Package ‘xmeta’

September 7, 2023

Type Package

Title A Toolbox for Multivariate Meta-Analysis

Version 1.3.2

Date 2023-09-04

Imports plotrix, aod, glmml, numDeriv, metafor, mvmeta, stats, MASS


Depends R (>= 3.5.0)

License GPL (>= 2)

LazyLoad no

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NeedsCompilation no

Repository CRAN

URL https://github.com/Penncil/xmeta

BugReports https://github.com/Penncil/xmeta/issues

Encoding UTF-8

RoxygenNote 7.2.3

Date/Publication 2023-09-07 02:10:02 UTC
Description

The package `xmeta` consists of a collection of functions for making inference and detecting publication bias in multivariate meta-analysis (MMA).

Inference

The aim of the estimation methods is to estimate the coefficients $\beta$ and the components of the between-study (co)variance matrix $\Psi$ for multivariate random-effects meta-analysis. One major challenge in MMA is the standard inference procedures, such as the maximum likelihood or maximum restricted likelihood inference, require the within-study correlations, which are usually unavailable. Different estimators with and without the knowledge of within study correlation are implemented in the package `xmeta`. The estimation methods available in function `mmeta` are:

- Restricted maximum likelihood for MMA with continuous outcomes
- Composite likelihood method for MMA with continuous outcomes
- Method of moment for MMA with continuous outcomes
- Improved method for Riley model for MMA with continuous outcomes
- Marginal bivariate normal model for MMA with binary outcomes
- Marginal beta-binomial model for MMA with binary outcomes
- Hybrid model for disease prevalence along with sensitivity and specificity for diagnostic test accuracy
- Trivariate model for multivariate meta-analysis of diagnostic test accuracy
Small study effects

Detecting and accounting for small study effects are challenging in MMA setting. The multivariate nature is often not fully accounted for by the existing univariate methods. The score test for detecting small study effects in MMA when the within-study correlations are unknown is implemented in the function `msset`.

Galaxy Plot

A New Visualization Tool of Bivariate Meta-Analysis Studies. This function `galaxy` returns the galaxy plot to visualize bivariate meta-analysis data, which faithfully retains the information in two separate funnel plots, while providing useful insights into outcome correlations, between-study heterogeneity and joint asymmetry. Galaxy plot is the counterpart of the funnel plot in the multivariate setting. The galaxy plot is an intuitive visualization tool that can aid in interpretation of results of multivariate meta-analysis. It preserves all of the information presented by separate funnel plots for each outcome while elucidating more complex features that may only be revealed by examining the joint distribution of the bivariate outcomes.

Publication bias in bivariate meta-analysis

The function `galaxy.trimfill` implements a bivariate T&F method accounting for publication bias in bivariate meta-analysis, based on symmetry of the galaxy plot. The bivariate T&F method assumes studies are suppressed based on a weighted sum of the two outcomes. We use a searching algorithm to find the optimal direction which gives the most trimmed studies. This is based on the observation that the closer a direction is to the truth, the more studies are expected to be trimmed along that direction.

ca125

**Recurrent ovarian carcinoma study**

**Description**

A meta-analysis of 52 studies that were reported between January 1995 and November 2007.

**Format**

The data frame contains the following columns:

- `n` total number of subjects
- `PiY` disease prevalence
- `SeY` true positive
- `n1` subjects with disease
- `SpY` true negative
- `n1` health individuals

**Note**

The dataset ca125 is used to conduct multivariate meta-analysis of diagnostic test accuracy.
References


See Also

mmeta, summary.mmeta

Examples

data(ca125)
summary(ca125)

dat.gen Generate bivariate meta analysis studies

Description

Generate bivariate meta analysis studies based on random-effects model, some studies with smallest weighted sum of the two outcomes are suppressed.

Usage

dat.gen(
  m.o,  
m.m,  
s.m,  
angle.LC = pi/4,  
mybeta,  
tau.sq,  
rho.w,  
rho.b,  
s.min = 0.01,  
m.m.o = 0,  
s2.dist = 2,  
verbose = F
)

Arguments

m.o  number of observed studies
m.m  number of missing / suppressed studies
s.m  vector of the mean of the variances of the two outcomes
angle.LC  direction of suppressing line, default is pi/4, i.e. the studies on the left bottom corner are missing
mybeta  the true center of the effect sizes
tau.sq  between-study variance, the larger it is the more heterogeneity.
rho.w  within-study correlation of the two outcomes
rho.b  between-study correlation of the two outcomes
s.min  minimum of the variances of the outcomes, default is 0.01
m.m.o  number of studies on one side of the suppressing line been observed, i.e. non-deterministic suppressing, default is 0, i.e. deterministic suppressing
s2.dist  options for generating the outcomes’ variances. 1=runif, 2=runif^2, 3=runif^4, 4=rnorm
verbose  logical, galaxy plot the studies? Default FALSE

Author(s)
Chongliang Luo, Yong Chen

References

Description
A new visualization method that simultaneously presents the effect sizes of bivariate outcomes and their standard errors in a two-dimensional space.

Usage
galaxy(data, y1, s1, y2, s2, scale1, scale2, scale.adj, corr, group, study.label, annotate, xlab, ylab, main, legend.pos)

Arguments
data  dataset with at least 4 columns for the effect sizes of the two outcomes and their standard errors
y1  column name for outcome 1, default is ‘y1’
s1  column name for standard error of y1, default is ‘s1’
y2  column name for outcome 2, default is ‘y2’
s2  column name for standard error of y2, default is ‘s2’
scale1 parameter for the length of the cross hair: the ellipse width is scale1 / s1 * scale.adj
scale2 parameter for the length of the cross hair: the ellipse height is scale2 / s2 * scale.adj
scale.adj a pre-specified parameter to adjust for scale1 and scale2
corr column name for within-study correlation
group column name for study group
study.label column name for study label
annotate logical specifying whether study label should be added to the plot, default is FALSE.
xlab x axis label, default y1
ylab y axis label, default y2
main main title
legend.pos The position of the legend for study groups if group is specified, see legend, default is 'bottomright'.

Details
This function returns the galaxy plot to visualize bivariate meta-analysis data, which faithfully retains the information in two separate funnel plots, while providing useful insights into outcome correlations, between-study heterogeneity and joint asymmetry. Galaxy plot: a new visualization tool of bivariate meta-analysis studies. Funnel plots have been widely used to detect small study effects in the results of univariate meta-analyses. However, there is no existing visualization tool that is the counterpart of the funnel plot in the multivariate setting. We propose a new visualization method, the galaxy plot, which can simultaneously present the effect sizes of bivariate outcomes and their standard errors in a two-dimensional space. The galaxy plot is an intuitive visualization tool that can aid in interpretation of results of multivariate meta-analysis. It preserves all of the information presented by separate funnel plots for each outcome while elucidating more complex features that may only be revealed by examining the joint distribution of the bivariate outcomes.

Author(s)
Chuan Hong, Chongliang Luo, Yong Chen

References

Examples
data(sim_dat)
galaxy(data=sim_dat, scale.adj = 0.9, corr = 'corr', group = 'subgroup', study.label = 'study.id', annotate = TRUE, main = 'galaxy plot')
Description

Bivariate T&F method accounting for small-study effects in bivariate meta-analysis, based on symmetry of the galaxy plot.

Usage

galaxy.trimfill(
  y1,
  v1,
  y2,
  v2,
  n.grid = 12,
  angle,
  estimator,
  side,
  rho = 0,
  method = "mm",
  method.uni = "DL",
  maxiter = 20,
  var.names = c("y1", "y2"),
  scale = 0.02,
  verbose = FALSE
)

Arguments

- **y1**: vector of the effect size estimates of the first outcome
- **v1**: estimated variance of y1
- **y2**: vector of the effect size estimates of the second outcome
- **v2**: estimated variance of y2
- **n.grid**: number of grid (equally spaced) candidate directions that the optimal projection direction are searched among, see Details
- **angle**: angles of candidate projection directions not by grid, this will overwrite n.grid
- **estimator**: estimator used for the number of trimmed studies in univariate T&F on the projected studies, one of c('R0', 'L0', 'Q0')
- **side**: either "left" or "right", indicating on which side of the galaxy plot the missing studies should be imputed. If null determined by the univariate T&F
- **rho**: correlation between y1 and y2 when computing the variance of the projected studies. Default is the estimated cor(y1, y2)
- **method**: method to estimate the center for the bivariate outcomes. Default is 'mm', i.e. random-effects model
method.uni  method to estimate the center for the univariate projected studies using a univariate T&F procedure. Default is 'DL', i.e. fixed-effects model
maxiter  max number of iterations used in the univariate T&F. Default is 20.
var.names  names of the two outcomes used in the galaxy plot (if plotted). Default is c('y1', 'y2')
scale  constant scale for plotting the galaxy plot for the bivariate studies, Default is 0.02.
verbose  plot the galaxy plot? Default is FALSE.

Details
The bivariate T&F method assumes studies are suppressed based on a weighted sum of the two outcomes, i.e. the studies with smallest values of \( z_i = c_1 \cdot y_{1i} + c_2 \cdot y_{2i}, i=1,...,N \) are suppressed. We use a searching algorithm to find the optimal ratio of \( c_1 \) and \( c_2 \) (i.e. a direction), which gives the most trimmed studies. This is based on the observation that the closer a direction is to the truth, the more studies are expected to be trimmed along that direction. We set a sequence of equally-spaced candidate directions with angle \( a_m = m \cdot \pi / M \), and \( (c_1, c_2) = (\cos(a_m), \sin(a_m)), m=1,...,M \).

Value
List with component:
- res  a data.frame of 9 columns and n.grid rows. Each row is the result for projection along one candidate grid direction, and the columns are named: 'y1.c', 'y2.c' for projected bivariate center, 'y1.f', 'y2.f' for bivariate center using filled studies, 'k0', 'se.k0' for estimated number of trimmed studies and its standard error, 'se.y1.f', 'se.y2.f' for standard errors of 'y1.f', 'y2.f', 'side.left' for the estimated side
- ID.trim  list of vectors of ids of studies been trimmed along each of the candidate direction.

Author(s)
Chongliang Luo, Yong Chen

References

Examples
```r
## Not run:
require(MASS)
require(mvmeta)
require(metafor)
set.seed(123)
mydata <- dat.gen(m.o=50, m.m=20, # # observed studies, # missing studies
                 s.m= c(0.5, 0.5), # c(mean(s1), mean(s2))
```

galaxy.trimfill

angle.LC = pi/4,  # suppress line direction
mybeta=c(2,2),  # true effect size
tau.sq=c(0.1, 0.1),  # true between-study var
rho.w=0.5, rho.b=0.5,  # true within-study and between-study corr
s.min = 0.1,  # s1i ~ Unif(s.min, 2*s.m[1]-s.min)
verbose = TRUE)

y1 <- mydata$mydat.sps$y1
y2 <- mydata$mydat.sps$y2
v1 <- mydata$mydat.sps$s1^2
v2 <- mydata$mydat.sps$s2^2

## unadjusted est
mv_obs <- mvmeta(cbind(y1, y2), cbind(v1, v2), method='mm')
c(mv.obs$coef)
# 2.142687 2.237741

estimator <- 'R0'
## univariate T&F based on y1 or y2
y1.rma <- rma(y1, v1, method='FE')
y2.rma <- rma(y2, v2, method='FE')
y1.tf <- trimfill.rma(y1.rma, estimator = estimator, method.fill = 'DL')
y2.tf <- trimfill.rma(y2.rma, estimator = estimator, method.fill = 'DL')
c(y1.tf$beta, y2.tf$beta)
# 2.122231 2.181333

## bivariate T&F method (based on galaxy plot)
tf.grid <- galaxy.trimfill(y1, v1, y2, v2, n.grid = 12,
estimator=estimator, side='left',
method.uni = 'FE',
method = 'mm',
rho=0.5, maxiter=100, verbose=FALSE)
tf.grid$res
tf.grid$res[which(tf.grid$res$k0==max(tf.grid$res$k0)),3:5]
#  y1.f  y2.f k0
# 2.053306 2.162347 14

## less bias by the proposed bivariate T&F method
rbind(true = c(2,2),
unadjusted=c(mv.obs$coef),
tf.uni = c(y1.tf$beta, y2.tf$beta),
tf.biv = tf.grid$res[which(tf.grid$res$k0==max(tf.grid$res$k0)),3:4])

## unlike the univariate T&Fs, biv T&F obtains one estimate of # missing studies
c(k0.true = 20,
k0.tf.uni.y1 = y1.tf$k0,
k0.tf.uni.y2 = y2.tf$k0,
k0.tf.biv = tf.grid$res[which(tf.grid$res$k0==max(tf.grid$res$k0)),5])
# k0.true k0.tf.uni.y1 k0.tf.uni.y2 k0.tf.biv
# 20 2 8 14
Description

Methods for multivariate random-effects meta-analysis

Usage

mmeta(data, rhow, type, k, method)

Arguments

data: dataset
rhow: within-study correlation
type: either "continuous" or "binary", indicating the type of outcomes.
k: integer indicating the number of outcomes

Details

Inference on the multivariate random-effects meta-analysis for both continuous and binary outcomes

The function can be used in meta-analyses with continuous outcomes and binary outcomes (e.g., mean differences, diagnostic test results in diagnostic accuracy studies, the exposure status of both cases and controls in case-control studies and so on). Different estimators with and without the knowledge of within-study correlations are implemented in this function. The estimation methods include

- Restricted maximum likelihood for MMA with continuous outcomes (nn.reml)
- Composite likelihood method for MMA with continuous outcomes (nn.cl)
- Moment of method for MMA with continuous outcomes (nn.mom)
- Improved method for Riley model for MMA with continuous outcomes (nn.rs)
- Marginal bivariate normal model for MMA with binary outcomes (bn.cl)
- Marginal beta-binomial model for MMA with binary outcomes (bb.cl)
- Hybrid model for disease prevalence along with sensitivity and specificity for diagnostic test accuracy (tb.cl)
- Trivariate model for multivariate meta-analysis of diagnostic test accuracy (tn.cl)
**Value**

An object of class "mmeta". The object is a list containing the following components:

- `beta` estimated coefficients of the model.
- `beta.cov` covariance matrix of the coefficients.

**Multivariate random-effects meta analysis**

We consider a meta-analysis with m studies where two outcomes in each study are of interest. For the i-th study, denote $Y_{ij}$ and $s_{ij}$ the summary measure for the j-th outcome of interest and associated standard error respectively, both assumed known, $i = 1, \ldots, m,$ and $j = 1, 2$. Each summary measure $Y_{ij}$ is an estimate of the true effect size $\theta_{ij}$. To account for heterogeneity in effect size across studies, we assume $\theta_{ij}$ to be independently drawn from a common distribution with overall effect size $\beta_j$ and between study variance $\tau_j^2, j = 1, 2$. Under normal distribution assumption for $Y_{ij}$ and $\theta_{ij}$, the general bivariate random-effects meta-analysis can be written as

$$
\begin{pmatrix}
Y_{i1} \\
Y_{i2}
\end{pmatrix}
\sim N
\left( 
\begin{pmatrix}
\theta_{i1} \\
\theta_{i2}
\end{pmatrix},
\Delta_i
\right),
\Delta_i = 
\begin{pmatrix}
\sigma_{i1}^2 \\
\sigma_{i2}^2
\end{pmatrix}
\begin{pmatrix}
s_{i1}^2 & s_{i1}s_{i2}\rho_{Wi_i} \\
s_{i1}s_{i2}\rho_{Wi_i} & s_{i2}^2
\end{pmatrix},
$$

where $\Delta_i$ and $\Omega$ are the respective within-study and between-study covariance matrices, and $\rho_{Wi_i}$ and $\rho_{B}$ are the respective within-study and between-study correlations.

**Restricted maximum likelihood for MMA**

When the within-study correlations are known, inference on the overall effect sizes $\beta_1$ and $\beta_2$ or their comparative measures (e.g., $\beta_1 - \beta_2$) can be based on the marginal distribution of $(Y_{i1}, Y_{i2})$

$$
\begin{pmatrix}
Y_{i1} \\
Y_{i2}
\end{pmatrix}
\sim N
\left( 
\begin{pmatrix}
\beta_1 \\
\beta_2
\end{pmatrix},
V_i
\right),
V_i = \Delta_i + \Omega = 
\begin{pmatrix}
\sigma_{i1}^2 + \tau_1^2 & s_{i1}s_{i2}\rho_{Wi_i} + \tau_1\tau_2\rho_{B} \\
s_{i1}s_{i2}\rho_{Wi_i} + \tau_1\tau_2\rho_{B} & s_{i2}^2 + \tau_2^2
\end{pmatrix}.
$$

For simplicity of notation, denote $Y_i = (Y_{i1}, Y_{i2})^T$, $\beta = (\beta_1, \beta_2)^T$, $\eta_1 = (\beta_1, \tau_1^2)^T$ and $\eta_2 = (\beta_2, \tau_2^2)^T$. The restricted likelihood of $(\eta_1, \eta_2, \rho_B)$ can be written as

$$
\log L(\eta_1, \eta_2, \rho_B) = -\frac{1}{2} \left[\log \left( \sum_{i=1}^m V_i^{-1} \right) + \sum_{i=1}^m \log |V_i| + (Y_i - \beta)^T V_i^{-1} (Y_i - \beta) \right].
$$

The parameters $(\eta_1, \eta_2, \rho_B)$ can be estimated by the restricted maximum likelihood (REML) approach as described in Van Houwelingen et al. (2002). The REML method for MMA is specified via method argument (method="nn.reml").

The standard inference procedures, such as the maximum likelihood or maximum restricted likelihood inference, require the within-study correlations, which are usually unavailable. In case within-study correlations are unknown, then one can leave the $\rho_w$ argument unspecified, and specify a method that does not require the within-study correlations via method argument.
Composite likelihood method for MMA with continuous outcomes

Chen et al. (2014) proposed a pseudolikelihood method for MMA with unknown within-study correlation. The pseudolikelihood method does not require within-study correlations, and is not prone to singular covariance matrix problem. In addition, it can properly estimate the covariance between pooled estimates for different outcomes, which enables valid inference on functions of pooled estimates, and can be applied to meta-analysis where some studies have outcomes MCAR. This composite likelihood method for MMA is specified via method argument (method="nn.cl").

Moment of method for MMA with continuous outcomes

Chen et al. (2015) proposed a simple non-iterative method that can be used for the analysis of multivariate meta-analysis datasets that has no convergence problems and does not require the use of within-study correlations. The strategy is to use standard univariate methods for the marginal effects but also provides valid joint inference for multiple parameters. This method method can directly handle missing outcomes under missing completely at random assumption. This moment of method for MMA is specified via method argument (method="nn.mom")

Improved method for Riley model for MMA with continuous outcomes

Riley et al.(2008) proposed a working model and an overall synthesis correlation parameter to account for the marginal correlation between outcomes, where the only data needed are those required for a separate univariate random-effects meta-analysis. As within-study correlations are not required, the Riley method is applicable to a wide variety of evidence synthesis situations. However, the standard variance estimator of the Riley method is not entirely correct under many important settings. As a consequence, the coverage of a function of pooled estimates may not reach the nominal level even when the number of studies in the multivariate meta-analysis is large. Hong et al. (2015) improved the Riley method by proposing a robust variance estimator, which is asymptotically correct even when the model is misspecified (i.e., when the likelihood function is incorrect). The improved method for Riley model MMA is specified via method argument (method="nn.rs")

Marginal bivariate normal model for MMA with binary outcomes

Diagnostic systematic review is a vital step in the evaluation of diagnostic technologies. In many applications, it involves pooling pairs of sensitivity and specificity of a dichotomized diagnostic test from multiple studies. Chen et al. (2014) proposed a composite likelihood method for bivariate meta-analysis in diagnostic systematic reviews. The idea of marginal bivariate normal model for MMA with binary outcomes is to construct a composite likelihood (CL) function by using an independent working assumption between sensitivity and specificity. There are three immediate advantages of using this CL method. First, the non-convergence or non positive definite covariance matrix problem is resolved since there is no correlation parameter involved in the CL. Secondly, because the two-dimensional integration involved in the standard likelihood is substituted by one-dimensional integrals, the approximation errors are substantially reduced. Thirdly, the inference based on the CL only relies on the marginal normality of logit sensitivity and specificity. Hence the proposed method can be more robust than the standard likelihood inference to mis-specifications of the joint distribution assumption. This method is specified via method argument (method="bn.c1")
Marginal beta-binomial model for MMA with binary outcomes

When conducting a meta-analysis of studies with bivariate binary outcomes, challenges arise when the within-study correlation and between-study heterogeneity should be taken into account. Chen et al. (2015) proposed a marginal beta-binomial model for the meta-analysis of studies with binary outcomes. This model is based on the composite likelihood approach, and has several attractive features compared to the existing models such as bivariate generalized linear mixed model (Chu and Cole, 2006) and Sarmanov beta-binomial model (Chen et al., 2012). The advantages of the proposed marginal model include modeling the probabilities in the original scale, not requiring any transformation of probabilities or any link function, having closed-form expression of likelihood function, and no constraints on the correlation parameter. More importantly, since the marginal beta-binomial model is only based on the marginal distributions, it does not suffer from potential misspecification of the joint distribution of bivariate study-specific probabilities. Such misspecification is difficult to detect and can lead to biased inference using current methods. This method is specified via method argument (method="bb.cl")

Hybrid model for disease prevalence along with sensitivity and specificity for diagnostic test accuracy

Meta-analysis of diagnostic test accuracy often involves mixture of case-control and cohort studies. The existing bivariate random effects models, which jointly model bivariate accuracy indices (e.g., sensitivity and specificity), do not differentiate cohort studies from case-control studies, and thus do not utilize the prevalence information contained in the cohort studies. The trivariate generalized linear mixed models are only applicable to cohort studies, and more importantly, they assume the common correlation structure across studies, and the trivariate normality on disease prevalence, test sensitivity and specificity after transformation by some pre-specified link functions. In practice, very few studies provide justifications of these assumptions, and sometimes these assumptions are violated. Chen et al. (2015) evaluated the performance of the commonly used random effects model under violations of these assumptions and propose a simple and robust method to fully utilize the information contained in case-control and cohort studies. The proposed method avoids making the aforementioned assumptions and can provide valid joint inferences for any functions of overall summary measures of diagnostic accuracy. This method is specified via method argument (method="tb.cl")

Trivariate model for multivariate meta-analysis of diagnostic test accuracy

The standard methods for evaluating diagnostic accuracy only focus on sensitivity and specificity and ignore the information on disease prevalence contained in cohort studies. Consequently, such methods cannot provide estimates of measures related to disease prevalence, such as population averaged or overall positive and negative predictive values, which reflect the clinical utility of a diagnostic test. Chen et al. (2014) proposed a hybrid approach that jointly models the disease prevalence along with the diagnostic test sensitivity and specificity in cohort studies, and the sensitivity and specificity in case-control studies. In order to overcome the potential computational difficulties in the standard full likelihood inference of the proposed hybrid model, an alternative inference procedure was proposed based on the composite likelihood. Such composite likelihood based inference does not suffer computational problems and maintains high relative efficiency. In addition, it is more robust to model mis-specifications compared to the standard full likelihood inference. This method is specified via method argument (method="tn.cl")
Author(s)

Yong Chen, Yulun Liu

References


Examples

data(prostate)
fit.nn=mmeta(data=prostate, type="continuous", k=2, method="nn.cl")
summary(fit.nn)

rhow=runif(dim(prostate)[1], -0.2, 0.8)
fit.reml=mmeta(data=prostate, rhow=rhow, type="continuous", k=2, method="nn.reml")
print(fit.reml)

data(nat2)
fit.bb=mmeta(data=nat2, type="binary", k=2, method="bb.cl")
summary(fit.bb)

data(ca125)
fit.tb=mmeta(data=ca125, type="binary", k=2, method="tb.cl")
summary(fit.tb)

Description

Testing and correcting for small study effects of multivariate meta-analysis
**Usage**

```r
msset(data, nm.y1, nm.s1, nm.y2, nm.s2, method, type, k)
```

**Arguments**

- `data` : dataset
- `nm.y1` : column name for outcome 1
- `nm.s1` : column name for standard error of outcome 1
- `nm.y2` : column name for outcome 2
- `nm.s2` : column name for standard error of outcome 2
- `method` : "nn.cl" indicating the score test for detecting small study effects of MMA
- `type` : either "continuous" or "binary" indicating the type of outcomes
- `k` : integer indicating the number of outcomes

**Details**

This function returns the test statistics for testing small study effects of multivariate meta-analysis using regression method.

**Value**

`msset.TS` returns the test statistic and p value of the score test.

**A score test for detecting small study effects in multivariate meta-analysis**

Small study effects occur when smaller studies show different, often larger, treatment effects than large ones, which may threaten the validity of systematic reviews and meta-analyses. Detecting small study effects in a multivariate meta-analysis setting remains an untouched research area. Hong et al. (2019) propose a pseudolikelihood-based score test for detecting small study effects in multivariate random-effects meta-analysis. This is the first test for detecting small study effects in multivariate meta-analysis setting.

**Author(s)**

Chuan Hong

**References**


**Examples**

```r
data(prostate)
fit.msset=msset(data=prostate, nm.y1="y1", nm.s1="s1", nm.y2="y2", nm.s2="s2",
method = "nn.cl", type = "continuous", k=2)
summary(fit.msset)
```
A meta-analysis of the association between N-acetyltransferase 2 acetylation status and colorectal cancer

Description

A meta-analysis of 20 published case-control studies from January 1985 to October 2001

Format

The data frame contains the following columns:

- \( y_1 \): acetylator status (exposed) in control group
- \( n_1 \): total number of subjects in control group
- \( y_2 \): acetylator status (exposed) in case group
- \( n_2 \): total number of subjects in case group

Note

The dataset nat2 is used to conduct marginal bivariate normal model for MMA with binary outcomes

References


See Also

\texttt{mmeta, summary.mmeta}

Examples

data(nat2)
summary(nat2)
Comparison between overall survival and disease-free survival for prostate cancer

Description

Results from five randomized clinical trials published between 1988 and 2011

Format

The data frame contains the following columns:

- **y1**: log-hazard ratio estimates comparing combined therapy using Goserelin acetate with radiotherapy with respect to overall survival
- **s1**: within-study standard error for outcome 1
- **y2**: log-hazard ratio estimates comparing combined therapy using Goserelin acetate with radiotherapy with respect to disease-free survival
- **s2**: within-study standard error for outcome 2

Note

The dataset prostate is used to conduct bivariate random-effects meta-analysis when the within-study correlations are unknown.

References


See Also

- `mmeta`, `summary.mmeta`

Examples

```r
data(prostate)
summary(prostate)
```
**sim_dat**  
*Simulated data*

**Description**
A simulated dataset for galaxy function

**Format**
The data frame contains the following columns:

- **study.id**  
  study id
- **y1**  
  effect size for the first outcome
- **s1**  
  within-study standard error for the first outcome
- **y2**  
  effect size for the second outcome
- **s2**  
  within-study standard error for the second outcome
- **corr**  
  within-study correlation
- **subgroup**  
  subgroup of the studies

**Note**
The dataset `sim_dat` is used to illustrate the galaxy plot.

**See Also**
galaxy

**Examples**
```r
data(sim_dat)
summary(sim_dat)
```

---

**summary.mmeta**  
*Summarize the objects mmeta*

**Description**
Summarize a model of class mmeta fitted by mmeta.

**Usage**
```r
## S3 method for class 'mmeta'
summary(object,...)
```
Arguments

object  
an object inheriting from class mmeta.

...  
additional arguments; currently none is used.

Value

A list with the following components: coefficients, covariance matrix.

References


See Also

mmeta

Examples

data(prostate)
fit.nn=mmeta(data=prostate, type="continuous", k=2, method="nn.cl")
summary(fit.nn)

summary.msset Summarize the objects msset

Description

Summarize a model of class msset fitted by msset.
Usage

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

Arguments

- `object` an object inheriting from class `msset`.
- `...` additional arguments; currently none is used.

Value

A list with the following components: test statistics (msset) and p-value.

References


See Also

- `msset`

Examples

```r
data(prostate)
fit.msset=msset(data=prostate, nm.y1="y1", nm.s1="s1", nm.y2="y2", nm.s2="s2",
method = "nn.cl", type = "continuous", k=2)
summary(fit.msset)
```

---

**trimfill_rma**

*Trim&fill method for univariate meta analysis*

Description

Modified metafor::trimfill.rma.uni to avoid the invalid sqrt in k0 calculation when estimator == "Q0"

Usage

```r
trimfill_rma(
  x,
  side,
  estimator = "L0",
  maxiter = 100,
  method.trim = NULL,
  method.fill = NULL,
  verbose = FALSE,
  ilim
)
```
Arguments

- **x**: an object of class "rma.uni".
- **side**: the same as in metafor::trimfill
- **estimator**: the same as in metafor::trimfill
- **maxiter**: the same as in metafor::trimfill
- **method.trim**: the model used in rma.uni() for estimating the center when trimming studies, default is x$method
- **method.fill**: the model used in rma.uni() for estimating the center after filling studies, default is x$method
- **verbose**: the same as in metafor::trimfill
- **ilim**: limits for the imputed values as in metafor::trimfill. If unspecified, no limits are used.

Details

It is recommend using fixed-effects for method.trim and random-effects for method.fill when heterogeneity exists.

Value

the same as in metafor::trimfill

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Index

* Multivariate meta-analysis
  msset, 14

* Score test
  msset, 14

* Small study effects
  msset, 14

* datasets
  ca125, 3
  nat2, 16
  prostate, 17
  sim_dat, 18

* meta-analysis of diagnostic accuracy study
  mmeta, 10

* multivariate meta-analysis
  mmeta, 10

* package
  xmeta-package, 2

* random-effects
  mmeta, 10

* summary
  summary.mmeta, 18
  summary.msset, 19

ca125, 3

dat.gen, 4

galaxy, 3, 5, 18
  galaxy.trimfill, 7

mmeta, 2, 4, 10, 16, 17, 19
msset, 3, 14, 20

nat2, 16

prostate, 17

sim_dat, 18
summary.mmeta, 4, 16, 17, 18
summary.msset, 19

trimfill_rma, 20

xmeta (xmeta-package), 2
xmeta-package, 2

22