease

Description

A dataset of seven trials investigating four treatments and placebo for Parkinson's disease. The outcome is mean off-time reduction.

Format

- parkinson: A network meta-analysis dataset containing fifteen rows of arm-based data (mean, standard deviation, and sample size).
- parkinson_diff: A network meta-analysis dataset containing fifteen rows of contrast-based data.
- parkinson_shared: A network meta-analysis dataset containing mixed arm-based and contrastbased data.

Source

Franchini et al. (2012), *Accounting for correlation in network meta-analysis with multi-arm trials*, Research Synthesis Methods, 3(2):142-160. [doi:10.1002/jrsm.1049]

30 plotCovariateEffect

plotCovariateEffect Plot treatment effects versus covariate values

Description

The plot will show the median treatment effect and the 95% credible interval on the y-axis and the covariate value on the x-axis. One plot page will be generated per pair of treatments.

Usage

Arguments

result	Results object - created by mtc.result
t1	A list of baseline treatments to calculate treatment effects against. Will be extended to match the length of t2.
t2	A list of treatments to calculate the effects for. Will be extended to match the length of t1. If left empty and t1 is a single treatment, effects of all treatments except t1 will be calculated.
xlim	The x-axis limits.
ylim	The y-axis limits.
ask	If TRUE, a prompt will be displayed before generating the next page of a multipage plot.

Details

Default x-axis limits will be set to three standard deviations above and below the centering value of the covariate. The y-axis limits will be set based on the minimum and maximum 95% CrI limits among the set of effects computed.

Value

None.

Author(s)

Gert van Valkenhoef

See Also

relative.effect, mtc.run

rank.probability 31

|--|

Description

Rank probabilities indicate the probability for each treatment to be best, second best, etc.

Usage

```
rank.probability(result, preferredDirection=1, covariate=NA)

## S3 method for class 'mtc.rank.probability'
print(x, ...)

## S3 method for class 'mtc.rank.probability'
plot(x, ...)

sucra(ranks)
rank.quantiles(ranks, probs=c("2.5%"=0.025, "50%"=0.5, "97.5%"=0.975))
```

Arguments

result	Object of S3 class mtc.result to be used in creation of the rank probability table
preferredDirec	ction
	Preferential direction of the outcome. Set 1 if higher values are preferred, -1 if lower values are preferred.
covariate	(Regression analyses only) Value of the covariate at which to compute rank probabilities.
х	An object of S3 class rank.probability.
• • •	Additional arguments.
ranks	A ranking matrix where the treatments are the rows (e.g. the result of rank.probability).

Details

probs

For each MCMC iteration, the treatments are ranked by their effect relative to an arbitrary baseline. A frequency table is constructed from these rankings and normalized by the number of iterations to give the rank probabilities.

Probabilities at which to give quantiles.

Value

rank.probability: A matrix (with class mtc.rank.probability) with the treatments as rows and the ranks as columns. sucra: A vector of SUCRA values. rank.quantiles: A matrix with treatments as rows and quantiles as columns, giving the quantile ranks (by default, the median and 2.5% and 97.5% ranks).

32 read.mtc.network

Author(s)

Gert van Valkenhoef, Joël Kuiper

See Also

```
relative.effect
```

Examples

```
model <- mtc.model(smoking)</pre>
# To save computation time we load the samples instead of running the model
## Not run: results <- mtc.run(model)</pre>
results <- dget(system.file("extdata/luades-smoking.samples.gz", package="gemtc"))
ranks <- rank.probability(results)</pre>
print(ranks)
## Rank probability; preferred direction = 1
         [,1]
                  [,2]
                          [,3]
## A 0.000000 0.003000 0.105125 0.891875
## B 0.057875 0.175875 0.661500 0.104750
## C 0.228250 0.600500 0.170875 0.000375
## D 0.713875 0.220625 0.062500 0.003000
print(sucra(ranks))
##
                       В
                                  С
                                              D
           Α
## 0.03670833 0.39591667 0.68562500 0.88175000
print(rank.quantiles(ranks))
   2.5% 50% 97.5%
## A
        3
## B
                  4
        1
            3
## C
                  3
        1
            2
## D
                  3
        1
            1
plot(ranks) # plot a cumulative rank plot
plot(ranks, beside=TRUE) # plot a 'rankogram'
```

read.mtc.network

Read mtc.network from legacy GeMTC XML files (.gemtc)

Description

Read legacy GeMTC XML (.gemtc) files.

Usage

```
read.mtc.network(file)
```

relative.effect 33

Arguments

file

Path to the file to read (read.mtc.network).

Details

read.mtc.network exists to migrate data from the older GeMTC XML format, and requires the XML package to be installed.

Support for writing XML files has been removed. Instead, use native R methods to read and write networks (e.g. save/load or dput/dget).

Value

read.mtc.network returns an object of S3 class mtc.network.

Author(s)

Gert van Valkenhoef

See Also

mtc.network

Examples

```
# Read an example GeMTC XML file
file <- system.file("extdata/luades-smoking.gemtc", package="gemtc")
network <- read.mtc.network(file)

# Summarize the network (generate some interesting network properties)
summary(network)
## $Description
## [1] "MTC dataset: Smoking cessation rates"
##
## $`Studies per treatment`
## A B C D
## 19 6 19 6
##
## $`Number of n-arm studies`
## 2-arm 3-arm
## 22 2</pre>
```

relative.effect

Calculating relative effects

Description

Calculates the relative effects of pairs of treatments.

34 relative.effect

Usage

```
relative.effect(result, t1, t2 = c(), preserve.extra = TRUE, covariate = NA)
```

Arguments

result	An object of S3 class mtc.result to derive the relative effects from.
t1	A list of baselines to calculate a relative effects against. Will be extended to match the length of t2.
t2	A list of treatments to calculate the relative effects for. Will be extended to match the length of t1. If left empty and t1 is a single treatment, relative effects of all treatments except t1 will be calculated.
preserve.extra	Indicates whether to preserve extra parameters such as the sd.d.
covariate	(Regression analyses only) Value of the covariate at which to compute relative effects.

Value

Returns an mtc.results object containing the calculated relative effects.

Note that this method stores the raw samples, which may result in excessive memory usage. You may want to consider using relative.effect.table instead.

Author(s)

Gert van Valkenhoef, Joël Kuiper

See Also

```
rank.probability, relative.effect.table
```

Examples

```
model <- mtc.model(smoking)</pre>
# To save computation time we load the samples instead of running the model
## Not run: results <- mtc.run(model)</pre>
results <- dget(system.file("extdata/luades-smoking.samples.gz", package="gemtc"))</pre>
# Creates a forest plot of the relative effects
forest(relative.effect(results, "A"))
summary(relative.effect(results, "B", c("A", "C", "D")))
## Iterations = 5010:25000
## Thinning interval = 10
## Number of chains = 4
## Sample size per chain = 2000
## 1. Empirical mean and standard deviation for each variable,
##
      plus standard error of the mean:
##
                     SD Naive SE Time-series SE
##
            Mean
```

relative.effect.table 35

```
## d.B.A -0.4965 0.4081 0.004563
                                     0.004989
## d.B.C 0.3394 0.4144 0.004634
                                     0.004859
## d.B.D 0.6123 0.4789 0.005354
                                     0.005297
## sd.d 0.8465 0.1913 0.002139
                                     0.002965
## 2. Quantiles for each variable:
           2.5%
                    25%
                            50%
## d.B.A -1.3407 -0.7530 -0.4910 -0.2312 0.2985
## d.B.C -0.4809 0.0744 0.3411 0.5977 1.1702
## d.B.D -0.3083 0.3005 0.6044 0.9152 1.5790
## sd.d 0.5509 0.7119 0.8180 0.9542 1.2827
```

relative.effect.table Table of relative effects

Description

Generates a table of the relative effects of all pairs of treatments. Unlike relative.effect, this method stores summaries only, not raw samples.

Usage

```
relative.effect.table(result, covariate=NA)

## S3 method for class 'mtc.relative.effect.table'
print(x, ...)

## S3 method for class 'mtc.relative.effect.table'
forest(x, t1, use.description=FALSE, ...)

## S3 method for class 'mtc.relative.effect.table'
as.data.frame(x, ...)
```

Arguments

result An object of S3 class mtc.result to derive the relative effects from.

covariate (Regression analyses only) Value of the covariate at which to compute relative effects.

x An object of S3 class mtc.relative.effect.table.

t1 Baseline treatment for the Forest plot.

use.description
Display treatment descriptions instead of treatment IDs.

Additional arguments.

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Value

Returns an mtc.relative.effect.table object containing the quantiles of the calculated relative effects of all pair-wise comparisons among the treatments.

The result will be pretty printed as an n-by-n table of relative treatment effects. It can also be used to produce Forest plots against any arbitrary baseline. Finally, the as.data.frame generic method makes it possible to export the table for use in Excel or other spreadsheet software, using the core R methods write.csv or write.csv2.

Author(s)

Gert van Valkenhoef

See Also

```
relative.effect
```

Examples

```
model <- mtc.model(smoking)
# To save computation time we load the samples instead of running the model
## Not run: results <- mtc.run(model)
results <- dget(system.file("extdata/luades-smoking.samples.gz", package="gemtc"))
# Creates a forest plot of the relative effects
tbl <- relative.effect.table(results)
# Print the n*n table
print(tbl)
# Plot effect relative to treatment "C"
forest(tbl, "C")
# Write to CSV (e.g. to import to Excel, then use in a Word table)
## Not run: write.csv(tbl, "smoking-effects.csv")
# Note: use write.csv2 for Western European locales</pre>
```

smoking

Psychological treatments to aid smoking cessation

Description

A dataset of 24 trials investigating four psychological treatments and no treatment for smoking cessation. The outcome is the number of people who stopped smoking.

Format

A network meta-analysis dataset containing 50 rows of arm-based data (responders and sample size).

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Source

Lu and Ades (2006), Assessing Evidence Inconsistency in Mixed Treatment Comparisons, Journal of the American Statistical Society, 101(474):447-459. [doi:10.1198/016214505000001302]

Hasselblad (1998), *Meta-analysis of multitreatment studies*, Medical Decision Making 18(1):37-43. [doi:10.1177/0272989X9801800110]

thrombolytic

Thrombolytic treatment after acute myocardial infarction

Description

A dataset of 28 trials investigating eight thrombolytic treatments administered after a myocardial infarction. The outcome is mortality after 30-35 days.

Format

A network meta-analysis dataset containing 58 rows of arm-based data (responders and sample size).

Source

Lu and Ades (2006), Assessing Evidence Inconsistency in Mixed Treatment Comparisons, Journal of the American Statistical Society, 101(474):447-459. [10.1198/016214505000001302]

Boland et al. (2003), Early thrombolysis for the treatment of acute myocardial infarction: a systematic review and economic evaluation, Health Technology Assessment 7(15):1-136. [doi:10.3310/hta7150]

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