Package: metamisc (via r-universe)

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Type Package

Title Meta-Analysis of Diagnosis and Prognosis Research Studies

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Description Facilitate frequentist and Bayesian meta-analysis of diagnosis and prognosis research studies. It includes functions to summarize multiple estimates of prediction model discrimination and calibration performance (Debray et al., 2019) [<doi:10.1177/0962280218785504>](https://doi.org/10.1177/0962280218785504). It also includes functions to evaluate funnel plot asymmetry (Debray et al., 2018) [<doi:10.1002/jrsm.1266>](https://doi.org/10.1002/jrsm.1266). Finally, the package provides functions for developing multivariable prediction models from datasets with clustering (de Jong et al., 2021) [<doi:10.1002/sim.8981>](https://doi.org/10.1002/sim.8981).

Imports metafor (>= 2.0.0), mvtnorm, lme4, dplyr, plyr, methods, pROC, ggplot2

Depends R ($>= 3.5.0$), stats, graphics

Suggests coda, runjags, rjags, ggmcmc, logistf $(>= 1.23)$, test that $(>= 1.23)$ 1.0.2), knitr, rmarkdown, formatR

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Encoding UTF-8

URL <https://github.com/smartdata-analysis-and-statistics/metamisc>,

<https://smartdata-analysis-and-statistics.github.io/metamisc/>

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NeedsCompilation no

VignetteBuilder knitr

BugReports <https://github.com/smartdata-analysis-and-statistics/metamisc/issues>

Repository https://smartdata-analysis-and-statistics.r-universe.dev

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metamisc-package *Meta-Analysis of Diagnosis and Prognosis Research Studies*

Description

Facilitate frequentist and Bayesian meta-analysis of diagnosis and prognosis research studies. It includes functions to summarize multiple estimates of prediction model discrimination and calibration performance (Debray et al., 2019) <doi:10.1177/0962280218785504>. It also includes functions to evaluate funnel plot asymmetry (Debray et al., 2018) <doi:10.1002/jrsm.1266>. Finally, the package provides functions for developing multivariable prediction models from datasets with clustering (de Jong et al., 2021) <doi:10.1002/sim.8981>.

Details

The following functionality is currently implemented: univariate meta-analysis of summary data ([uvmeta](#page-65-1)), bivariate meta-analysis of correlated outcomes ([riley](#page-52-1)), meta-analysis of prediction model performance ([valmeta](#page-70-1)), evaluation of funnel plot asymmetry ([fat](#page-19-1)).

The metamisc package also provides a comprehensive framework for developing prediction models when patient-level data from multiple studies or settings are available ([metapred](#page-38-1)).

Author(s)

Thomas Debray <thomas.debray@gmail.com>, Valentijn de Jong <Valentijn.M.T.de.Jong@gmail.com>

References

de Jong VMT, Moons KGM, Eijkemans MJC, Riley RD, Debray TPA. Developing more generalizable prediction models from pooled studies and large clustered data sets. Stat Med. 2021;40(15):3533– 59.

Debray TPA, Moons KGM, Ahmed I, Koffijberg H, Riley RD. A framework for developing, implementing, and evaluating clinical prediction models in an individual participant data meta-analysis. Stat Med. 2013;32(18):3158–80.

Debray TPA, Damen JAAG, Riley R, Snell KIE, Reitsma JB, Hooft L, et al. A framework for metaanalysis of prediction model studies with binary and time-to-event outcomes. Stat Methods Med Res. 2019 Sep;28(9):2768–86.

Debray TPA, Damen JAAG, Snell KIE, Ensor J, Hooft L, Reitsma JB, et al. A guide to systematic review and meta-analysis of prediction model performance. BMJ. 2017;356:i6460.

Debray TPA, Moons KGM, Riley RD. Detecting small-study effects and funnel plot asymmetry in meta-analysis of survival data: a comparison of new and existing tests. Res Syn Meth. 2018;9(1):41–50.

Riley RD, Moons K, Snell KIE, Ensor J, Hooft L, Altman D, et al. A guide to systematic review and meta-analysis of prognostic factor studies. BMJ. 2019;364:k4597.

Riley RD, Tierney JF, Stewart LA. Individual participant data meta-analysis: a handbook for healthcare research. Hoboken, NJ: Wiley; 2021. ISBN: 978-1-119-33372-2.

Schmid CH, Stijnen T, White IR. Handbook of meta-analysis. First edition. Boca Raton: Taylor and Francis; 2020. ISBN: 978-1-315-11940-3.

Steyerberg EW, Nieboer D, Debray TPA, Van Houwelingen JC. Assessment of heterogeneity in an individual participant data meta-analysis of prediction models: An overview and illustration. Stat Med. 2019;38(22):4290–309.

See Also

[fat](#page-19-1), [metapred](#page-38-1), [riley](#page-52-1), [uvmeta](#page-65-1), [valmeta](#page-70-1)

acplot *Plot the autocorrelation of a Bayesian meta-analysis model*

Description

Function to display autocorrelation of a fitted Bayesian meta-analysis model.

Usage

```
acplot(...)
```
Arguments

... Additional arguments, which are currently ignored.

acplot.mcmc.list 5

Details

This is a generic function.

Value

A ggplot object.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

acplot.mcmc.list *Plot the autocorrelation of a Bayesian meta-analysis model*

Description

Function to display autocorrelation of a fitted Bayesian meta-analysis model.

Usage

```
## S3 method for class 'mcmc.list'
acplot(x, P, nLags = 50, greek = FALSE, ...)
```
Arguments

Value

A ggplot object.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

Function to display autocorrelation of a fitted Bayesian meta-analysis model.

Usage

```
## S3 method for class 'uvmeta'
acplot(x, \ldots)
```
Arguments

Details

Results are displayed for the estimated mean (mu) and standard-deviation (tau) of the meta-analysis model.

Value

A ggplot object.

An object of class ggplot

Author(s)

Thomas Debray <thomas.debray@gmail.com>

Examples

Not run: data(Roberts)

```
fit <- with(Roberts, uvmeta(r=SDM, r.se=SE, labels=rownames(Roberts), method="BAYES"))
acplot(fit)
```
End(Not run)

Function to display autocorrelation of a fitted Bayesian meta-analysis model.

Usage

```
## S3 method for class 'valmeta'
acplot(x, \ldots)
```
Arguments

Details

Results are displayed for the estimated mean (mu) and standard-deviation (tau) of the meta-analysis model.

Value

A ggplot object.

An object of class ggplot

Author(s)

Thomas Debray <thomas.debray@gmail.com>

Examples

```
## Not run:
data(EuroSCORE)
fit <- valmeta(cstat=c.index, cstat.se=se.c.index, cstat.cilb=c.index.95CIl,
              cstat.ciub=c.index.95CIu, N=n, O=n.events,
              data=EuroSCORE, method="BAYES", slab=Study)
acplot(fit)
## End(Not run)
```
The function calculates (transformed versions of) the concordance (c-) statistic with the corresponding sampling variance.

Usage

```
ccalc(
  cstat,
  cstat.se,
  cstat.cilb,
  cstat.ciub,
  cstat.cilv,
  sd.LP,
  N,
  O,
  Po,
  data,
  slab,
  subset,
  g = NULL,level = 0.95,approx.se.method = 4,
  ...
)
```
Arguments

Details

The c-statistic is a measure of discrimination, and indicates the ability of a prediction model to distinguish between patients developing and not developing the outcome. The c-statistic typically ranges from 0.5 (no discriminative ability) to 1 (perfect discriminative ability).

By default, the function ccalc will derive the c-statistic of each study, together with the corresponding standard error and 95% confidence interval. However, it is also possible to calculate transformed versions of the c-statistic. Appropriate standard errors are then derived using the Delta method. For instance, the logit transformation can be applied by specifying $g = "log(cstat/(1-cstat))"$.

Restoring the c-statistic: For studies where the c-statistic is missing, it is estimated from the standard deviation of the linear predictor (theta.source="std.dev(LP)"). The corresponding method is described by White et al. (2015).

Restoring the standard error of the c-statistic: When missing, the standard error of the cstatistic can be estimated from the confidence interval. Alternatively, the standard error can be approximated from a combination of the reported c-statistic, the total sample size and the total number of events (Newcombe, 2006). This can be achieved by adopting (a modification of) the method proposed by Hanley and McNeil, as specified in approx.se.method.

Value

An object of class c("mm_perf","data.frame") with the following columns:

"theta" The (transformed) c-statistics.

"theta.se" Standard errors of the (transformed) c-statistics.

- "theta.cilb" Lower confidence interval of the (transformed) c-statistics. The level is specified in level. Intervals are calculated on the same scale as theta by assuming a Normal distribution.
- "theta.ciub" Upper confidence interval of the (transformed) c-statistics. The level is specified in level. Intervals are calculated on the same scale as theta by assuming a Normal distribution.

"theta.source" Method used for calculating the (transformed) c-statistic.

"theta.se.source" Method used for calculating the standard error of the (transformed) c-statistic.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

References

Debray TPA, Damen JAAG, Snell KIE, Ensor J, Hooft L, Reitsma JB, et al. A guide to systematic review and meta-analysis of prediction model performance. BMJ. 2017;356:i6460.

Debray TPA, Damen JAAG, Riley R, Snell KIE, Reitsma JB, Hooft L, et al. A framework for metaanalysis of prediction model studies with binary and time-to-event outcomes. Stat Methods Med Res. 2018; In press.

Hanley JA, McNeil BJ. The meaning and use of the area under a receiver operating characteristic (ROC) curve. *Radiology*. 1982; 143(1):29–36.

Newcombe RG. Confidence intervals for an effect size measure based on the Mann-Whitney statistic. Part 2: asymptotic methods and evaluation. *Stat Med*. 2006; 25(4):559–73.

Snell KI, Ensor J, Debray TP, Moons KG, Riley RD. Meta-analysis of prediction model performance across multiple studies: Which scale helps ensure between-study normality for the C statistic and calibration measures? *Statistical Methods in Medical Research*. 2017.

White IR, Rapsomaniki E, the Emerging Risk Factors Collaboration. Covariate-adjusted measures of discrimination for survival data. *Biom J*. 2015;57(4):592–613.

Examples

```
######### Validation of prediction models with a binary outcome #########
data(EuroSCORE)
# Calculate the c-statistic and its standard error
est1 <- ccalc(cstat = c.index, cstat.se = se.c.index, cstat.cilb = c.index.95CIl,
           cstat.ciub = c.index.95CIu, N = n, 0 = n.events, data = EuroSCORE, slab = Study)est1
# Calculate the logit c-statistic and its standard error
est2 <- ccalc(cstat = c.index, cstat.se = se.c.index, cstat.cilb = c.index.95CIl,
           cstat.ciub = c.index.95CIu, N = n, O = n.events, data = EuroSCORE, slab = Study,
              g = "log(cstat/(1-cstat))")est2
# Display the results of all studies in a forest plot
plot(est1)
```


Description

A meta-analysis of nine clinical trials investigating the effect of taking diuretics during pregnancy on the risk of pre-eclampsia.

Usage

data(Collins)

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Format

A data frame with 9 observations on the following 2 variables.

logOR a numeric vector with treatment effect sizes (log odds ratio)

SE a numeric vector with the standard error of the treatment effect sizes

Source

Collins, R., Yusuf, S., Peto, R. Overview of randomised trials of diuretics in pregnancy. *British Medical Journal* 1985, 290, 17–23.

Hardy, R.J. Thompson, S.G. A likelihood approach to meta-analysis with random effects. *Statistics in Medicine* 1996; 15:619–629.

Examples

data(Collins)

cor2cov *Convert a correlation matrix into a covariance matrix*

Description

Convert a correlation matrix into a covariance matrix

Usage

cor2cov(sigma, cormat)

Arguments

Value

The covariance matrix

Author(s)

Thomas Debray <thomas.debray@gmail.com>

Data frame with treatment differences in CD4 cell count.

Usage

```
data("Daniels")
```
Format

A data frame with 15 observations on the following 2 variables.

- Y1 Treatment differences for the log hazard ratio for the development of AIDS or death over 2 years.
- vars1 Error variances of Y1.
- Y2 Difference in mean change in CD4 cell count between baseline and 6 month for studies of the AIDS Clinical Trial Group

vars2 Error variances of Y2.

Details

The Daniels data comprises 15 phase II/III randomized clinical trials of the HIV Disease Section of the Adult AIDS Clinical Trials Group of the National Institutes of Health, which had data available as of May 1996, which had at least six months of follow-up on some patients and in which at least one patient developed AIDS or died. The data were previously used by Daniels and Hughes (1997) to assess whether the change in CD4 cell count is a surrogate for time to either development of AIDS or death in drug trials of patients with HIV.

Source

Daniels MJ, Hughes MD. Meta-analysis for the evaluation of potential surrogate markers. *Statistics in Medicine* 1997; 16: 1965–1982.

dplot *Posterior distribution of estimated model parameters*

Description

Generate a plot of the posterior distribution

Usage

dplot(...)

dplot.mcmc.list 13

Arguments

... Additional arguments, which are currently ignored.

Details

This is a generic function.

Value

A ggplot object.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

dplot.mcmc.list *Posterior distribution of estimated model parameters*

Description

Generate a plot of the posterior distribution

Usage

```
## S3 method for class 'mcmc.list'
dplot(x, P, plot_type = "dens", ...)
```
Arguments

Value

A ggplot object.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

Function to generate plots for the prior and posterior distribution of a Bayesian meta-analysis.

Usage

```
## S3 method for class 'uvmeta'
dplot(x, par, distr_type, plot_type = "dens", ...)
```
Arguments

Value

An object of class ggplot

Author(s)

Thomas Debray <thomas.debray@gmail.com>

Examples

```
## Not run:
data(Roberts)
fit <- with(Roberts, uvmeta(r=SDM, r.se=SE, method="BAYES"))
dplot(fit)
dplot(fit, distr_type = "posterior")
dplot(fit, par = "tau", distr_type = "prior")
dplot(fit, plot_type = "hist")
## End(Not run)
```


Function to generate plots for the prior and posterior distribution of a Bayesian meta-analysis.

Usage

S3 method for class 'valmeta' $dplot(x, par, distr_type, plot_type = "dens", ...)$

Arguments

Value

A ggplot object.

An object of class ggplot

Author(s)

Thomas Debray <thomas.debray@gmail.com>

Examples

```
## Not run:
data(EuroSCORE)
# Meta-analysis of the concordance statistic
fit <- valmeta(cstat=c.index, cstat.se=se.c.index, cstat.cilb=c.index.95CIl,
               cstat.ciub=c.index.95CIu, N=n, O=n.events,
               data=EuroSCORE, method="BAYES", slab=Study)
dplot(fit)
dplot(fit, distr_type = "posterior")
dplot(fit, par = "tau", distr_type = "prior")
```
Meta-analysis of the O:E ratio

```
EuroSCORE.new <- EuroSCORE
EuroSCORE.new$n[c(1, 2, 5, 10, 20)] <- NA
pars <- list(hp.tau.dist="dhalft", # Prior for the between-study standard deviation
            hp.tau.sigma=1.5, # Standard deviation for 'hp.tau.dist'
            hp.tau.df=3, # Degrees of freedom for 'hp.tau.dist'
         hp.tau.max=10) # Maximum value for the between-study standard deviation
fit2 <- valmeta(measure="OE", O=n.events, E=e.events, N=n, data=EuroSCORE.new,
               method="BAYES", slab=Study, pars=pars)
dplot(fit2, plot_type = "hist")
## End(Not run)
```


DVTipd *Hypothetical dataset for diagnosis of Deep Vein Thrombosis (DVT)*

Description

A hypothetical dataset with 500 subjects suspected of having deep vein thrombosis (DVT).

Usage

data(DVTipd)

Format

A data frame with 500 observations of 16 variables.

sex gender (0=female, 1=male) malign active malignancy (0=no active malignancy, 1=active malignancy) par paresis (0=no paresis, 1=paresis) surg recent surgery or bedridden tend tenderness venous system oachst oral contraceptives or hst leg entire leg swollen notraum absence of leg trauma calfdif3 calf difference >= 3 cm pit pitting edema vein vein distension altdiagn alternative diagnosis present histdvt history of previous DVT ddimdich dichotimized D-dimer value dvt final diagnosis of DVT study study indicator

DVTmodels and the set of the set of

Details

Hypothetical dataset derived from the Individual Participant Data Meta-Analysis from Geersing *et al* (2014). The dataset consists of consecutive outpatients with suspected deep vein thrombosis, with documented information on the presence or absence of proximal deep vein thrombosis (dvt) by an acceptable reference test. Acceptable such tests were either compression ultrasonography or venography at initial presentation, or, if venous imaging was not performed, an uneventful followup for at least three months.

Source

Geersing GJ, Zuithoff NPA, Kearon C, Anderson DR, Ten Cate-Hoek AJ, Elf JL, et al. Exclusion of deep vein thrombosis using the Wells rule in clinically important subgroups: individual patient data meta-analysis. *BMJ*. 2014;348:g1340.

Examples

```
data(DVTipd)
str(DVTipd)
summary(apply(DVTipd,2,as.factor))
```

```
## Develop a prediction model to predict presence of DVT
model.dvt <- glm("dvt~sex+oachst+malign+surg+notraum+vein+calfdif3+ddimdich",
                  family=binomial, data=DVTipd)
```

```
summary(model.dvt)
```
DVTmodels *Risk prediction models for diagnosing Deep Venous Thrombosis (DVT)*

Description

Previously published prediction models for predicting the presence of DVT.

Usage

```
data(DVTmodels)
```
Format

An object of the class litmodels with the following information for each literature model: the study-level descriptives ("descriptives"), the regression coefficient or weight for each predictor ("weights") and the error variance for each regression coefficient or weight ("weights.var").

Details

Previously, several models (Gagne, Oudega) and score charts (Wells, modified Wells, and Hamilton) have been published for evaluating the presence of DVT in suspected patients. These models combine information on mulitple predictors into a weighted sum, that can subsequently be used to obtain estimates of absolute risk. See DVTipd for more information on the predictors.

Source

Wells PS, Anderson DR, Bormanis J, Guy F, Mitchell M, Gray L, Clement C, Robinson KS, Lewandowski B. Value of assessment of pretest probability of deep-vein thrombosis in clinical management. *Lancet* 1997; 350(9094):1795–1798. DOI: 10.1016/S0140-6736(97)08140-3.

Wells PS, Anderson DR, Rodger M, Forgie M, Kearon C, Dreyer J, Kovacs G, Mitchell M, Lewandowski B, Kovacs MJ. Evaluation of D-dimer in the diagnosis of suspected deep-vein thrombosis. *The New England Journal of Medicine* 2003; 349(13):1227–1235. DOI: 10.1056/NEJMoa023153.

Gagne P, Simon L, Le Pape F, Bressollette L, Mottier D, Le Gal G. Clinical prediction rule for diagnosing deep vein thrombosis in primary care. *La Presse Medicale* 2009; 38(4):525–533. DOI: 10.1016/j.lpm.2008.09.022.

Subramaniam RM, Snyder B, Heath R, Tawse F, Sleigh J. Diagnosis of lower limb deep venous thrombosis in emergency department patients: performance of Hamilton and modified Wells scores. *Annals of Emergency Medicine* 2006; 48(6):678–685. DOI: 10.1016/j.annemergmed.2006.04.010.

Oudega R, Moons KGM, Hoes AW. Ruling out deep venous thrombosis in primary care. a simple diagnostic algorithm including D-dimer testing. *Thrombosis and Haemostasis* 2005; 94(1):200– 205. DOI: 10.1160/TH04-12-0829.

References

Debray TPA, Koffijberg H, Nieboer D, Vergouwe Y, Steyerberg EW, Moons KGM. Meta-analysis and aggregation of multiple published prediction models. *Stat Med*. 2014 Jun 30;33(14):2341–62.

Debray TPA, Koffijberg H, Vergouwe Y, Moons KGM, Steyerberg EW. Aggregating published prediction models with individual participant data: a comparison of different approaches. *Stat Med*. 2012 Oct 15;31(23):2697–712.

See Also

[DVTipd](#page-15-1)

Examples

data(DVTmodels)

EuroSCORE *Predictive performance of EuroSCORE II*

Description

This data set contains estimates on the predictive performance of the European system for cardiac operative risk evaluation (EuroSCORE II) in patients undergoing cardiac surgery. Results are based on the original development study and 22 validations identified by Guida *et al*.

Usage

data("EuroSCORE")

EuroSCORE 19

Format

A data frame with 23 observations on the following 13 variables.

Study a vector with the first author of each validation study

- n a numeric vector with the total number of patients on which performance estimates are based
- n.events a numeric vector with the total number of observed events
- c.index a numeric vector with the estimated concordance statistic of each validation
- se.c.index a numeric vector with the standard error of the concordance statistics
- c.index.95CIl a numeric vector with the lower bound of the 95% confidence interval of the estimated concordance statistics
- c.index.95CIu a numeric vector with the upper bound of the 95% confidence interval of the estimated concordance statistics
- Po a numeric vector with the overall observed event probability of each validation
- Pe a numeric vector with the overall expected event probability of each validation
- SD.Pe a numeric vector with the standard error of Pe
- e.events a numeric vector with the total number of expected events in each validation
- multicentre a logical vector describing whether the study was a multicentre study

mean.age a numeric vector describing the mean age of the patients

- sd.age a numeric vector with the spread of the age of the patients
- pts.before.2010 a logical vector describing whether studies included patients before 2010 (i.e., before EuroSCORE II was developed)

Details

Published in 2012, EuroSCORE II was developed using logistic regression in a dataset comprising 16,828 adult patients undergoing major cardiac surgery from 154 hospitals in 43 countries over a 12-week period (May-July) in 2010. EuroSCORE II was developed to predict in-hospital mortality for patients undergoing any type of cardiac surgery. In 2014, a systematic review of published evidence on the performance value of the euroSCORE II was undertaken by Guida *et al*. Twenty-two validations, including more 145,592 patients from 21 external validation articles (one study included two validations) and a split-sample validation contained within original development article were included in the review; 23 validation studies in total.

Source

Guida P, Mastro F, Scrascia G, Whitlock R, Paparella D. Performance of the European System for Cardiac Operative Risk Evaluation II: a meta-analysis of 22 studies involving 145,592 cardiac surgery procedures. *J Thorac Cardiovasc Surg*. 2014; 148(6):3049–3057.e1.

Nashef SAM, Roques F, Sharples LD, Nilsson J, Smith C, Goldstone AR, et al. EuroSCORE II. *Eur J Cardiothorac Surg*. 2012; 41(4):734-744; discussion 744-745.

Examples

data(EuroSCORE)

The presence of small-study effects is a common threat to systematic reviews and meta-analyses, especially when it is due to publication bias, which occurs when small primary studies are more likely to be reported (published) if their findings were positive. The presence of small-study effects can be verified by visual inspection of the funnel plot, where for each included study of the metaanalysis, the estimate of the reported effect size is depicted against a measure of precision or sample size. The premise is that the scatter of plots should reflect a funnel shape, if small-study effects do not exist. However, when small studies are predominately in one direction (usually the direction of larger effect sizes), asymmetry will ensue.

The [fat](#page-19-1) function implements several tests for detecting funnel plot asymmetry, which can be used when the presence of between-study heterogeneity in treatment effect is relatively low.

Usage

fat(b, b.se, n.total, d.total, d1, d2, method = $"E-FIV"$)

Arguments

Details

Egger regression method: A common approach to test the presence of small-study effects is to estimate a regression model where the standardized effect estimate (effect/SE) is regressed on a measure of precision (1/SE), (method="E-UW", Egger 1997). It is possible to allow for betweenstudy heterogeneity by adopting a multiplicative overdispersion parameter by which the variance in each study is multiplied (method="E-FIV", Sterne 2000).

Unfortunately, it has been demonstrated that the aforementioned two tests are biased because: (i) the independent variable is subject to sampling variability; (ii) the standardized treatment effect is

correlated with its estimated precision; and (iii) for binary data, the independent regression variable is a biased estimate of the true precision, with larger bias for smaller sample sizes (Macaskill et al. 2001).

Macaskill regression method: The standard approach estimates a regression model with the effect size as a function of the study size (method="M-FIV", Macaskill et al. 2001). Each study is weighted by the precision of the treatment effect estimate to allow for possible heteroscedasticity. An alternative approach is to weight each study by a pooled' estimate of the outcome proportion (method="M-FPV")

For studies with zero events, a continuity correction is applied by adding 0.5 to all cell counts.

Peters regression method: This approach (method="P-FPV") estimates a regression model with the treatment effect as a function of the inverse of the total sample size (Peters et al. 2006). For studies with zero events, a continuity correction is applied by adding 0.5 to all cell counts.

Debray regression method: This approach was proposed for survival data, and uses the total number of events as independent variable in the weighted regression model (Debray et al. 2017). The study weights are based on the inverse variance (method="D-FIV") or on an approximation thereof (method="D-FAV").

Value

a list containing the following entries:

- "pval" A two-sided P-value indicating statistical significance of the funnel plot asymettry test. Values below the significance level (usually defined as 10%) support the presence of funnel plot asymmetry, and thus small-study effects.
- "model" A fitted glm object, representing the estimated regression model used for testing funnel plot asymmetry.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

References

Debray TPA, Moons KGM, Riley RD. Detecting small-study effects and funnel plot asymmetry in meta-analysis of survival data: a comparison of new and existing tests. Res Syn Meth. 2018;9(1):41–50.

Egger M, Davey Smith G, Schneider M, Minder C. Bias in meta-analysis detected by a simple, graphical test. *BMJ*. 1997;315(7109):629–34.

Macaskill P, Walter SD, Irwig L. A comparison of methods to detect publication bias in metaanalysis. *Stat Med*. 2001;20(4):641–54.

Peters JL, Sutton AJ, Jones DR, Abrams KR, Rushton L. Comparison of two methods to detect publication bias in meta-analysis. *JAMA*. 2006 Feb 8;295(6):676–80.

Sterne JA, Gavaghan D, Egger M. Publication and related bias in meta-analysis: power of statistical tests and prevalence in the literature. *J Clin Epidemiol*. 2000;53(11):1119–29.

See Also

[plot.fat](#page-45-1)

Examples

```
data(Fibrinogen)
b <- log(Fibrinogen$HR)
b.se <- ((log(Fibrinogen$HR.975) - log(Fibrinogen$HR.025))/(2*qnorm(0.975)))
n.total <- Fibrinogen$N.total
d.total <- Fibrinogen$N.events
fat(b=b, b.se=b.se)
fat(b=b, b.se=b.se, d.total=d.total, method="D-FIV")
# Note that many tests are also available via metafor
require(metafor)
fat(b=b, b.se=b.se, n.total=n.total, method="M-FIV")
regtest(x=b, sei=b.se, ni=n.total, model="lm", predictor="ni")
```
Fibrinogen *Meta-analysis of the association between plasma fibrinogen concentration and the risk of coronary heath disease*

Description

The Fibrinogen data set is a meta-analysis of 31 studies in which the association between plasma fibrinogen concentration and the risk of coronary heath disease (CHD) was estimated.

Usage

data("Fibrinogen")

Format

A data frame with 5 variables:

- N.total a numeric vector describing the total number of patients for each study
- N.events a numeric vector describing the number of observed events within each study
- HR a numeric vector describing the estimated hazard ratio of each study
- HR.025 a numeric vector describing the lower boundary of the 95% confidence interval of HR
- HR.975 a numeric vector describing the upper boundary of the 95% confidence interval of HR

fitted.metapred 23

Source

Fibrinogen Studies Collaboration. Collaborative meta-analysis of prospective studies of plasma fibrinogen and cardiovascular disease. *Eur J Cardiovasc Prev Rehabil*. 2004 Feb;11(1):9-17.

Thompson S, Kaptoge S, White I, Wood A, Perry P, Danesh J, et al. Statistical methods for the time-to-event analysis of individual participant data from multiple epidemiological studies. *Int J Epidemiol*. 2010 Oct;39(5):1345-59.

Examples

```
data(Fibrinogen)
## maybe str(Fibrinogen) ; plot(Fibrinogen) ...
```
fitted.metapred *Extract Model Fitted Values*

Description

Extract the fitted values of a metapred object. By default returns fitted values of the model in the cross-validation procedure.

Usage

```
## S3 method for class 'metapred'
fitted(
 object,
 select = "cv",step = NULL,
 model = NULL,as.stratified = TRUE,
  type = "response",
  ...
)
```
Arguments

Details

Function still under development, use with caution.

Only returns type = "response".

Author(s)

Valentijn de Jong

forest *Forest plot*

Description

Generate a forest plot by specifying the various effect sizes, confidence intervals and summary estimate.

Usage

forest(...)

Arguments

... Additional arguments, which are currently ignored.

Details

This is a generic function. See [forest.default](#page-24-1) for making forest plots of summary statistics, [for](#page-26-1)[est.metapred](#page-26-1) for plotting [metapred](#page-38-1) objects, and [forest.mp.cv.val](#page-27-1) for plotting mp.cv.val objects.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

Valentijn de Jong <Valentijn.M.T.de.Jong@gmail.com>

Generate a forest plot by specifying the various effect sizes, confidence intervals and summary estimate.

Usage

```
## Default S3 method:
forest(
  theta,
  theta.ci.lb,
  theta.ci.ub,
  theta.slab,
  theta.summary,
  theta.summary.ci.lb,
  theta.summary.ci.ub,
  theta.summary.pi.lb,
  theta.summary.pi.ub,
  title,
  sort = "asc",theme = thene_bw(),
  predint.linetype = 1,
  xlim,
 xlab = ",
  refline = 0,
  label.summary = "Summary Estimate",
  label.predint = "Prediction Interval",
  nrows.before.summary = 1,
  study.digits = 2,
  study.shape = 15,
  col.diamond = "white",
  col.predint = "black",
  size.study = 0.5,
  size.predint = 1,
  lty.ref = "dotted",
  ...
```
)

Arguments

Value

An object of class ggplot

forest.metapred 27

Author(s)

Thomas Debray <thomas.debray@gmail.com>

forest.metapred *Forest plot of a metapred fit*

Description

Draw a forest plot of the performance of an internally-externally cross-validated model. By default the final model is shown.

Usage

```
## S3 method for class 'metapred'
forest(x, perfFUN = 1, step = NULL, method = "REML", model = NULL, ...)
```
Arguments

Author(s)

Valentijn de Jong <Valentijn.M.T.de.Jong@gmail.com>

Examples

data(DVTipd)

```
# Internal-external cross-validation of a pre-specified model 'f'
f <- dvt ~ histdvt + ddimdich + sex + notraum
fit <- metapred(DVTipd, strata = "study", formula = f, scope = f, family = binomial)
```
Display the model's external performance (expressed as mean squared error by default)

28 Framingham

```
# for each study
forest(fit)
```
forest.mp.cv.val *Forest plot of a validation object.*

Description

Draw a forest plot of the performance of an internally-externally cross-validated model.

Usage

```
## S3 method for class 'mp.cv.val'
forest(x, perfFUN = 1, method = "REML", xlab = NULL, ...)
```
Arguments

Author(s)

Valentijn de Jong <Valentijn.M.T.de.Jong@gmail.com>

Description

This data set contains estimates on the performance of the Framingham model for predicting coronary heart disease in male populations (Wilson 1998). Results are based on the original development study and 20 validations identified by Damen *et al* (BMC Med, 2017).

Framingham 29

Usage

data("Framingham")

Format

A data frame with 24 observations on the following 19 variables.

AuthorYear a vector describing the study authors

- n a numeric vector with the total number of patients on which performance estimates are based
- n.events a numeric vector with the total number of observed events
- c.index a numeric vector with the estimated concordance statistic of each validation

se.c.index a numeric vector with the standard error of the concordance statistics

- c.index.95CIl a numeric vector with the lower bound of the 95% confidence interval of the estimated concordance statistics
- c.index.95CIu a numeric vector with the upper bound of the 95% confidence interval of the estimated concordance statistics
- Po a numeric vector with the overall observed event probability of each validation
- Pe a numeric vector with the overall expected event probability of each validation
- t.val a numeric vector describing the time period in which predictive performance was assessed for each validation
- mean_age a numeric vector describing the mean age of the patients
- sd_age a numeric vector with the spread of the age of the patients
- mean_SBP a numeric vector with the mean systolic blood pressure in the validation studies (mm Hg)
- sd_SBP a numeric vector with the spread of systolic blood pressure in the validation studies
- mean_total_cholesterol a numeric vector with the mean total cholesterol in the validation studies (mg/dL)
- sd_total_cholesterol a numeric vector with the spread of total cholesterol in the validation studies
- mean_hdl_cholesterol a numeric vector with the mean high-density lipoprotein cholesterol in the validation studies (mg/dL)
- sd_hdl_cholesterol a numeric vector with the spread of high-density lipoprotein cholesterol in the validation studies
- pct_smoker a numeric vector with the percentage smokers in the validation studies

Details

The Framingham Risk Score allows physicians to predict 10-year coronary heart disease (CHD) risk in patients without overt CHD. It was developed in 1998 from a middle-aged white population sample, and has subsequently been validated across different populations. The current dataset contains the original (internal validation) results, as well as 23 external validations which were identified through a systematic review. In this review, studies were eligible for inclusion if they described the validation of the original Framingham model and assessed its performance for fatal or nonfatal CHD in males from a general population setting.

Source

Damen JAAG, Hooft L, Schuit E, Debray TPA, Collins GS, Tzoulaki I, et al. Prediction models for cardiovascular disease risk in the general population: systematic review. *BMJ*. 2016;i2416.

Damen JAAG, Pajouheshnia R, Heus P, Moons KGM, Reitsma JB, Scholten RJPM, et al. Performance of the Framingham risk models and Pooled Cohort Equations: a systematic review and meta-analysis. *BMC Med*. 2017;17(1):109.

Wilson PW, D'Agostino RB, Levy D, Belanger AM, Silbershatz H, Kannel WB. Prediction of coronary heart disease using risk factor categories. *Circulation*. 1998; 97(18):1837–47.

Examples

data(Framingham)

gelmanplot *Gelman-Rubin-Brooks plot*

Description

This plot shows the evolution of Gelman and Rubin's shrink factor as the number of iterations increases. The code is adapted from the R package coda.

Usage

gelmanplot(...)

Arguments

... Additional arguments which are currently not used

Value

A ggplot object.

An object of class ggplot

Author(s)

Thomas Debray <thomas.debray@gmail.com>

This plot shows the evolution of Gelman and Rubin's shrink factor as the number of iterations increases. The code is adapted from the R package coda.

Usage

```
## S3 method for class 'mcmc.list'
gelmanplot(
  x,
  P,
  confidence = 0.95,
  max.bins = 50,
  autoburnin = TRUE,
  greek = FALSE,
  ...
\overline{\phantom{a}}
```
Arguments

Value

A ggplot object.

An object of class ggplot

Author(s)

Thomas Debray <thomas.debray@gmail.com>

gelmanplot.uvmeta *Gelman-Rubin-Brooks plot*

Description

This plot shows the evolution of Gelman and Rubin's shrink factor as the number of iterations increases. The code is adapted from the R package coda.

Usage

```
## S3 method for class 'uvmeta'
gelmanplot(x, confidence = 0.95, ...)
```
Arguments

Value

A ggplot object.

An object of class ggplot

Author(s)

Thomas Debray <thomas.debray@gmail.com>

gelmanplot.valmeta *Gelman-Rubin-Brooks plot*

Description

This plot shows the evolution of Gelman and Rubin's shrink factor as the number of iterations increases. The code is adapted from the R package coda.

Usage

```
## S3 method for class 'valmeta'
gelmanplot(x, confidence = 0.95, ...)
```
gen 33 and 33

Arguments

Value

A ggplot object. An object of class ggplot

Author(s)

Thomas Debray <thomas.debray@gmail.com>

Examples

```
## Not run:
data(EuroSCORE)
# Meta-analysis of the concordance statistic
fit <- valmeta(cstat=c.index, cstat.se=se.c.index, cstat.cilb=c.index.95CIl,
               cstat.ciub=c.index.95CIu, N=n, O=n.events,
               data=EuroSCORE, method="BAYES", slab=Study)
gelmanplot(fit)
## End(Not run)
```
gen *Generalizability estimates*

Description

Obtain generalizability estimates from a model fit.

Usage

```
gen(object, ...)
generalizability(object, ...)
```
Arguments

object A model fit object, either a [metapred](#page-38-1) or subset(metapred) object. ... By default, the final model is selected. This parameter allows other arguments to be passed to [subset.metapred](#page-61-1) such that the generalizability estimates of other steps/models may be returned..

Details

With named values or indices for parameter genFUN one or more estimates of generalizability can be selected. Use genFUN = θ to select all.

Author(s)

Valentijn de Jong

impact *IMPACT data*

Description

The IMPACT dataset comprises 15 studies of patients suffering from traumatic brain injury, including individual patient data from 11 randomized controlled trials and four observational studies.

Usage

data("impact")

Format

A data frame with 11022 observations on the following 11 variables.

name Name of the study type Type of study, RCT: randomized controlled trial,OBS: observational cohort age Age of the patient motor_score Glasgow Coma Scale motor score pupil Pupillary reactivity ct Marshall Computerized Tomography classification hypox Hypoxia (0=no, 1=yes) hypots Hypotension (0=no, 1=yes) tsah Traumatic subarachnoid hemorrhage (0=no, 1=yes) edh Epidural hematoma (0=no, 1=yes) mort 6-month mortality (0=alive, 1=dead)

Details

The included studies were part of the IMPACT project, where a total of 25 prognostic factors were considered for prediction of 6-month mortality. Missing values were imputed using the study as a fixed effect in the imputation model (Steyerberg et al, 2008).

Source

Steyerberg EW, Nieboer D, Debray TPA, Van Houwelingen JC. Assessment of heterogeneity in an individual participant data meta-analysis of prediction models: An overview and illustration. *Stat Med*. 2019;38(22):4290–309.

References

Murray GD, Butcher I, McHugh GS, et al. Multivariable prognostic analysis in traumatic brain injury: results from the IMPACT study. *J Neurotrauma*. 2007;24(2):329–337.

Steyerberg EW, Mushkudiani N, Perel P, et al. Predicting outcome after traumatic brain injury: development and international validation of prognostic scores based on admission characteristics. *PLOS Med*. 2008;5(8):e165.

Examples

data(impact) by(impact, impact\$name, summary) # Plot the distribution of age by study library(ggplot2) $e \leftarrow \text{gplot}(\text{impact}, \text{aes}(x = \text{name}, y = \text{age}))$ $e + geom_violin(aes(fill = type), trim = FALSE) +$ theme(axis.text.x = element_text(angle = 45)) + xlab("Study")

impute_conditional_mean

Impute missing values by their conditional mean

Description

This function imputes missing values by their conditional mean

Usage

```
impute_conditional_mean(x, mu, Sigma)
```
Arguments

Value

A vector where missing values for 'x' have been replaced by their conditional mean

Author(s)

Thomas Debray <thomas.debray@gmail.com>

Examples

```
# Define the population means
mu \leftarrow c(0, 1, 2)# Define the covariance of the population
Sigma \leftarrow diag(1,3)Sigma[1,2] <- Sigma[2,1] <- 0.3
Sigma[2,3] <- Sigma[3,2] <- 0.1
Sigma[1,3] <- Sigma[3,1] <- -0.2
# Generate a 'random' sample from the population that is partially observed
x \leq -c(NA, 2, 4)# Impute the missing values
impute_conditional_mean (x=x, mu=mu, Sigma=Sigma)
```
inv.logit *Apply the inverse logit tranformation*

Description

Transforms a linear predictor into a probability.

Usage

inv.logit(x)

Arguments

x A vector of numerics (between -Inf and Inf)

Value

A vector of numerics between 0 and 1.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

See Also

[logit](#page-36-1)
Description

Data frame with diagnostic accuracy data from exercise electrocardiography.

Usage

```
data("Kertai")
```
Format

One data frame with 4 variables.

- TP integer. number of true positives
- FN integer. number of false negatives
- FP integer. number of false positives
- TN integer. number of true negatives

Details

The Kertai data set is a meta-analysis of prognostic test studies and comprises 7 studies where the diagnostic test accuracy of exercise electrocardiography for predicting cardiac events in patients undergoing major vascular surgery was measured.

Source

Kertai MD, Boersma E, Bax JJ, Heijenbrok-Kal MH, Hunink MGM, L'talien GJ, Roelandt JRTC, van Urk H, Poldermans D. A meta-analysis comparing the prognostic accuracy of six diagnostic tests for predicting perioperative cardiac risk in patients undergoing major vascular surgery. *Heart* 2003; 89: 1327–1334.

Jackson D, Riley RD, & White IW. Multivariate meta-analysis: Potential and promise. *Statistics in Medicine* 2010; 30: 2481–2498.

logit *Apply logit tranformation*

Description

Transforms values between 0 and 1 to values between -Inf and Inf.

Usage

logit(x)

Arguments

Value

A vector of numerics (between -Inf and Inf).

Author(s)

Thomas Debray <thomas.debray@gmail.com>

See Also

[inv.logit](#page-35-0)

logLik.riley *Print the log-likelihood*

Description

This function provides the (restricted) log-likelihood of a fitted model.

Usage

```
## S3 method for class 'riley'
logLik(object, ...)
```
Arguments

Value

Returns an object of class logLik. This is the (restricted) log-likelihood of the model represented by object evaluated at the estimated coefficients. It contains at least one attribute, "df" (degrees of freedom), giving the number of (estimated) parameters in the model.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

References

Riley RD, Thompson JR, Abrams KR. An alternative model for bivariate random-effects metaanalysis when the within-study correlations are unknown. *Biostatistics* 2008; 9: 172–186.

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Examples

```
data(Daniels)
fit <- riley(Daniels,control=list(maxit=10000))
logLik(fit)
```
ma *Random effects meta-analysis*

Description

Meta-analysis of the performance or coefficients of a metapred object. Caution: it is still under development.

Usage

ma(object, method, ...)

Arguments

Details

Produces different object types depending on input.

Author(s)

Valentijn de Jong

Description

Generalized stepwise regression for obtaining a prediction model that is validated with (stepwise) internal-external cross-validation, in or to obtain adequate performance across data sets. Requires data from individuals in multiple studies.

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Usage

```
metapred(
  data,
  strata,
  formula,
  estFUN = "glm",scope = NULL,retest = FALSE,
  max.steps = 1000,
  center = FALSE,
  recal.int = FALSE,
  cvFUN = NULL,
  cv.k = NULL,metaFUN = NULL,
  meta.method = NULL,
  predFUN = NULL,
  perfFUN = NULL,
  genFUN = NULL,
  selFUN = "which.min",
  gen.of.perf = "first",
  ...
\mathcal{L}
```

```
Arguments
```


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Details

Use [subset.metapred](#page-61-0) to obtain an individual prediction model from a metapred object.

Note that formula.changes is currently unordered; it does not represent the order of changes in the stepwise procedure.

metapred is still under development, use with care.

Value

A list of class metapred, containing the final model in global.model, and the stepwise tree of estimates of the coefficients, performance measures, generalizability measures in stepwise.

Author(s)

Valentijn de Jong <Valentijn.M.T.de.Jong@gmail.com>

References

Debray TPA, Moons KGM, Ahmed I, Koffijberg H, Riley RD. A framework for developing, implementing, and evaluating clinical prediction models in an individual participant data meta-analysis. *Stat Med*. 2013;32(18):3158-80.

de Jong VMT, Moons KGM, Eijkemans MJC, Riley RD, Debray TPA. Developing more generalizable prediction models from pooled studies and large clustered data sets. *Stat Med*. 2021;40(15):3533– 59.

Riley RD, Tierney JF, Stewart LA. Individual participant data meta-analysis: a handbook for healthcare research. Hoboken, NJ: Wiley; 2021. ISBN: 978-1-119-33372-2.

Schmid CH, Stijnen T, White IR. Handbook of meta-analysis. First edition. Boca Raton: Taylor and Francis; 2020. ISBN: 978-1-315-11940-3.

See Also

[forest.metapred](#page-26-0) for generating a forest plot of prediction model performance

Examples

```
data(DVTipd)
## Not run:
# Explore heterogeneity in intercept and assocation of 'ddimdich'
glmer(dvt \sim 0 + cluster + (ddimdich|study), family = binomial(), data = DVTipd)
## End(Not run)
# Scope
f \le - dvt \sim histdvt + ddimdich + sex + notraum
# Internal-external cross-validation of a pre-specified model 'f'
fit <- metapred(DVTipd, strata = "study", formula = f, scope = f, family = binomial)
fit
# Let's try to simplify model 'f' in order to improve its external validity
metapped(DVTipd, strata = "study", formula = f, family = binomial)# We can also try to build a generalizable model from scratch
## Not run:
# Some additional examples:
metapred(DVTipd, strata = "study", formula = dvt \sim 1, scope = f, family = binomial) # Forwards
```
oecalc 43

```
metapred(DVTipd, strata = "study", formula = f, scope = f, family = binomial) # no selection
metapred(DVTipd, strata = "study", formula = f, max.steps = 0, family = binomial) # no selection
metapred(DVTipd, strata = "study", formula = f, recal.int = TRUE, family = binomial)
metapred(DVTipd, strata = "study", formula = f, meta.method = "REML", family = binomial)
## End(Not run)
# By default, metapred assumes the first column is the outcome.
newdat <- data.frame(dvt=0, histdvt=0, ddimdich=0, sex=1, notraum=0)
fitted \leq predict(fit, newdata = newdat)
```
oecalc *Calculate the total O:E ratio*

Description

This function calculates (transformed versions of) the ratio of total number of observed versus expected events with the corresponding sampling variance.

Usage

oecalc(OE, OE.se, OE.cilb, OE.ciub, OE.cilv, EO, EO.se, citl, citl.se, N, O, E, Po, Po.se, Pe, data, slab, $add = 1/2,$ $g = NULL,$ $level = 0.95$

Arguments

)

OE vector with the estimated ratio of total observed versus total expected events

Value

An object of class c("mm_perf","data.frame") with the following columns:

"theta" The (transformed) O:E ratio.

"theta.se" Standard errors of the (transformed) O:E ratio.

- "theta.cilb" Lower confidence interval of the (transformed) O:E ratios. The level is specified in level. Intervals are calculated on the same scale as theta by assuming a Normal distribution.
- "theta.ciub" Upper confidence interval of the (transformed) c-statistics. The level is specified in level. Intervals are calculated on the same scale as theta by assuming a Normal distribution.

"theta.source" Method used for calculating the (transformed) O:E ratio.

"theta.se.source" Method used for calculating the standard error of the (transformed) O:E ratio.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

References

Debray TPA, Damen JAAG, Snell KIE, Ensor J, Hooft L, Reitsma JB, et al. A guide to systematic review and meta-analysis of prediction model performance. *BMJ*. 2017;356:i6460.

Debray TPA, Damen JAAG, Riley R, Snell KIE, Reitsma JB, Hooft L, et al. A framework for metaanalysis of prediction model studies with binary and time-to-event outcomes. *Stat Methods Med Res*. 2019 Sep;28(9):2768–86.

Snell KI, Ensor J, Debray TP, Moons KG, Riley RD. Meta-analysis of prediction model performance across multiple studies: Which scale helps ensure between-study normality for the C statistic and calibration measures? *Stat Methods Med Res*. 2017.

Examples

```
######### Validation of prediction models with a binary outcome #########
data(EuroSCORE)
# Calculate the total O:E ratio and its standard error
est1 <- oecalc(0 = n.events, E = e.events, N = n, data = EuroSCORE, slab = Study)
est1
# Calculate the log of the total O:E ratio and its standard error
est2 <- oecalc(0 = n.events, E = e.events, N = n, data = EuroSCORE, slab = Study, g = "log(OE)")
est2
# Display the results of all studies in a forest plot
plot(est1)
```
perf *Performance estimates*

Description

Obtain performance estimates from a model fit.

Usage

```
perf(object, ...)
performance(object, ...)
```
Arguments

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46 plot.fat

Author(s)

Valentijn de Jong

plot.fat *Display results from the funnel plot asymmetry test*

Description

Generates a funnel plot for a fitted fat object.

Usage

```
## S3 method for class 'fat'
plot(
 x,
 ref,
 confint = TRUE,
 confint.level = 0.1,
 confint.col = "skyblue",
 confint.alpha = 0.5,
 confint.density = NULL,
 xlab = "Effect size",
  add.pval = TRUE,...
\mathcal{L}
```
Arguments

Author(s)

Thomas Debray <thomas.debray@gmail.com>

Frantisek Bartos <f.bartos96@gmail.com>

Examples

```
data(Fibrinogen)
b <- log(Fibrinogen$HR)
b.se <- ((log(Fibrinogen$HR.975) - log(Fibrinogen$HR.025))/(2*qnorm(0.975)))
n.total <- Fibrinogen$N.total
# A very simple funnel plot
plot(fat(b=b, b.se=b.se), xlab = "Log hazard ratio")
# Plot the funnel for an alternative test
plot(fat(b=b, b.se=b.se, n.total=n.total, method="M-FIV"), xlab = "Log hazard ratio")
```
plot.mm_perf *Forest Plots*

Description

Function to create forest plots for objects of class "mm_perf".

Usage

S3 method for class 'mm_perf' $plot(x, \ldots)$

Arguments

Details

The forest plot shows the performance estimates of each study with corresponding confidence intervals.

Value

An object of class ggplot

Author(s)

Thomas Debray <thomas.debray@gmail.com>

References

Lewis S, Clarke M. Forest plots: trying to see the wood and the trees. *BMJ*. 2001; 322(7300):1479– 80.

Examples

data(EuroSCORE)

```
# Calculate the c-statistic and its standard error
est1 <- ccalc(cstat = c.index, cstat.se = se.c.index, cstat.cilb = c.index.95CIl,
           cstat.ciub = c.index.95CIu, N = n, 0 = n.events, data = EuroSCORE, slab = Study)plot(est1)
```

```
# Calculate the total O:E ratio and its standard error
est2 <- oecalc(0 = n.events, E = e.events, N = n, data = EuroSCORE, slab = Study)
plot(est2)
```


Description

Generates a forest plot for each outcome of the bivariate meta-analysis.

Usage

```
## S3 method for class 'riley'
plot(x, title, sort = "asc", xlim, refline, ...)
```
Arguments

plot.uvmeta 49

Author(s)

Thomas Debray <thomas.debray@gmail.com>

References

Riley RD, Thompson JR, Abrams KR. An alternative model for bivariate random-effects metaanalysis when the within-study correlations are unknown. *Biostatistics* 2008; 9: 172–186.

Examples

```
data(Scheidler)
```

```
#Perform the analysis
fit <- riley(Scheidler[which(Scheidler$modality==1),])
plot(fit)
require(ggplot2)
plot(fit, sort="none", theme=theme_gray())
```
plot.uvmeta *Forest Plots*

Description

Function to create forest plots for objects of class "uvmeta".

Usage

```
## S3 method for class 'uvmeta'
plot(x, sort = "asc", ...)
```
Arguments

Details

The forest plot shows the performance estimates of each validation with corresponding confidence intervals. A polygon is added to the bottom of the forest plot, showing the summary estimate based on the model. A 95% prediction interval is added by default for random-effects models, the dotted line indicates its (approximate) bounds

Full lines indicate confidence intervals or credibility intervals (in case of a Bayesian meta-analysis). Dashed lines indicate prediction intervals. The width of all intervals is defined by the significance level chosen during meta-analysis.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

References

- Lewis S, Clarke M. Forest plots: trying to see the wood and the trees. *BMJ*. 2001; 322(7300):1479– 80.
- Riley RD, Higgins JPT, Deeks JJ. Interpretation of random effects meta-analyses. *BMJ*. 2011 342:d549–d549.

Examples

```
data(Roberts)
```

```
# Frequentist random-effects meta-analysis
fit <- with(Roberts, uvmeta(r=SDM, r.se=SE, labels=rownames(Roberts)))
plot(fit)
```
plot.valmeta *Forest Plots*

Description

Function to create forest plots for objects of class "valmeta".

Usage

S3 method for class 'valmeta' $plot(x, \ldots)$

Arguments

Details

The forest plot shows the performance estimates of each validation with corresponding confidence intervals. A polygon is added to the bottom of the forest plot, showing the summary estimate based on the model. A 95% prediction interval is added by default for random-effects models, the dotted line indicates its (approximate) bounds.

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Value

An object of class ggplot

Author(s)

Thomas Debray <thomas.debray@gmail.com>

References

Debray TPA, Damen JAAG, Snell KIE, Ensor J, Hooft L, Reitsma JB, et al. A guide to systematic review and meta-analysis of prediction model performance. *BMJ*. 2017;356:i6460.

Lewis S, Clarke M. Forest plots: trying to see the wood and the trees. *BMJ*. 2001; 322(7300):1479– 80.

Riley RD, Higgins JPT, Deeks JJ. Interpretation of random effects meta-analyses. *BMJ*. 2011 342:d549–d549.

See Also

When a Bayesian meta-analysis was conducted, the prior and posterior distribution can be visualized using [dplot.valmeta](#page-14-0). Further, the running means and the presence of autocorrelation can be inspected using [rmplot.valmeta](#page-57-0) and, respectively, [acplot.valmeta](#page-6-0).

Examples

```
data(EuroSCORE)
fit <- valmeta(cstat=c.index, cstat.se=se.c.index, cstat.cilb=c.index.95CIl,
               cstat.ciub=c.index.95CIu, N=n, O=n.events, data=EuroSCORE)
plot(fit)
library(ggplot2)
plot(fit, theme=theme_grey())
```
predict.riley *Prediction Interval*

Description

Calculates a prediction interval for the summary parameters of Riley's alternative model for bivariate random-effects meta-analysis. This interval predicts in what range future observations will fall given what has already been observed.

Usage

```
## S3 method for class 'riley'
predict(object, ...)
```
52 recalibrate \sim 52 recalibrate \sim 52 recalibrate \sim 52 recalibrate \sim 52 recalibrate

Arguments

Details

Prediction intervals are based on Student's t-distribution with (numstudies - 5) degrees of freedom. The width of the interval is specified by the significance level chosen during meta-analysis.

Value

Data frame containing prediction intervals with the summary estimates beta1 and beta2 (for effect size data), or with the mean sensitivity and false positive rate (for diagnostic test accuracy data).

Author(s)

Thomas Debray <thomas.debray@gmail.com>

recalibrate *Recalibrate a Prediction Model*

Description

recalibrate is used to recalibrate a prediction model of classes metapred, glm or lm.

Usage

```
recalibrate(object, newdata, f = -1, estFUN = NULL, ...)
```
Arguments

Details

Currently only the coefficients are updated and the variances and other aspects are left untouched. For updating the entire model and all its statistics, see [update.](#page-0-0)

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Value

Recalibrated model fit object, of the same class as object. Generally, updated coefficients can be retrieved with coef().

Examples

```
data(DVTipd)
DVTipd$cluster <- 1:4 # Add a fictional clustering to the data set.
# Suppose we estimated the model in three studies:
DVTipd123 <- DVTipd[DVTipd$cluster <= 3, ]
mp <- metamisc:::metapred(DVTipd123, strata = "cluster", f = dvt ~ vein + malign,
family = binomial)
# and now want to recalibrate it for the fourth:
DVTipd4 <- DVTipd[DVTipd$cluster == 4, ]
metamisc:::recalibrate(mp, newdata = DVTipd4)
```
riley *Fit the alternative model for bivariate random-effects meta-analysis*

Description

This function fits the alternative model for bivariate random-effects meta-analysis when the withinstudy correlations are unknown. This bivariate model was proposed by Riley et al. (2008) and is similar to the general bivariate random-effects model (van Houwelingen et al. 2002), but includes an overall correlation parameter rather than separating the (usually unknown) within- and betweenstudy correlation. As a consequence, the alternative model is not fully hierarchical, and estimates of additional variation beyond sampling error (psi) are not directly equivalent to the between-study variation (tau) from the general model. This model is particularly useful when there is large withinstudy variability, few primary studies are available or the general model estimates the between-study correlation as 1 or -1.

Usage

```
riley(X, slab, optimization = "Nelder-Mead", control = list(), pars, ...)
```
Arguments

X data frame containing integer variables Y1, vars1, Y2 and vars2, where the columns Y1 and Y2 represent the effect sizes of outcome 1 and, respectively, outcome 2. The columns vars1 and vars2 represent the error variances of Y1 and, respectively, Y2. Alternatively, when considering a meta-analysis of diagnostic test accuracy data, the columns TP, FN, FP and TN may be specified. Corresponding values then represent the number of true positives, the number of false negatives, the number of false positives and, respectively, the number of true negatives.

```
slab Optional vector specifying the label for each study
```


Details

Parameters are estimated by iteratively maximizing the restriced log-likelihood using the Newton-Raphson procedure. The results from a univariate random-effects meta-analysis with a methodof-moments estimator are used as starting values for beta1, beta2, psi1 and psi2 in the optim command. Standard errors for all parameters are obtained from the inverse Hessian matrix.

Meta-analysis of effect sizes: The following parameters are estimated by iteratively maximizing the restriced log-likelihood using the Newton-Raphson procedure: pooled effect size for outcome 1 (beta1), pooled effect size for outcome 2 (beta2), additional variation of beta1 beyond sampling error (psi1), additional variation of beta2 beyond sampling error (psi2) and the correlation rho between psi1 and psi2.

Meta-analysis of diagnostic test accuracy: Although the model can also be used for diagnostic test accuracy data when substantial within-study correlations are expected, assuming zero withinstudy correlations (i.e. applying Reitsma's approach) is usually justified (Reitsma et al. 2005, Daniels and Hughes 1997, Korn et al. 2005, Thompson et al. 2005, Van Houwelingen et al. 2002).

A logit transformation is applied to the sensitivities ans false positive rates of each study, in order to meet the normality assumptions. When zero cell counts occur, continuity corrections may be required. The correction value can be specified using correction (defaults to 0.5). Further, when the argument correction.control is set to "all" (the default) the continuity correction is added to the whole data if only one cell in one study is zero. If correction.control="single" the correction is only applied to rows of the data which have a zero.

The following parameters are estimated: logit of sensitivity (beta1), logit of false positive rate (beta2), additional variation of beta1 beyond sampling error (psi1), additional variation of beta2 beyond sampling error (psi2) and the correlation (rho) between psi1 and psi2.

Value

An object of the class riley for which many standard methods are available. A warning message is casted when the Hessian matrix contains negative eigenvalues, which implies that the identified solution is a saddle point and thus not optimal.

Note

The overall correlation parameter rho is transformed during estimation to ensure that corresponding values remain between -1 and 1. The transformed correlation rhoT is given as $logit((rho+1)/2)$. During optimization, the starting value for rhoT is set to 0. The standard error of rho is derived from rhoT using the Delta method. Similarly, the Delta methods is used to derive the standard error of the sensitivity and false positive rate from beta1 and, respectively, beta2.

Algorithms for dealing with missing data are currently not implemented, but Bayesian approaches will become available in later versions.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

References

- Korn EL, Albert PS, McShane LM. Assessing surrogates as trial endpoints using mixed models. *Statistics in Medicine* 2005; 24: 163–182.
- Nelder JA, Mead R. A simplex algorithm for function minimization. *Computer Journal* (1965); 7: 308–313.
- Reitsma J, Glas A, Rutjes A, Scholten R, Bossuyt P, Zwinderman A. Bivariate analysis of sensitivity and specificity produces informative summary measures in diagnostic reviews. *Journal of Clinical Epidemiology* 2005; 58: 982–990.
- Riley RD, Thompson JR, Abrams KR. An alternative model for bivariate random-effects metaanalysis when the within-study correlations are unknown. *Biostatistics* 2008; 9: 172–186.
- Thompson JR, Minelli C, Abrams KR, Tobin MD, Riley RD. Meta-analysis of genetic studies using mendelian randomization–a multivariate approach. *Statistics in Medicine* 2005; 24: 2241–2254.
- van Houwelingen HC, Arends LR, Stijnen T. Advanced methods in meta-analysis: multivariate approach and meta-regression. *Statistics in Medicine* 2002; 21: 589–624.

Examples

```
data(Scheidler)
data(Daniels)
data(Kertai)
# Meta-analysis of potential surrogate markers data
# The results obtained by Riley (2008) were as follows:
# beta1 = -0.042 (SE = 0.063),
# beta2 = 14.072 (SE = 4.871)
# rho = -0.759## Not run:
fit1 <- riley(Daniels) #maxit reached, try again with more iterations
## End(Not run)
fit1 <- riley(Daniels, control=list(maxit=10000))
summary(fit1)
# Meta-analysis of prognostic test studies
fit2 <- riley(Kertai)
fit2
```
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```
# Meta-analysis of computed tomography data
ds <- Scheidler[which(Scheidler$modality==1),]
fit3 <- riley(ds)
fit3
```
rmplot *Plot the running means of a Bayesian meta-analysis model*

Description

Function to display running means of a fitted Bayesian meta-analysis model.

Usage

rmplot(...)

Arguments

... Additional arguments, which are currently ignored.

Details

This is a generic function.

Value

A ggplot object.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

rmplot.mcmc.list *Plot the running means of a Bayesian meta-analysis model*

Description

Function to display running means of a fitted Bayesian meta-analysis model.

Usage

```
## S3 method for class 'mcmc.list'
rmplot(x, P, greek = FALSE, ...)
```
rmplot.uvmeta 57

Arguments

Value

A ggplot object.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

Description

Function to display running means of a fitted Bayesian meta-analysis model.

Usage

```
## S3 method for class 'uvmeta'
rmplot(x, \ldots)
```
Arguments

Details

Results are displayed for the estimated mean (mu) and standard-deviation (tau) of the meta-analysis model.

Value

A ggplot object.

An object of class ggplot

Author(s)

Thomas Debray <thomas.debray@gmail.com>

Examples

```
## Not run:
data(Roberts)
```

```
fit <- with(Roberts, uvmeta(r=SDM, r.se=SE, labels=rownames(Roberts), method="BAYES"))
rmplot(fit)
```
End(Not run)

rmplot.valmeta *Plot the running means of a Bayesian meta-analysis model*

Description

Function to display running means of a fitted Bayesian meta-analysis model.

Usage

```
## S3 method for class 'valmeta'
rmplot(x, \ldots)
```
Arguments

Details

Results are displayed for the estimated mean (mu) and standard-deviation (tau) of the meta-analysis model.

Value

A ggplot object.

An object of class ggplot

Author(s)

Thomas Debray <thomas.debray@gmail.com>

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Examples

```
## Not run:
data(EuroSCORE)
fit <- valmeta(cstat=c.index, cstat.se=se.c.index, cstat.cilb=c.index.95CIl,
               cstat.ciub=c.index.95CIu, N=n, O=n.events,
               data=EuroSCORE, method="BAYES", slab=Study)
rmplot(fit)
## End(Not run)
```
Roberts *Roberts data*

Description

Data frame with summary data from 14 comparative studies.

Usage

data("Roberts")

Format

One data frame with 2 variables.

SDM Effect sizes (standardized differences in means)

SE Standard error of the effect sizes

Details

The Roberts data set is a meta-analysis of 14 studies comparing 'set shifting' ability (the ability to move back and forth between different tasks) in people with eating disorders and healthy controls.

Source

Roberts ME, Tchanturia K, Stahl D, Southgate L, Treasure J. A systematic review and meta-analysis of set-shifting ability in eating disorders. *Psychological Medicine* 2007, 37: 1075–1084.

Higgins JPT, Thompson SG, Spiegelhalter DJ. A re-evaluation of random-effects meta-analysis. *Journal of the Royal Statistical Society. Series A (Statistics in Society)* 2009, 172: 137–159.

Description

Data frame with diagnostic accuracy data from three imaging techniques for the diagnosis of lymph node metastasis in women with cervical cancer.

Usage

data("Scheidler")

Format

One data frame with 6 variables.

author string . author of article

modality integer . type of test (1=CT, 2=LAG, 3=MRI)

- TP integer. number of true positives
- FN integer. number of false negatives
- FP integer. number of false positives
- TN integer. number of true negatives

Details

The Scheidler data comprises the results from a meta-analysis where three imaging techniques for the diagnosis of lymph node metastasis in women with cervical cancer are compared. Forty-four studies in total were included: 17 studies evaluated lymphangiography, another 17 studies examined computed tomography and the remaining 10 studies focused on magnetic resonance imaging. Diagnosis of metastatic disease by lymphangiography (LAG) is based on the presence of nodal-filling defects, whereas computed tomography (CT) and magnetic resonance imaging (MRI) rely on nodal enlargement.

Source

Scheidler J, Hricak H, Yu KK, Subak L, Segal MR. Radiological evaluation of lymph node metastases in patients with cervical cancer. A meta-analysis. *Journal of the American Medical Association* 1997; 278: 1096–1101.

Reitsma J, Glas A, Rutjes A, Scholten R, Bossuyt P, Zwinderman A. Bivariate analysis of sensitivity and specificity produces informative summary measures in diagnostic reviews. *Journal of Clinical Epidemiology* 2005; 58: 982–990.

Description

Obtain standard errors or variances of a model fit

Usage

```
se(object, ...)
variances(object, ...)
tau(object, ...)
tau2(object, ...)
```
Arguments

Value

For se the standard errors of object, and for variances the variances. For tau the heterogeneity of the coefficients.

Author(s)

Valentijn de Jong

stackedglm *Stacked Regression*

Description

This function combines one or more existing prediction models into a so/called meta-model.

Usage

```
stackedglm(models, family = binomial, data)
```
Arguments

Author(s)

Thomas Debray <thomas.debray@gmail.com>

subset.metapred *Subsetting metapred fits*

Description

Return a model from the cross-validation procedure or the final 'global' model. Caution: This function is still under development.

Usage

```
## S3 method for class 'metapred'
subset(
 x,
 select = "cv",step = NULL,
 model = NULL,stratum = NULL,
 add = TRUE,...
)
```
Arguments

summary.riley 63

Value

An object of class mp.cv for select $=$ "cv" and an object of class mp.global for select $=$ "global". In both cases, additional data is added to the resulting object, thereby making it suitable for further methods.

Author(s)

Valentijn de Jong

Examples

```
data(DVTipd)
DVTipd$cluster <- letters[1:4] # Add a fictional clustering to the data.
mp <- metapred(DVTipd, strata = "cluster", formula = dvt ~ histdvt + ddimdich, family = binomial)
subset(mp) # best cross-validated model
subset(mp, select = "global") # Final model fitted on all strata.
subset(mp, step = 1) # The best model of step 1
subset(mp, step = 1, model = "histdvt") # The model in which histdvt was removed, in step 1.
```


Description

Parameter summaries Provides the summary estimates of the alternative model for bivariate randomeffects meta-analysis by Riley et al. (2008) with their corresponding standard errors (derived from the inverse Hessian). For confidence intervals, asymptotic normality is assumed.

Usage

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

Details

For meta-analysis of diagnostic test accuracy data, beta1 equals the logit sensitivity (Sens) and beta2 equals the logit false positive rate (FPR).

Value

array with confidence intervals for the estimated model parameters. For diagnostic test accuracy data, the resulting summary sensitivity and false positive rate are included.

Note

For the overall correlation (rho) confidence intervals are derived using the transformation $logit$ ((rho+1)/2). Similarly, the logit transformation is used to derive confidence intervals for the summary sensitivity and false positive rate.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

References

Riley RD, Thompson JR, Abrams KR. An alternative model for bivariate random-effects metaanalysis when the within-study correlations are unknown. *Biostatistics* 2008; 9: 172–186.

summary.uvmeta *Summarizing Univariate Meta-Analysis Models*

Description

This function provides summary estimates of a fitted univariate meta-analysis model.

Usage

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

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Author(s)

Thomas Debray <thomas.debray@gmail.com>

References

- Borenstein M, Hedges LV, Higgins JPT, Rothstein HR. A basic introduction to fixed-effect and random-effects models for meta-analysis. *Research Synthesis Methods* 2010; 1: 97–111.
- DerSimonian R, Laird N. Meta-analysis in clinical trials. *Controlled Clinical Trials* 1986; 7: 177–188.
- Riley RD, Higgins JPT, Deeks JJ. Interpretation of random effects meta-analyses. *British Medical Journal* 2011; 342: d549.

See Also

[uvmeta](#page-65-0)

Tzoulaki *The incremental value of cardiovascular risk factors*

Description

Tzoulaki et al. (2009) reviewed studies that evaluated various candidate prognostic factors in their ability to improve prediction of coronary heart disease (CHD) or other outcomes beyond what the Framingham risk score (FRS) can achieve.

Usage

data("Tzoulaki")

Format

A data frame containing data from 27 studies on the following 2 variables.

PubmedID a character vector with the Pubmed ID of the study

N a numeric vector describing the study size

- N.events a numeric vector describing the observed number of events
- FRS.orig.refitted a boolean vector describing whether the coefficients of the original Framingham Risk Score (FRS) were re-estimated
- FRS.modif.refitted a boolean vector describing whether the coefficients of the modified Framingham Risk Score were re-estimated
- predictors a character vector indicating which new risk factor(s) were included in the modified FRS
- outcome a character vector indicating the primary outcome being predicted
- AUC.orig a numeric vector describing the Area under the ROC curve (AUC) of the original FRS model
- AUC.orig.CIl a numeric vector describing the lower boundary of the 95% confidence interval of the AUC of the original FRS model
- AUC.orig.CIu a numeric vector describing the upper boundary of the 95% confidence interval of the AUC of the original FRS model
- AUC.modif a numeric vector describing the Area under the ROC curve (AUC) of the modified FRS model that includes one or more new risk factors
- AUC.modif.CIl a numeric vector describing the lower boundary of the 95% confidence interval of the AUC of the modified FRS model
- AUC.modif.CIu a numeric vector describing the upper boundary of the 95% confidence interval of the AUC of the modified FRS model
- pval.AUCdiff a numeric vector with the p-value of the difference between AUC.orig and AUC.modif
- sign.AUCdiff a boolean vector indicating whether the difference between AUC.orig and AUC.modif is below 0.05

Source

Tzoulaki I, Liberopoulos G, Ioannidis JPA. Assessment of claims of improved prediction beyond the Framingham risk score. *JAMA*. 2009 Dec 2;302(21):2345–52.

Examples

```
data(Tzoulaki)
## maybe str(Tzoulaki) ; plot(Tzoulaki) ...
```
uvmeta *Univariate meta-analysis*

Description

This function summarizes multiple estimates for a single parameter by assuming a fixed (i.e. common) effect or random effects across studies. The summary estimate is obtained by calculating a weighted mean that accounts for sample size and (in case random effects are assumed) for betweenstudy heterogeneity.

Usage

```
uvmeta(
  r,
  r.se,
  r.vi,
 method = "REML".test = "knha",
  labels,
  na.action,
  n.chains = 4,
  pars,
```
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```
verbose = FALSE,
```
... \mathcal{L}

Arguments

Details

Unless specified otherwise, all meta-analysis models assume random effects and are fitted using restricted maximum likelihood estimation with the metafor package (Viechtbauer 2010). Further, confidence intervals for the average performance are based on the Hartung-Knapp-Sidik-Jonkman method, to better account for the uncertainty in the estimated between-study heterogeneity (Debray 2016). A Bayesian meta-analysis can be performed by specifying method="BAYES". In that case, the R packages runjags and riags must be installed.

For random-effects models, a prediction interval for the pooled effect size is displayed. This interval predicts in what range future effect sizes will fall given what has already been observed (Higgins 2009, Riley 2011).

Bayesian meta-analysis models: For Bayesian meta-analysis models that involve the Gibbs sampler (method="BAYES"), the R packages runjags and rjags must be installed. The Bayesian approach uses an uninformative Normal prior for the mean and a uniform prior for the betweenstudy variance of the pooled effect size (Higgins 2009). By default, the Normal prior has a mean of 0 and a variance of 1000. These hyperparameters can, however, be altered through the variables hp.mu.mean and hp.mu.var in the argument pars. The prior distribution of the between-study standard deviation is given by a uniform distribution, by default bounded between 0 and 100.

Value

An object of the class uvmeta for which many standard methods are available.

- "data" array with (transformed) data used for meta-analysis, and method(s) used for restoring missing information.
- "method" character string specifying the meta-analysis method.
- "est" estimated performance statistic of the model. For Bayesian meta-analysis, the posterior median is returned.
- "se" standard error (or posterior standard deviation) of the summary estimate.
- "tau2" estimated amount of (residual) heterogeneity. Always 0 when method="FE". For Bayesian meta-analysis, the posterior median is returned.
- "se.tau2" estimated standard error (or posterior standard deviation) of the between-study variation.
- "ci.lb" lower bound of the confidence (or credibility) interval of the summary estimate
- "ci.ub" upper bound of the confidence (or credibility) interval of the summary estimate
- "**pi.lb**" lower bound of the (approximate) prediction interval of the summary estimate
- "pi.ub" upper bound of the (approximate) prediction interval of the summary estimate

"fit" the full results from the fitted model

"slab" vector specifying the label of each study.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

References

Biggerstaff BJ, Tweedie RL. Incorporating variability in estimates of heterogeneity in the random effects model in meta-analysis. *Statistics in Medicine* 1997; 16: 753–768.

Borenstein M, Hedges LV, Higgins JPT, Rothstein HR. A basic introduction to fixed-effect and random-effects models for meta-analysis. *Research Synthesis Methods* 2010; 1: 97–111. [doi:10.1002](https://doi.org/10.1002/jrsm.12)/ [jrsm.12](https://doi.org/10.1002/jrsm.12)

DerSimonian R, Laird N. Meta-analysis in clinical trials. *Controlled Clinical Trials* 1986; 7: 177– 188.

Graham PL, Moran JL. Robust meta-analytic conclusions mandate the provision of prediction intervals in meta-analysis summaries. *Journal of Clinical Epidemiology* 2012; 65: 503–510.

Higgins JPT, Thompson SG. Quantifying heterogeneity in a meta-analysis. *Statistics in Medicine* 2002; 21: 1539–1558.

Higgins JPT, Thompson SG, Spiegelhalter DJ. A re-evaluation of random-effects meta-analysis. *J R Stat Soc Ser A Stat Soc*. 2009;172:137–59. [doi:10.1111/j.1467985X.2008.00552.x](https://doi.org/10.1111/j.1467-985X.2008.00552.x)

Riley RD, Higgins JPT, Deeks JJ. Interpretation of random effects meta-analyses. *British Medical Journal* 2011; 342: d549. [doi:10.1136/bmj.d549](https://doi.org/10.1136/bmj.d549)

Viechtbauer W. Conducting Meta-Analyses in R with the metafor Package. *Journal of Statistical Software*. 2010; 36. [doi:10.18637/jss.v036.i03](https://doi.org/10.18637/jss.v036.i03)

Examples

```
data(Roberts)
```

```
# Frequentist random-effects meta-analysis
fit1 <- with(Roberts, uvmeta(r=SDM, r.se=SE, labels=rownames(Roberts)))
summary(fit1)
plot(fit1) #show a forest plot
fit1
## Not run:
# Bayesian random effects meta-analysis
fit2 <- with(Roberts, uvmeta(r=SDM, r.se=SE, labels=rownames(Roberts), method="BAYES"))
plot(fit2)
```
End(Not run)

uvmeta-class *Class "uvmeta". Result of a univariate meta-analysis.*

Description

This class encapsulates results of a univariate meta-analysis.

Objects from the Class

Objects can be created by calls of the form [uvmeta](#page-65-0).

- call: (language) The call to [uvmeta](#page-65-0).
- data: (data frame) The data used for the meta-analysis.
- results: (data frame) Contains the pooled effect size (mu), the between-study variability (tausq), Cochran's Q statistic (Q) and Higgins' and Thompson's I square statistic (Isq). For each estimate, error variances are provided with predefined confidence (method="MOM") or credibility (method="bayes") intervals.
- model: (character) The meta-analysis model used.
- method: (character) The estimator used.
- na.action: (character) Information from the action which was applied to object if NAs were handled specially, or NULL.
- df: (numeric) Degrees of freedom.
- numstudies: (numeric) The amount of studies used in the meta-analysis.
- pred.int: (data frame) A prediction interval, predicting in what range future effect sizes will fall given what has already been observed (based on a Student's t-distribution, cfr. Riley 2011)
- formula: (character) If a formula was specified, a character vector giving the formula and parameter specifications.

Methods

print signature(object = "uvmeta"): Print object summary.

- forest signature(object = "uvmeta"): Plot a forest plot with the summary estimate.
- summary signature(object = "uvmeta"): Generate object summary.

Examples

data(Collins)

```
#Extract effect size and error variance
r <- Collins$logOR
vars <- Collins$SE**2
```

```
#Frequentist random-effects meta-analysis
fit1 <- uvmeta(r,vars)
```

```
#Extract results
fit1$results
```
Slots

Description

This function provides summary estimates for the concordance statistic, the total observed-expected ratio or the calibration slope. Where appropriate, data transformations are applied and missing information is derived from available quantities. Unless specified otherwise, all meta-analysis models assume random effects and are fitted using restricted maximum likelihood estimation with the metafor package (Viechtbauer 2010). Further, confidence intervals for the average performance are based on the Hartung-Knapp-Sidik-Jonkman method. When conducting a Bayesian meta-analysis, the R packages runjags and rjags must be installed.

Usage

```
valmeta(
  measure = "cstat",
  cstat,
  cstat.se,
  cstat.cilb,
  cstat.ciub,
  cstat.cilv,
  sd.LP,
  OE,
  OE.se,
  OE.cilb,
  OE.ciub,
  OE.cilv,
  citl,
  citl.se,
  N,
  O,
  E,
  Po,
  Po.se,
  Pe,
  data,
  method = "REML",test = "knha",verbose = FALSE,
  slab,
  n.chains = 4,
  pars,
  ...
)
```
Arguments

Details

Meta-analysis of the concordance statistic: A summary estimate for the concorcance (c-) statistic can be obtained by specifying measure="cstat". The c-statistic is a measure of discrimination, and indicates the ability of a prediction model to distinguish between patients developing and not developing the outcome. The c-statistic typically ranges from 0.5 (no discriminative ability) to 1 (perfect discriminative ability). When missing, the c-statistic and/or its standard error are derived from other reported information. See [ccalc](#page-7-0) for more information.

By default, it is assumed that the logit of the c-statistic is Normally distributed within and across studies (pars\$model.cstat = "normal/logit"). Alternatively, it is possible to assume that the raw c-statistic is Normally distributed across studies pars\$model.cstat = "normal/identity".

Meta-analysis of the total observed versus expected ratio: A summary estimate for the total observed versus expected (O:E) ratio can be obtained by specifying measure="OE". The total O:E ratio provides a rough indication of the overall model calibration (across the entire range of predicted risks). When missing, the total O:E ratio and/or its standard error are derived from other reported information. See [oecalc](#page-42-0) for more information.

For frequentist meta-analysis, within-study variation can either be modeled using a Normal (model.oe = "normal/log" or model.oe = "normal/identity") or a Poisson distribution (model.oe = "poisson/log"). When performing a Bayesian meta-analysis, all data are modeled using a one-stage random effects (hierarchical related regression) model. In particular, a binomial distribution (if O, E and N is known), a Poisson distribution (if only O and E are known) or a Normal distribution (if OE and OE.se or OE.95CI are known) is selected separately for each study.

Bayesian meta-analysis: All Bayesian meta-analysis models assume the presence of random effects. Summary estimates are based on the posterior mean. Credibility and prediction intervals are directly obtained from the corresponding posterior quantiles.

The prior distribution for the (transformed) performance estimate is modeled using a Normal distribution, with mean hp.mu.mean (defaults to 0) and variance hp.mu.var (defaults to 1000). For meta-analysis of the total O:E ratio, the maximum value for hp.mu.var is 100.

By default, the prior distribution for the between-study standard deviation is modeled using a uniform distribution (hp.tau.dist="dunif"), with boundaries hp.tau.min and hp.tau.max. Alternative choices are a truncated Student-t distribution (hp.tau.dist="dhalft") with a mean of hp.tau.mean, a standard deviation of hp.tau.sigma and hp.tau.df degrees of freedom. This distribution is again restricted to the range hp.tau.min to hp.tau.max.

Value

An object of class valmeta with the following elements:

- "data" array with (transformed) data used for meta-analysis, and method(s) used for restoring missing information.
- "measure" character string specifying the performance measure that has been meta-analysed.
- "method" character string specifying the meta-analysis method.
- "model" character string specifying the meta-analysis model (link function).
- "est" summary estimate for the performance statistic. For Bayesian meta-analysis, the posterior median is returned.
- "ci.lb" lower bound of the confidence (or credibility) interval of the summary performance estimate.
- "ci.ub" upper bound of the confidence (or credibility) interval of the summary performance estimate.
- "pi.lb" lower bound of the (approximate) prediction interval of the summary performance estimate.
- "pi.ub" upper bound of the (approximate) prediction interval of the summary performance estimate.

"fit" the full results from the fitted model.

"slab" vector specifying the label of each study.

Note

The width of calculated confidence, credibility and prediction intervals can be specified using level in the pars argument (defaults to 0.95).

References

Debray TPA, Damen JAAG, Snell KIE, Ensor J, Hooft L, Reitsma JB, et al. A guide to systematic review and meta-analysis of prediction model performance. *BMJ*. 2017;356:i6460. [doi:10.1136/](https://doi.org/10.1136/bmj.i6460) [bmj.i6460](https://doi.org/10.1136/bmj.i6460)

Debray TPA, Damen JAAG, Riley R, Snell KIE, Reitsma JB, Hooft L, et al. A framework for metaanalysis of prediction model studies with binary and time-to-event outcomes. *Stat Methods Med Res*. 2019;28:2768–86. [doi:10.1177/0962280218785504](https://doi.org/10.1177/0962280218785504)

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Riley RD, Tierney JF, Stewart LA. Individual participant data meta-analysis: a handbook for healthcare research. Hoboken, NJ: Wiley; 2021. ISBN: 978-1-119-33372-2.

Steyerberg EW, Nieboer D, Debray TPA, van Houwelingen HC. Assessment of heterogeneity in an individual participant data meta-analysis of prediction models: An overview and illustration. *Stat Med*. 2019; 38:4290–309. [doi:10.1002/sim.8296](https://doi.org/10.1002/sim.8296)

Viechtbauer W. Conducting Meta-Analyses in R with the metafor Package. *Journal of Statistical Software*. 2010; 36. [doi:10.18637/jss.v036.i03](https://doi.org/10.18637/jss.v036.i03)

See Also

[ccalc](#page-7-0) to calculate concordance statistics and corresponding standard errors, [oecalc](#page-42-0) to calculate the total O:E ratio and corresponding standard errors, plot. valmeta to generate forest plots

Examples

```
######### Validation of prediction models with a binary outcome #########
data(EuroSCORE)
# Meta-analysis of the c-statistic (random effects)
fit <- valmeta(cstat=c.index, cstat.se=se.c.index, cstat.cilb=c.index.95CIl,
              cstat.ciub=c.index.95CIu, cstat.cilv=0.95, N=n, O=n.events,
              slab=Study, data=EuroSCORE)
plot(fit)
# Nearly identical results when we need to estimate the SE
valmeta(cstat=c.index, N=n, O=n.events, slab=Study, data=EuroSCORE)
# Two-stage meta-analysis of the total O:E ratio (random effects)
valmeta(measure="OE", O=n.events, E=e.events, N=n, slab=Study, data=EuroSCORE)
valmeta(measure="OE", O=n.events, E=e.events, data=EuroSCORE)
valmeta(measure="OE", Po=Po, Pe=Pe, N=n, data=EuroSCORE)
## Not run:
# One-stage meta-analysis of the total O:E ratio (random effects)
valmeta(measure="OE", O=n.events, E=e.events, data=EuroSCORE, method="ML",
       pars=list(model.oe="poisson/log"))
# Bayesian random effects meta-analysis of the c-statistic
fit2 <- valmeta(cstat=c.index, cstat.se=se.c.index, cstat.cilb=c.index.95CIl,
                cstat.ciub=c.index.95CIu, cstat.cilv=0.95, N=n, O=n.events,
                data=EuroSCORE, method="BAYES", slab=Study)
# Bayesian one-stage random effects meta-analysis of the total O:E ratio
# Consider that some (but not all) studies do not provide information on N
# A Poisson distribution will be used for studies 1, 2, 5, 10 and 20
# A Binomial distribution will be used for the remaining studies
EuroSCORE.new <- EuroSCORE
EuroSCORE.new$n[c(1, 2, 5, 10, 20)] <- NA
pars <- list(hp.tau.dist="dhalft", # Prior for the between-study standard deviation
            hp.tau.sigma=1.5, # Standard deviation for 'hp.tau.dist'
            hp.tau.df=3, # Degrees of freedom for 'hp.tau.dist'
```

```
hp.tau.max=10, # Maximum value for the between-study standard deviation
            seed=5) # Set random seed for the simulationsfit3 <- valmeta(measure="OE", O=n.events, E=e.events, N=n, data=EuroSCORE.new,
       method="BAYES", slab=Study, pars=pars)
plot(fit3)
print(fit3$fit$model) # Inspect the JAGS model
print(fit3$fit$data) # Inspect the JAGS data
## End(Not run)
######### Validation of prediction models with a time-to-event outcome #########
data(Framingham)
# Meta-analysis of total O:E ratio after 10 years of follow-up
valmeta(measure="OE", Po=Po, Pe=Pe, N=n, data=Framingham)
```
vcov.riley *Calculate Variance-Covariance Matrix for a Fitted Riley Model Object*

Description

Returns the variance-covariance matrix of the main parameters of a fitted model object.

Usage

S3 method for class 'riley' vcov(object, ...)

Arguments

Details

The variance-covariance matrix is obtained from the inverse Hessian as provided by optim.

Value

A matrix of the estimated covariances between the parameter estimates in the Riley model: logit of sensitivity (mu1), logit of false positive rate (mu2), additional variation of mu1 beyond sampling error (psi1), additional variation of mu2 beyond sampling error (psi2) and a transformation of the correlation between psi1 and psi2 (rhoT). The original correlation is given as inv. logit(rhoT)*2-1.

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Note

A warning message is casted when the Hessian matrix contains negative eigenvalues. This implies that the identified minimum for the (restricted) negative log-likelihood is a saddle point, and that the solution is therefore not optimal.

Author(s)

Thomas Debray <thomas.debray@gmail.com>

References

Riley, RD., Thompson, JR., & Abrams, KR. (2008). "An alternative model for bivariate randomeffects meta-analysis when the within-study correlations are unknown." *Biostatistics*, 9, 172–186.

See Also

[riley](#page-52-0)

Zhang *Meta-analysis of the prognostic role of hormone receptors in endometrial cancer*

Description

This dataset comprises the results from 16 studies assessing the prognostic role of human epidermal growth factor receptor 2 (HER2) in endometrial cancer. These studies were previously identified in a systematic review by Zhang et al. to evaluate the overall risk of several hormone receptors for endometrial cancer survival.

Usage

data("Zhang")

Format

A data frame with 20 observations on the following 10 variables.

Study a factor with 16 levels to indicate the study

PrimaryAuthor a factor indicating the first author's last name

year a numeric vector indicating the publication year

Country a factor indicating the source country of the study data

Disease a factor indicating the studied disease. Possible levels are EC (endometrial cancer), EEC (endometrioid endometrial cancer) and UPSC (uterine papillary serous carcinoma)

N a numeric vector describing the total sample size of each study

HR a numeric vector describing the estimated hazard ratio of each study

HR.025 a numeric vector describing the lower boundary of the 95% confidence interval of HR

- HR.975 a numeric vector describing the upper boundary of the 95% confidence interval of HR
- outcome a factor indicating the studied outcome. Possible levels are OS (overall survival) and PFS (progression-free survival)

Details

Eligible studies were identified by searching the PubMed and EMBASE databases for publications from 1979 to May 2014. Data were collected from studies comparing overall survival or progression-free survival in patients with elevated levels of human epidermal growth factor receptor 2 with those in patients with lower levels.

Source

Zhang Y, Zhao D, Gong C, Zhang F, He J, Zhang W, et al. Prognostic role of hormone receptors in endometrial cancer: a systematic review and meta-analysis. World J Surg Oncol. 2015 Jun 25;13:208.

References

Riley RD, Jackson D, Salanti G, Burke DL, Price M, Kirkham J, et al. Multivariate and network meta-analysis of multiple outcomes and multiple treatments: rationale, concepts, and examples. BMJ. 2017 13;358:j3932.

Examples

data(Zhang)

```
# Display the hazard ratios for overall survival in a forest plot
ds <- subset(Zhang, outcome=="OS")
with(ds, forest(theta = HR, theta.ci.lb = HR.025, theta.ci.ub = HR.975,
              theta.slab = Study, xlab = "Hazard ratio of HER2 versus OS", refline = 1))
```
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