

Package ‘dosresmeta’

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Title Multivariate Dose-Response Meta-Analysis

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Description Estimates dose-response relations from summarized dose-response data and to combines them according to principles of (multivariate) random-effects models.

URL <https://alecric.github.io/software/dosresmeta.html>

BugReports <https://github.com/alecric/dosresmeta/issues>

License GPL-2

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dosresmeta-package *Multivariate dose-response meta-analysis*

Description

It consists of a collection of functions to estimate dose-response relations from summarized dose-response data for both continuous and binary outcomes, and to combine them according to principles of (multivariate) random-effects model.

Modeling framework

Dose-response meta-analysis represents a specific type of meta-analysis. Aim of such analysis is to reconstruct and combine study-specific curves from summarized dose-response data. Greenland and Longnecker originally developed the methodology in 1992 for pooling associations from epidemiological studies of binary outcomes. Extensions are currently proposed for other types of outcomes (e.g. continuous) from others study design, such as clinical trials.

The summarized dose-response data are most often presented in a tabular way, reporting the levels of the exposure (doses) and the corresponding outcome variable. The latter is usually expressed as contrast to the unexposed or baseline category (referent level). Examples are (log) relative risks, (log) odds ratios, (log) incidence rate ratios, mean differences, and standardized mean differences. Thus the outcome cannot be regarded as independent and a (co)variance matrix needs to be provided or approximated from the available data. See `covar.smd` and `covar.logrr` for more details.

Estimation procedure

The pooled dose-response association can be estimated using two different approaches. The former consists of a two-stage procedure, where the study-specific trend are first estimated and then pooled across studies. Assuming y_j is the vector of non-referent outcome values in each of $i = 1, \dots, m$ studies, and X_i the related matrix of p transformations of the exposure (typically $p = 1, 2$), the dose-response model can be written as

$$y_i = X_i \beta_i + \epsilon_i$$

with $S_i = (\text{co})\text{variance}$ of ϵ_i known (available or reconstructed from the available data). The β_i are then combined according to principles of (multivariate) random-effects meta-analytical models

$$\beta_i \sim N(\beta, V_i + \Psi)$$

where V_i and Ψ indicate, respectively, the within study (co)variance (obtained in the first stage analysis) and the between study (co)variance.

The alternative approach, instead, consists of a one-stage (also known as pool-first) procedure. The data are pooled by concatenating the vector y_i and vectors (or matrices) X_i . The (multivariate) random effects-model can be written as

$$y_i = X_i \beta + Z_i \eta_i + \epsilon_i$$

where β represents the fixed-effects parameter, η_i the vector (or matrix) of unobserved random-effects for the i -th study, and Z_i coincides with X_i . The marginal model has a co(variance) matrix equal to $\Sigma + Z_i \Psi Z_i^t$, where Σ is the block diagonal (co)variance with i -th diagonal block S_i .

The two approaches provide similar results, despite the two-stage procedure may be more stable and faster in terms of convergence. In both the procedures the aim is to estimate the coefficients β and, for random-effects models, the components of the between-study (co)variance matrix Ψ . Different estimators are implemented in the package. The estimation options available are

- Fixed-effects
- Maximum likelihood (ML)
- Restricted maximum likelihood (REML)
- Method of moments (currently available only for the two-stage procedure)

The fixed-effects model is fitted through generalized least squares (GLS), assuming the (co)variance structure, composed by the within-study error only, as completely known. Among random-effects models, ML and REML approaches provides fit criteria and inferential test derived from likelihood theory, such as AIC and likelihood ratio test, particularly useful in a one-stage procedure. Further details on estimation methods are given in the related help pages.

Functions and data included in the package

The structure of the package and the internal functions resemble those of the `mixmeta` package. See `mixmeta-package` for a general overview. The main function is `dosresmeta`, which performs the various models illustrated above. The function returns a list object of class "dosresmeta" (see `dosresmetaObject`).

The estimation is carried out internally through `dosresmeta.fit`, a wrapper which prepares the data and calls specific estimation functions for fitting the models, depending on the chosen procedure. For the two-stage procedure, the second part of the analysis is performed using the function `mixmeta.fit` while estimators for random-effects models are implemented in the functions `dosresmeta.ml` and `dosresmeta.reml` for (restricted) maximum likelihood. For likelihood-based methods, iterative optimizations algorithms are used for maximizing the (restricted) likelihood. Fitting parameter options are set by `dosresmeta.control`.

Method functions are available for objects of class "dosremeta" (see `dosresmetaObject` for a complete list). The method `summary` produces a list of class "summary.dosremeta" for summarizing the fit of the model and providing additional results. The method function `predict` computes predicted values, optionally for a set of new values of the predictors. `blup` gives the (empirical) best linear unbiased predictions for the unobserved random-effects. Other default or specific method functions for regression can be used on objects of class "dosremeta", such as `logLik`, `AIC` and `BIC`, among others. The method function `qttest.dosresmeta` (producing an object with class of the same name) performs the Cochran Q test for (residual) heterogeneity currently appropriate only for the two-stage approach.

Printing functions for the objects of classes defined above are also provided. Other functions are used internally in the source code, and not exported in the namespace. For users interested in getting into details of the package structure, these functions can be displayed using the triple colon (`':::'`) operator. For instance, `dosresmeta:::glsfit` displays the code of the function `glsfit`.

The package includes the datasets `alcohol_crc`, `alcohol_cvd`, `ari`, and `cc_ex` as data frames, which are used in the examples.

Use `citation("dosresmeta")` to cite this package.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

References

- Alessio Crippa, Nicola Orsini (2016). Multivariate Dose-Response Meta-Analysis: The dosresmeta R Package. *Journal of Statistical Software, Code Snippets*, 72(1), 1-15.doi:10.18637/jss.v072.c01
- Greenland, S., Longnecker, M. P. (1992). Methods for trend estimation from summarized dose-response data, with applications to meta-analysis. *American journal of epidemiology*, 135(11), 1301-1309.

Orsini, N., Bellocco, R., Greenland, S. (2006). Generalized least squares for trend estimation of summarized dose-response data. *Stata Journal*, 6(1), 40.

Orsini, N., Li, R., Wolk, A., Khudyakov, P., Spiegelman, D. (2012). Meta-analysis for linear and nonlinear dose-response relations: examples, an evaluation of approximations, and software. *American journal of epidemiology*, 175(1), 66-73.

Gasparrini, A., Armstrong, B., Kenward, M. G. (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*, 31(29), 3821-3839.

See Also

[dosresmeta](#) [mixmeta](#)

alcohol_crc

Eight published studies on the relation between alcohol intake and colorectal cancer

Description

The dataset reports the summarized dose-response results from eight prospective studies on the relation between alcohol intake and colorectal cancer risk.

Format

A data frame with 48 observations on the following 7 variables:

id	label for author's names (id variable).
type	code for study design.
dose	assigned dose levels.
cases	number of cases for each exposure level.
peryears	amount of person-time for each exposure level.
logrr	natural logarithm of the adjusted "relative risks".
se	standard error for the logarithm of the adjusted "relative risks".

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Orsini, N., Li, R., Wolk, A., Khudyakov, P., Spiegelman, D. (2012). Meta-analysis for linear and nonlinear dose-response relations: examples, an evaluation of approximations, and software. *American journal of epidemiology*, 175(1), 66-73.

alcohol_cvd	<i>Six published studies on the relation between alcohol intake and cardiovascular disease risk</i>
-------------	---

Description

The dataset reports the summarized dose-response results from six observational studies on the relation between alcohol intake and vascular disease risk. Four are case-control studies, two prospective (cumulative-incidence data).

Format

A data frame with 25 observations on the following 8 variables:

id	id of the studies included in the analysis.
author	names of the first author of the study.
type	code for study design.
dose	assigned dose levels.
cases	number of cases for each exposure level.
n	total number of subjects for each exposure level.
logrr	natural logarithm of the adjusted "relative risks".
se	standard error for the logarithm of the adjusted "relative risks".

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Liu, Q., Cook, N. R., Bergstrom, A., Hsieh, C. C. (2009). A two-stage hierarchical regression model for meta-analysis of epidemiologic nonlinear dose-response data. *Computational Statistics & Data Analysis*, 53(12), 4157-4167.

alcohol_esoph	<i>Fourteen case-control studies on the relation between alcohol consumption and esophageal cancer</i>
---------------	--

Description

The dataset reports the summarized dose-response results from fourteen case-control studies on the relation between alcohol consumption and esophageal squamous cell carcinoma.

Format

A data frame with 63 observations on the following 8 variables:

id	id of the studies included in the analysis.
author	names of the first author.
type	code for study design.
cases	number of cases for each exposure level.
n	total number of subjects for each exposure level.
dose	assigned dose levels.
logrr	natural logarithm of the adjusted odds ratio.
se	standard error for the logarithm of the adjusted odds ratio

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Rota M, Bellocco R, Scotti L, Tramacere I, Jenab M, Corrao G, La Vecchia C, Boffetta P, Bagnardi V. Random-effects meta-regression models for studying nonlinear dose-response relationship, with an application to alcohol and esophageal squamous cell carcinoma. *Statistics in medicine*. 2010 Nov 20;29(26):2679-87.

alcohol_lc	<i>Four published studies on the relation between alcohol intake and lung cancer</i>
------------	--

Description

The dataset reports the summarized dose-response results from four prospective studies on the relation between alcohol intake and lung cancer.

Format

A data frame with 20? observations on the following 7 variables:

id	label for author's names (id variable).
type	code for study design.
dose	assigned dose levels.
cases	number of cases for each exposure level.
peryears	amount of person-time for each exposure level.
logrr	natural logarithm of the adjusted "relative risks".
se	standard error for the logarithm of the adjusted "relative risks".

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Orsini, N., Li, R., Wolk, A., Khudyakov, P., Spiegelman, D. (2012). Meta-analysis for linear and nonlinear dose-response relations: examples, an evaluation of approximations, and software. *American journal of epidemiology*, 175(1), 66-73.

ari	<i>Five clinical trials on the relation between aripiprazole and schizophrenia</i>
-----	--

Description

The dataset reports the summarized dose-response results from five clinical trials on the relation between different levels of aripiprazole and severity of schizophrenia measured using the PANSS medical score.

Format

A data frame with 18 observations on the following 6 variables:

id	id of the studies included in the analysis.
author	names of the first author of the studies.
dose	assigned dose level of aripiprazole (0 for placebo group).
y	outcome variable: change in PANNS score after and before treatment.
sd	standard deviation of y for each exposure level.
n	total number of subjects for each exposure level.

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Crippa, A., Orsini, N. Dose-response meta-analysis of differences in means. *BMC medical research methodology*. 2016 Aug 2;16(1):91.

blup.dosresmeta	<i>Best Linear Unbiased Predictions from dosresmeta Models</i>
-----------------	--

Description

This method function computes (empirical) best linear unbiased predictions from fitted dose-response meta-analytical models represented in objects of class "dosresmeta".

Usage

```
## S3 method for class 'dosresmeta'  
blup(object, ...)
```

Arguments

object	objects of classe "dosresmeta".
...	further arguments passed to or from other methods.

Details

The method function blup produces (empirical) best linear unbiased predictions from dosresmeta objects. Predictions are expressed in terms of study-specific deviations as random effects. Predicted random effects from blup are a shrunk version of study-specific realizations, where study-specific predictions borrow strength from the assumption of an underlying distribution in a (usually hypothetical) population of studies. Blup are not available for fixed-effects models since they are meaningless in that context.

Examples

```
## Load data and run the linear and quadratic models  
data("alcohol_cvd")  
lin <- dosresmeta(formula = logrr ~ dose, type = type, id = id,  
                 se = se, cases = cases, n = n, data = alcohol_cvd)  
quadr <- dosresmeta(formula = logrr ~ dose + I(dose^2), type = type, id = id,  
                   se = se, cases = cases, n = n, data = alcohol_cvd)  
  
## blup prediction for the previous models  
blup(lin)  
blup(quadr)
```

bmi_rc	<i>Four case-control studies on the relation between Body Mass Index and renal cell cancer</i>
--------	--

Description

The dataset reports the summarized dose-response results from four cases-control studies on the relation Body Mass Index and renal cell cancer

Format

A data frame with 33 observations on the following 13 variables:

id	id of the studies included in the analysis.
author	names of the first author and year of publication.
type	code for study design.
interval	intervals for the categories of bmi.
bmi	assigned bmi levels.
case	number of cases for each exposure level.
control	number of controls for each exposure level.
n	total number of subjects for each exposure level.
or	adjusted odds ratios for each exposure level.
lb_or	lower bound for the confidence limits of the adjusted odds ratios.
ub_or	upper bound for the confidence limits of the adjusted odds ratios.
logor	natural logarithm of the adjusted odds ratios.
se_logor	standard error for the logarithm of the adjusted odds ratios.

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Liu, Q., Cook, N. R., Bergstrom, A., Hsieh, C. C. (2009). A two-stage hierarchical regression model for meta-analysis of epidemiologic nonlinear dose-response data. *Computational Statistics & Data Analysis*, 53(12), 4157-4167.

cc_ex	<i>Case-control data on alcohol and breast cancer risk</i>
-------	--

Description

The dataset reports the summarized dose-response results from a case-control study on alcohol and breast cancer, first presented by Rohan and McMichael.

Format

A data frame with 4 observations on the following 10 variables:

gday	label for exposure levels.
dose	assigned dose levels.
case	number of cases for each exposure level.
control	number of controls for each exposure level.
n	total number of subjects for each exposure level.
crudeor	unadjusted odds ratios for each exposure level.
adjrr	adjusted odds ratios for each exposure level.
lb	lower bound for the confidence limits of the adjusted odds ratios.
ub	upper bound for the confidence limits of the adjusted odds ratios.
logrr	natural logarithm of the adjusted odds ratios.

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Rohan, T. E., McMichael, A. J. (1988). Alcohol consumption and risk of breast cancer. *International journal of cancer*, 41(5), 695-699.

Greenland, S., Longnecker, M. P. (1992). Methods for trend estimation from summarized dose-response data, with applications to meta-analysis. *American journal of epidemiology*, 135(11), 1301-1309.

ci_ex	<i>Cumulative incidence data on high-fat dairy food and colorectal cancer risk</i>
-------	--

Description

The dataset reports the summarized dose-response results from a cumulative-incidence study on high-fat dairy food intake and risk of colorectal cancer, first presented by Larsson, Bergkvist, and Wolk (2005).

Format

A data frame with 5 observations on the following 8 variables:

dose	assigned dose levels.
case	number of cases for each exposure level.
n	total number of subjects for each exposure level.
adjrr	adjusted risk ratios for each exposure level.
lb	lower bound for the confidence limits of the adjusted risk ratios.

ub upper bound for the confidence limits of the adjusted risk ratios.
 logrr natural logarithm of adjusted risk ratios.
 se standard error for the logarithm of the adjusted risk ratios.

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Larsson, S. C., L. Bergkvist, and A. Wolk. (2005). High-fat dairy food and conjugated linoleic acid intakes in relation to colorectal cancer incidence in the Swedish Mammography Cohort. *American Journal of Clinical Nutrition* 82: 894-900.

Greenland, S., Longnecker, M. P. (1992). Methods for trend estimation from summarized dose-response data, with applications to meta-analysis. *American journal of epidemiology*, 135(11), 1301-1309.

coef.dosresmeta	<i>Extract Coefficients and (Co)Variance Matrix from dosresmeta Objects</i>
-----------------	---

Description

These method functions return the estimated fixed-effects coefficients and their (co)variance matrix for fitted dose-response meta-analytical models represented in objects of class "dosresmeta".

Usage

```
## S3 method for class 'dosresmeta'
coef(object, format = c("vector", "matrix"), ...)
```

```
## S3 method for class 'dosresmeta'
vcov(object, ...)
```

Arguments

object an object of class "dosresmeta".
 format format of the returned object.
 ... further arguments passed to or from other methods.

Value

For coef, a vector (default) or matrix with the estimated (fixed-effects) coefficients. For vcov, the (co)variance matrix of the estimated (fixed-effects) coefficients.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

See Also

[dosresmeta](#), [coef](#), [vcov](#), [logLik.dosresmeta](#)

Examples

```
## Load data and run the model
data("alcohol_cvd")
model <- dosresmeta(formula = logrr ~ dose + I(dose^2), type = type, id = id,
                   se = se, cases = cases, n = n, data = alcohol_cvd)

## Fixed-effect coefficients
coef(model)

## Fixed-effect (co)variance matrix
vcov(model)
```

coffee_cancer

Eight prospective studies on the relation between coffee consumption and cancer mortality

Description

The dataset reports the summarized dose-response results from eight prospective studies on the relation between coffee consumption and cancer mortality.

Format

A data frame with 59 observations on the following 11 variables:

id	id of the studies included in the analysis.
author	names of the first author.
year	year of publication.
type	code for study design.
dose	assigned dose levels.
cases	number of cases for each exposure level.
n	total number of subjects (type = "ci") or person-years (type = "ir") for each exposure level.
logrr	natural logarithm of the adjusted risk estimates.
se	standard error for the logarithm of the adjusted risk estimates.
gender	factor variable for the gender of the participants.
area	factor variable for the study location.

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Crippa A, Discacciati A, Larsson SC, Wolk A, Orsini N. Coffee Consumption and Mortality from All Causes, Cardiovascular Disease, and Cancer: A Dose-Response Meta-Analysis. *Am J Epidemiol.* 2014 Aug 24. pii: kwu194.

coffee_cvd

Thirteen prospective studies on the relation between coffee consumption and cardiovascular mortality

Description

The dataset reports the summarized dose-response results from thirteen prospective studies on the relation between coffee consumption and cardiovascular mortality.

Format

A data frame with 100 observations on the following 12 variables:

id	id of the studies included in the analysis.
author	names of the first author.
year	year of publication.
type	code for study design.
dose	assigned dose levels.
cases	number of cases for each exposure level.
n	total number of subjects (type = "ci") or person-years (type = "ir") for each exposure level.
logrr	natural logarithm of the adjusted risk estimates.
se	standard error for the logarithm of the adjusted risk estimates.
gender	character variable for the gender of the participants.
area	character variable for the study location.
smoking	character variable for the type of smoking adjustment.

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Crippa A, Discacciati A, Larsson SC, Wolk A, Orsini N. Coffee Consumption and Mortality from All Causes, Cardiovascular Disease, and Cancer: A Dose-Response Meta-Analysis. *Am J Epidemiol.* 2014 Aug 24. pii: kwu194.

coffee_mort	<i>Twenty-one prospective studies on the relation between coffee consumption and all-cause mortality</i>
-------------	--

Description

The dataset reports the summarized dose-response results from twenty-one prospective studies on the relation between coffee consumption and all-cause mortality.

Format

A data frame with 109 observations on the following 11 variables:

id	id of the studies included in the analysis.
author	names of the first author.
year	year of publication.
type	code for study design.
dose	assigned dose levels.
cases	number of cases for each exposure level.
n	total number of subjects (type = "ci") or person-years (type = "ir") for each exposure level.
logrr	natural logarithm of the adjusted risk estimates.
se	standard error for the logarithm of the adjusted risk estimates.
gender	factor variable for the gender of the participants.
area	factor variable for the study location.

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Crippa A, Discacciati A, Larsson SC, Wolk A, Orsini N. Coffee Consumption and Mortality from All Causes, Cardiovascular Disease, and Cancer: A Dose-Response Meta-Analysis. *Am J Epidemiol.* 2014 Aug 24. pii: kwu194.

coffee_mort_add	<i>Additional two prospective studies on the relation between coffee consumption and all-cause mortality</i>
-----------------	--

Description

The dataset reports the summarized dose-response results from two additional prospective studies on the relation between coffee consumption and all-cause mortality. The studies do not report information on the number of cases and participants/person-time.

Format

A data frame with 109 observations on the following 11 variables:

id	id of the studies included in the analysis.
author	names of the first author.
year	year of publication.
type	code for study design.
dose	assigned dose levels.
cases	number of cases for each exposure level.
n	total number of subjects (type = "ci") or person-years (type = "ir") for each exposure level.
logrr	natural logarithm of the adjusted risk estimates.
se	standard error for the logarithm of the adjusted risk estimates.
gender	factor variable for the gender of the participants.
area	factor variable for the study location.

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Crippa A, Discacciati A, Larsson SC, Wolk A, Orsini N. Coffee Consumption and Mortality from All Causes, Cardiovascular Disease, and Cancer: A Dose-Response Meta-Analysis. *Am J Epidemiol*. 2014 Aug 24. pii: kwu194.

coffee_stroke	<i>Eleven prospective studies on the relation between coffee consumption and stroke risk</i>
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Description

The dataset reports the summarized dose-response results from eleven prospective studies on the relation between coffee consumption and risk of stroke.

Format

A data frame with 68 observations on the following 12 variables:

id	id of the studies included in the analysis.
author	names of the first author of the studies.
type	code for study design.
dose	assigned dose levels.
cases	number of cases for each exposure level.
n	total number of subjects (type = "ci") or person-years (type = "ir") for each exposure level.
rr	adjusted risk estimates for each exposure level.
lb	lower bound for the confidence limits of the adjusted risk estimates.

ub upper bound for the confidence limits of the adjusted risk estimates.
 logrr natural logarithm of the adjusted risk estimates.
 se standard error for the logarithm of the adjusted risk estimates.
 nordic indicator variable for the study to be conducted in the nordic countries (1 = yes).

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Larsson, S. C., Orsini, N. (2011). Coffee consumption and risk of stroke: a dose-response meta-analysis of prospective studies. *American journal of epidemiology*, 174(9), 993-1001.

covar.logrr	<i>Computes the covariance matrix for a set of log relative risks</i>
-------------	---

Description

Reconstructs the covariance matrix for a set of (reported) log relative risks, given the number of cases and the number of total persons or person-years for each treatment (dose) level.

Usage

```
covar.logrr(cases, n, y, v, type, data, covariance = "gl")
```

Arguments

cases	a vector, defining the number of cases for each exposure level.
n	a vector, defining the total number of subjects for each exposure level. For incidence-rate data n indicates the amount of person-time within each exposure level.
y	a vector, defining the (reported) log relative risks.
v	a vector, defining the variances of the reported log relative risks.
type	a vector (or a character string), specifying the design of the study. Options are <code>cc</code> , <code>ir</code> , and <code>ci</code> , for case-control, incidence-rate, and cumulative incidence data, respectively.
data	an optional data frame (or object coercible by as.data.frame to a data frame) containing the variables in the previous arguments.
covariance	method to approximate the covariation among set of reported log relative risks, "gl" for the method proposed by Greenland and Longnecker (default), "h" for the method proposed by Hamling.

Details

This is an internal function called by [dosresmeta](#) to reconstruct the (co)variance matrix of the (adjusted) log relative risks. The function calls, depending on the chosen method, [gr1](#) (default) or [hamling](#) to reconstruct the effective counts corresponding to the (adjusted) log relative risks as well as their standard errors. From these it computes the covariance matrix; analytical formulas can be found in the referenced article.

Value

The (co)variance matrix of the log relative risks.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

References

Orsini, N., Li, R., Wolk, A., Khudyakov, P., Spiegelman, D. (2012). Meta-analysis for linear and nonlinear dose-response relations: examples, an evaluation of approximations, and software. *American journal of epidemiology*, 175(1), 66-73.

See Also

[gr1](#), [hamling](#), [covar.smd](#), [dosresmeta](#)

Examples

```
## Loading data
data("alcohol_cvd")

## Obtaining the (co)variance matrix of log RR for the first study (id = 1)
covar.logrr(y = logrr, v = I(se^2), cases = cases, n = n, type = type,
            data = subset(alcohol_cvd, id == 1))

## Obtaining the (co)variance matrices of log RR for all study
by(alcohol_cvd, alcohol_cvd$id, function(x)
   covar.logrr(y = logrr, v = I(se^2), cases = cases, n = n,
               type = type, data = x))

## Restructuring the previous results in a list of matrices
do.call("list", by(alcohol_cvd, alcohol_cvd$id, function(x)
   covar.logrr(y = logrr, v = I(se^2), cases = cases, n = n, type = type,
               data = x)))
```

covar.smd	<i>Computes mean and standardized mean differences for continuous outcome with corresponding co(variance) matrix</i>
-----------	--

Description

This internal function computes mean and standardized mean of a continuous outcome with the corresponding variances. It also reconstructs the covariance matrix from the available data.

Usage

```
covar.smd(y, sd, n, measure = "md", method = "cohens", data)
```

Arguments

y	a vector defining the mean outcome for each treatment level.
sd	a vector defining the standard deviation of the outcome for each treatment level.
n	a vector defining the number of subjects for each treatment level.
measure	character string, indicating the measure to be calculated. Options are md and smd for mean difference and standardized mean difference, respectively.
method	character string indicating the method to be used. Options are cohens, hedges, and glass.
data	an optional data frame (or object coercible by as.data.frame to a data frame) containing the variables in the previous arguments.

Details

This is an internal function called by [dosresmeta](#) to reconstruct the (co)variance matrix of the outcome variable. The function is expected to be extended and/or modified at every release of the package

Value

A list containing the following

y	mean or standardized mean differences for each treatment level, included the referent one (0 by calculation).
v	variances corresponding to the mean or standardized mean differences for each treatment level, included the referent one (0 by calculation).
S	co(variance) matrix for the non-referent mean or standardized mean differences.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

References

Cooper, H., Hedges, L. V., & Valentine, J. C. (Eds.). (2009). The handbook of research synthesis and meta-analysis. Russell Sage Foundation.

See Also

[covar.logrr](#), [dosresmeta](#)

Examples

```
## Loading the data
data("ari")

## Obtaining standardized mean differences, variances, and (co)varinace
## matrix for the first study (id = 1)
covar.smd(y, sd, n, measure = "smd", data = subset(ari, id == 1))

## Obtaining mean differences, variances, and (co)varinace matrices for the all the studies
cov.md <- by(ari, ari$id, function(x) covar.smd(y, sd, n, "md", data = x))

## Extracting mean differences
unlist(lapply(cov.md, function(x) x$y))
## Extracting variances for the mean differences
unlist(lapply(cov.md, function(x) x$v))
## List of the (co)variance matrices for the mean differences
lapply(cov.md, function(x) x$S)
```

dosresmeta

Multivariate Dose-Response Meta-Analysis

Description

The function `dosresmeta` estimates a dose-response curve from either single or multiple summarized dose-response data, taking into account the correlation among observations and heterogeneity across studies. The function `dosresmeta.fit` is a wrapper for actual fitting functions based on different estimation methods, usually called internally. See [dosresmeta-package](#) for an overview.

Usage

```
dosresmeta(
  formula,
  id,
  v,
  type,
  cases,
  n,
  sd,
```

```

    data,
    mod = ~1,
    intercept = F,
    center = T,
    se,
    lb,
    ub,
    covariance = "gl",
    method = "reml",
    proc = "2stage",
    Slist,
    method.smd = "cohen",
    control = list()
)

```

```
dosresmeta.fit(X, Z, y, Slist, id, method, control, proc, mod, v, data)
```

Arguments

formula	an object of class " formula " offering a symbolic description of the dose-response functional relation. Terms in the formula can be provided in the data below.
id	an vector to specify the id variable for the studies included in the analysis. Optional if estimating a dose-response model from a single study.
v	a vector to specify the variances of the reported outcome. Alternatively the user can provide the standard error in the se argument, or only for log relative risks, the confidence interval in the lb and ub arguments.
type	an optional vector (or a string) required when the outcome is log relative risks. It specifies the study-specific design. The values for case-control, incidence-rate, and cumulative incidence data are cc, ir, and ci, respectively.
cases	a vector to specify the number of cases for each exposure level. Required to reconstruct the (co)variance matrix for log relative risks.
n	a vector to specify the total number of subjects for each exposure level. Required to reconstruct the (co)variance matrix for log relative risks. For incidence-rate data n indicates the amount of person-time for each exposure level.
sd	a vector to specify the standard deviation. Required to reconstruct the (co)variance matrix for differences and standardized mean differences.
data	a data frame (or object coercible by <code>as.data.frame</code> to a data frame) containing the variables in the previous arguments.
mod	an object of class " formula " offering a symbolic description of the meta-regression model (by default <code>mod = ~ 1</code>). Terms in the formula can be provided in the data below.
intercept	a logical value to specify if an intercept term needs to be included in the model. See details.
center	a logical value to specify if the design matrix need to be center at the referent ones. See details.

se	an optional vector to specify the standard error of the reported log relative risks; needed if v is not provided.
lb	an optional vector to specify the lower bound of the confidence interval for the reported relative risks; needed if v and se are not provided.
ub	an optional vector to specify the upper bound of the confidence interval for the reported relative risks; needed if v and se are not provided.
covariance	method to approximate the (co)variance matrix of the outcome. Options are "gl" for the method proposed by Greenland and Longnecker (default) , "h" for the method proposed by Hamling, "md" for mean differences, "smd" for standardized mean differences, and "user" if provided by the user.
method	method used to estimate the (pooled) dose-response relation: "fixed" for fixed-effects models, "ml" or "reml" for random-effects models fitted through (restricted) maximum likelihood, and "mm" for random-effects models fitted through method of moments (currently available only for the two stages procedure).
proc	"2stage" (default) or "1stage" procedure. See dosresmeta-package for an overview.
Slist	list of approximated or given (co)variance matrices.
method.smd	character string indicating the method to be used. Options are cohens, hedges, and glass. Required only if covariance equal "smd".
control	list of parameters for controlling the fitting process. These are passed to dosresmeta.control by dosresmeta.fit to replace otherwise selected default values.
X	processed design matrix of fixed effects.
Z	processed design matrix of random effects.
y	processed outcome vector.

Details

The function defines all the elements required to estimate a dose-response association taking into account the correlation among the observations. If the (co)variance matrix is not provided then it is approximated depending of the type of outcome specified through the `covariance` argument. The dose-response model is specified in the `formula`. Typically the outcome is expressed as a contrast to a reference exposure level, so that the model does not have an intercept and the values in the design matrix need to be centered at the referent values, as described by Qin Liu et al, 2009. This is internally done, respectively, when `intercept = FALSE` and `center = TRUE` (default values).

The function calls the wrapper `dosresmeta.fit` to perform the actual fitting. The latter prepares the data and calls specific fitting functions, depending on the chosen procedure and method. For the two stages procedure, the second part of the analysis is performed using the function `mixmeta.fit` from the `mixmeta` package. Different estimator are implemented in the package. The estimation options available are

- Fixed-effects
- Maximum likelihood (ML)
- Restricted maximum likelihood (REML)
- Method of moments (currently available only for the two stage procedure)

The fitting procedure can be controlled through the additional terms specified in `control`, which are passed to the function [dosresmeta.control](#).

Value

The `dosresmeta` function typically returns a list of object of class `dosresmeta` representing the meta-analytical model fit, as described in [dosresmetaObject](#).

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

References

Greenland, S., Longnecker, M. P. (1992). Methods for trend estimation from summarized dose-response data, with applications to meta-analysis. *American journal of epidemiology*, 135(11), 1301-1309.

Orsini, N., Bellocco, R., Greenland, S. (2006). Generalized least squares for trend estimation of summarized dose-response data. *Stata Journal*, 6(1), 40.

Liu, Q., Cook, N. R., Bergstrom, A., Hsieh, C. C. (2009). A two-stage hierarchical regression model for meta-analysis of epidemiologic nonlinear dose-response data. *Computational Statistics & Data Analysis*, 53(12), 4157-4167.

Gasparri, A., Armstrong, B., Kenward, M. G. (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*, 31(29), 3821-3839.

See Also

[dosresmeta-package](#), [mixmeta](#), [covar.logrr](#), [covar.smd](#)

Examples

```
## First example: Single case-control study
## Linear trend estimation
data("cc_ex")

## Fitting the model
mod1 <- dosresmeta(formula = logrr ~ dose, type = "cc", cases = case,
                  n = n, lb = lb, ub = ub, data= cc_ex)
summary(mod1)
## Results
predict(mod1, delta = 1, expo = TRUE)

## Second example: Multiple studies
## Linear and quadratic trend using random-effects meta-analysis
data("alcohol_cvd")

## Linear trend
lin <- dosresmeta(formula = logrr ~ dose, type = type, id = id,
                 se = se, cases = cases, n = n, data = alcohol_cvd)
summary(lin)
## Predicted linear trend
predict(lin, delta = 1, expo = TRUE)
```

```

## Non-linear (quadratic) trend
quadr <- dosresmeta(formula = logrr ~ dose + I(dose^2), type = type, id = id,
                    se = se, cases = cases, n = n, data = alcohol_cvd)
summary(quadr)

## Graphical results
with(predict(quadr, expo = TRUE, order = TRUE), {
  plot(dose, pred, log = "y", type = "l",
        xlim = c(0, 45), ylim = c(.4, 2))
  lines(dose, ci.lb, lty = 2)
  lines(dose, ci.ub, lty = 2)
  rug(dose, quiet = TRUE)
})

## Third example: Continuous outcome (smd)
data("ari")
mod3 <- dosresmeta(formula = y ~ dose + I(dose^2), id = id,
                   sd = sd, n = n, covariance = "smd", data = ari)
summary(mod3)

## Graphical results
newdata <- data.frame(dose = seq(0, 30, 1))
with(predict(mod3, newdata, order = TRUE), {
  plot(dose, pred, type = "l",
        ylim = c(0, .6))
  lines(dose, ci.lb, lty = 2)
  lines(dose, ci.ub, lty = 2)
  rug(dose, quiet = TRUE)
})

```

dosresmeta.control *Ancillary Parameters for Controlling the Fit in dosresmeta Models*

Description

This internal function sets the parameter options used for fitting dose-response meta-analytical models, commonly to pre-specified default values. It is usually internally called by [dosresmeta.fit](#). The function has many more arguments than needed for dosresmeta. For more details see [mixmeta.control](#).

Usage

```

dosresmeta.control(
  optim = list(),
  showiter = FALSE,
  maxiter = 100,
  initPsi = NULL,
  Psifix = NULL,
  Scor = NULL,

```



```

    addSlist = NULL,
    inputna = FALSE,
    inputvar = 10^4,
    loglik.iter = "hybrid",
    igls.inititer = 10,
    gr = FALSE,
    hessian = FALSE,
    vc.adj = TRUE,
    reltol = sqrt(.Machine$double.eps),
    checkPD = NULL,
    set.negeigen = sqrt(.Machine$double.eps)
)

```

Arguments

optim	list of parameters passed to the control argument of the function optim, which performs the quasi-Newton optimization in likelihood-based random-effects models. See optim .
showiter	logical. If TRUE, the progress of iterative optimization is shown.
maxiter	positive integer value. Maximum number of iterations in methods involving optimization procedures.
initPsi	either a matrix or a vector of its lower triangular elements (with diagonal, taken by column) from which starting values of the parameters of the between-study (co)variance matrix are derived, used in the optimization procedure for likelihood-based random-effects models. If NULL (the default, and recommended), the starting value is created internally through an iterative generalized least square algorithm.
Psifix	either a matrix or a vector of its lower triangular elements (with diagonal, taken by column), or optionally a named list with one or more of such objects. Used to define fixed parts of the random-effects
Scor	either a scalar, vector or matrix representing the within-unit correlation(s) to be inputted when the covariances are not provided in multivariate models, and ignored if they are.
addSlist	a list of m matrices for the (outer-level) groups of units defining the (known) error (co)variance structure, when this cannot be passed through the argument S of mixmeta.
inputna	logical. If missing values must be internally inputted. To be used with caution.
inputvar	multiplier for inputting the missing variances, to be passed as an argument to inputna.
loglik.iter	iterative scheme used in in likelihood-based optimization routines. Options are "hybrid", "newton", and "igls" or "RIGLS".
igls.inititer	number of iteration of the iterative generalized least square algorithm to be run in the hybrid optimization procedure of linkelihood-based models to provide the starting value.
gr	indicates if the gradient of the (re)ml likelihood should be provided. FALSE by default.

hessian	logical. If TRUE, the Hessian matrix of the parameters estimated in the optimization process is computed and returned. Only applicable to likelihood-based estimation methods.
vc.adj	logical. If TRUE, an adjustment to the way the marginal variance part is computed in the (co)variance components estimator is applied in the variance components estimator.
reltol	relative convergence tolerance in methods involving optimization procedures. The algorithm stops if it is unable to reduce the value by a factor of $reltol * (abs(val) + reltol)$ at a step.
checkPD	logical. Determines if the semi-positiveness of within-unit error or random-effects (co)variance matrices must be checked.
set.negeigen	positive value. Value to which negative eigenvalues are to be set in estimators where such method is used to force semi-positive definiteness of the estimated between-study (co)variance matrix.

Value

A list with components named as the arguments.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

References

Gasparrini, A., Armstrong, B., Kenward, M. G. (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*, 31(29), 3821-3839.

See Also

[dosresmeta](#), [dosresmeta-package](#), [dosresmeta.control](#)

Examples

```
## Loading data
data("alcohol_cvd")

## print the iterations (see ?optim) and change the default for starting values
dosresmeta(formula = logrr ~ dose, type = type, id = id, se = se,
            cases = cases, n = n, data = alcohol_cvd, proc = "1stage",
            control = list(showiter = TRUE, igls.inititer = 20))
```

dosresmeta.fixed *Fixed-Effects Estimator for dosresmeta Models*

Description

This function implements a generalized least square estimator for fixed-effects dose-response meta-analysis. It is meant to be used internally and not directly run by the users.

Usage

```
dosresmeta.fixed(Xlist, Zlist, ylist, Slist, nalist, q, nall, control, ...)
```

Arguments

Xlist	a m-dimensional list of study-specific design matrices for the fixed-effects part of the model.
Zlist	a m-dimensional list of study-specific design matrices for the random-effects part of the model.
ylist	a m-dimensional list of study-specific of vectors of estimated outcomes.
Slist	a m-dimensional list of within-study (co)variance matrices of estimated outcomes.
nalist	a m-dimensional list of k-dimensional study-specific logical vectors, identifying missing outcomes.
q, nall	numeric scalars: number of predictors, number of observations (excluding missing).
control	list of parameters for controlling the fitting process, usually internally set to default values by <code>dosresmeta.control</code> .
...	further arguments passed to or from other methods. Currently not used.

Details

The estimation involves only the p fixed-effects coefficients. The routine is based on a standard generalized least square (GLS) algorithm implemented in the internal function `glsfit`. The between-study (co)variance matrix is set to zero, so the marginal (co)variance matrix, composed only by elements of the within-study component, is assumed as completely known. Similarly to the likelihood-based estimators implemented in `dosresmeta.ml` and `dosresmeta.reml`, the computation involves Cholesky and QR decompositions for computational stability and efficiency.

Value

This function returns an intermediate list object, whose components are then processed by `dosresmeta.fit`. Other components are added later through `mixmeta` to finalize an object of class "dosresmeta".

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

References

Gasparini, A., Armstrong, B., Kenward, M. G. (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*, 31(29), 3821-3839.

See Also

[dosresmeta](#), [dosresmeta-package](#), [dosresmeta.ml](#)

Examples

```
data("alcohol_crc")

## Fixed-effect dose-response model assuming linearity
dosresmeta(formula = logrr ~ dose, type = type, id = id, se = se,
            cases = cases, n = peryears, data = alcohol_crc, method = "fixed")
```

dosresmeta.ml

ML and REML Estimators for dosresmeta Models

Description

These functions implement maximum likelihood (ML) and restricted maximum likelihood (REML) estimators for random-effects dose-response meta-analysis. They are meant to be used internally and not directly run by the users.

Usage

```
dosresmeta.ml(Xlist, Zlist, ylist, Slist, nalist, q, nall, control, ...)
```

```
dosresmeta.reml(Xlist, Zlist, ylist, Slist, nalist, q, nall, control, ...)
```

Arguments

Xlist	a m-dimensional list of study-specific design matrices for the fixed-effects part of the model.
Zlist	a m-dimensional list of study-specific design matrices for the random-effects part of the model.
ylist	a m-dimensional list of study-specific of vectors of estimated outcomes.
Slist	a m-dimensional list of within-study (co)variance matrices of estimated outcomes.
nalist	a m-dimensional list of k-dimensional study-specific logical vectors, identifying missing outcomes.
q, nall	numeric scalars: number of predictors, number of observations (excluding missing).
control	list of parameters for controlling the fitting process, usually internally set to default values by <code>dosresmeta.control</code> .
...	further arguments passed to or from other methods. Currently not used.

dosresmetaObject *dosresmeta Object*

Description

An object returned by `dosresmeta` function, inheriting from class "dosresmeta", and representing a fitted dose-response (meta-analytical) model.

Value

Objects of class "dosresmeta" are lists with defined components. Dimensions of such components differs according to the chosen procedure. For the one-stage analysis the dimensions refer to a one dimensional outcome, p predictors and m studies used for fitting the model. For the two-stage analysis the dimensions refer to p outcome parameters, no predictor (only the intercept) and m studies. The following components needs to be included in a legitimate dosresmeta object:

<code>coefficients</code>	a p -dimensional vector of the fixed-effects coefficients.
<code>vcov</code>	estimated $p \times p$ (co)variance matrix of the fixed-effects coefficients.
<code>Psi</code>	for random-effects models, the estimated $p \times p$ between-study (co)variance matrix.
<code>residuals</code>	a vector of residuals, that is observed minus fitted values.
<code>fitted.values</code>	a vector of of fitted mean values.
<code>df.residual</code>	the residual degrees of freedom.
<code>rank</code>	the numeric rank of the fitted model.
<code>logLik</code>	the (restricted) log-likelihood of the fitted model.
<code>converged, niter</code>	for models with iterative estimation methods, logical scalar indicating if the algorithm eventually converged.
<code>control</code>	a list with the values of the control arguments used, as returned by <code>dosresmeta.control</code> .
<code>method</code>	the estimation method.
<code>dim</code>	list with the following scalar components: m (number of studies included in estimation), k (number of outcome parameters).
<code>df</code>	list with the following scalar components: <code>nall</code> (number of observations used for estimation, excluding missing values).
<code>lab</code>	list with the following label vectors: p for the p predictors (including intercept).
<code>model</code>	the model frame used for fitting.
<code>call</code>	the function call.
<code>formula</code>	the model supplied.
<code>terms</code>	the <code>terms</code> object representing the fitted model.
<code>proc</code>	the estimation procedure.
<code>center</code>	if the desing matrix had been centered.
<code>covariance</code>	how the (co)variance had been approximated.
<code>Slist</code>	list of approximated (co)variance matrices.
<code>id</code>	identification vector of the studies.
<code>v</code>	variances of the outcome values

Methods

A number of methods functions are available for `dosresmeta` objects, most of them common to other regression functions. Specifically-written method functions are defined for `predict` (standard predictions). The `qtest` method performs the Cochran Q test for heterogeneity only for a two-stage

analysis. Other methods have been produced for summary, logLik, coef, and vcov. Printing functions for the objects of classes defined above are also provided. All the methods above are visible (exported from the namespace) and documented. In additions, several default method functions for regression are also applicable to objects of class "mixmeta", such as fitted, residuals, AIC, BIC and update, among others.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

See Also

[dosresmeta](#), [dosresmeta-package](#), [mixmetaObject](#)

fish_ra	<i>Six studies on the relation between fish consumption and rheumatoid arthritis risk</i>
---------	---

Description

The dataset reports the summarized dose-response results from six studies on the relation between fish consumption and rheumatoid arthritis risk

Format

A data frame with 22 observations on the following 12 variables:

id	id of the studies included in the analysis.
author	names of the first author.
year	year of publication.
type	code for study design.
cases	number of cases for each exposure level.
n	total number of subjects (type = "ci") or person-years (type = "ir") for each exposure level.
dose	assigned dose levels.
rr	adjusted risk estimates.
lrr	lower bound for the confidence limits of the adjusted risk estimates.
urr	upper bound for the confidence limits of the adjusted risk estimates.
logrr	natural logarithm of the adjusted odds ratio.
se	standard error for the logarithm of the adjusted odds ratio

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Di Giuseppe D, Crippa A, Orsini N, Wolk A. Fish consumption and risk of rheumatoid arthritis: a dose-response meta-analysis. *Arthritis research & therapy*. 2014 Sep 30;16(5):446.

fpgrid

Grid with combinations of p for two-order fractional polynomials

Description

Computes the different combinations of p usefull for evaluating two-order fractional polynomials.

Usage

```
fpgrid(p = c(-2, -1, -0.5, 0, 0.5, 1, 2, 3))
```

Arguments

p a numeric vector with the coefficient to be combined.

Value

A data.frame with the different combinations of p.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

References

Royston, Patrick, and Douglas G. Altman. "Regression using fractional polynomials of continuous covariates: parsimonious parametric modelling." *Applied Statistics* (1994): 429-467.

Examples

```
grd <- fpgrid()
head(grd)
```

fracpol*Fractional Polynomials*

Description

Two-order fractional polynomials transformation for continuous covariates.

Usage

```
fracpol(x, p = c(1, 1), shift, scale, scaling = TRUE)
```


Arguments

x	a numeric vector.
p	a vector of length 2 with the powers of x to be included.
shift	optional scalar representing the shift, if scaling = TRUE. If not specified it is set internally equal to 0.
scale	optional scalar representing the scale, if scaling = TRUE. If not specified it is set internally equal to 1.
scaling	a logical indicating if the measurements are scaled prior to model fitting.

Details

The fracpol is based on the FP function in the mboost package. See help(FP) for more details.

Value

A matrix including the transformations corresponding to the input values.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

References

Royston, Patrick, and Douglas G. Altman. "Regression using fractional polynomials of continuous covariates: parsimonious parametric modelling." *Applied Statistics* (1994): 429-467.

See Also

mboost, rcs.eval

Examples

```
## Load data and run the model
data("alcohol_cvd")

with(alcohol_cvd, fracpol(dose, p = c(.5, .5)))

model <- dosresmeta(formula = logrr ~ fracpol(dose, p = c(.5, .5)), type = type, id = id,
                    se = se, cases = cases, n = n, data = alcohol_cvd)

## Test for significance of the overall dose-response association
waldtest(b = coef(model), Sigma = vcov(model), Terms = 1:nrow(vcov(model)))
```

gof	<i>Computes statistics to evaluate the goodness-of-fit from dosresmeta Objects</i>
-----	--

Description

This function computes statistics to evaluate the goodness-of-fit for dose-response meta-analysis. It implements the deviance test, the coefficient of determination, and a dataframe useful for a decorrelated residuals-versus-exposure plot. See reference for more details

Usage

```
gof(object, fixed = TRUE)

## S3 method for class 'gof.dosresmeta'
print(x, digits = 3, ...)
```

Arguments

object	an object of class dosresmeta produced by dosresmeta .
fixed	logical for selecting fixed model. By default equal to TRUE.
x	an object of class gof.dosresmeta produced by gof.
digits	an integer specifying the number of digits to which printed results must be rounded.
...	further arguments passed to or from other methods.

Value

A list of class gof.dosresmeta containing the following

tdata	a dataframe with the decorrelated variables (y^* , X^* , e^*).
R2	Coefficient of determination R^2 .
deviance	Deviance test.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

References

Discacciati A, Crippa A, Orsini N. Goodness of fit tools for dose-response meta-analysis of binary outcomes. Research synthesis methods. 2015 Jan 1.

Examples

```
## Loading the data
data("milk_ov")

## Linear dose-response model
lin <- dosresmeta(formula = logrr ~ dose, type = type, id = id,
                 se = se, cases = case, n = n, data = milk_ov)

## Display goodness-of-fit statistics
gof(lin)

## Meta-regression model
lin_reg <- dosresmeta(formula = logrr ~ dose, type = type, id = id,
                    se = se, cases = case, n = n, data = milk_ov,
                    mod = ~ type)

## Display goodness-of-fit statistics for meta-regression model
gof(lin_reg)
```

grl	<i>Approximating effective-counts as proposed by Greenland & Longnecker</i>
-----	---

Description

Reconstructs the set of pseudo-numbers (or 'effective' numbers) of cases and non-cases consistent with the input data (log relative risks). The method was first proposed in 1992 by Greenland and Longnecker.

Usage

```
grl(y, v, cases, n, type, data, tol = 1e-05)
```

Arguments

y	a vector, defining the (reported) log relative risks.
v	a vector, defining the variances of the reported log relative risks.
cases	a vector, defining the number of cases for each exposure level.
n	a vector, defining the total number of subjects for each exposure level. For incidence-rate data n indicates the amount of person-time within each exposure level.
type	a vector (or a character string), specifying the design of the study. Options are <code>cc</code> , <code>ir</code> , and <code>ci</code> , for case-control, incidence-rate, and cumulative incidence data, respectively.
data	an optional data frame (or object coercible by <code>as.data.frame</code> to a data frame) containing the variables in the previous arguments.
tol	define the tolerance.

Details

The function reconstructs the effective counts corresponding to the multivariable adjusted log relative risks as well as their standard errors. A unique solution is guaranteed by keeping the margins of the table of pseudo-counts equal to the margins of the crude or unadjusted data (Greenland and Longnecker 1992). See the referenced article for a complete description of the algorithm implementation.

Value

The results are returned structured in a matrix

- A approximated number of effective cases.
- N approximated total number of effective subjects.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

References

Greenland, S., Longnecker, M. P. (1992). Methods for trend estimation from summarized dose-response data, with applications to meta-analysis. *American journal of epidemiology*, 135(11), 1301-1309.

Orsini, N., Li, R., Wolk, A., Khudyakov, P., Spiegelman, D. (2012). Meta-analysis for linear and nonlinear dose-response relations: examples, an evaluation of approximations, and software. *American journal of epidemiology*, 175(1), 66-73.

See Also

[covar.logrr](#), [hamling](#)

Examples

```
## Loading data
data("alcohol_cvd")

## Obtaining pseudo-counts for the first study (id = 1)
grl(y = logrr, v = I(se^2), cases = cases, n = n, type = type,
    data = subset(alcohol_cvd, id == 1))

## Obtaining pseudo-counts for all study
by(alcohol_cvd, alcohol_cvd$id, function(x)
    grl(y = logrr, v = I(se^2), cases = cases, n = n, type = type, data = x))

## Restructuring the previous results in a matrix
do.call("rbind", by(alcohol_cvd, alcohol_cvd$id, function(x)
    grl(y = logrr, v = I(se^2), cases = cases, n = n, type = type, data = x)))
```

`hamling`*Approximating effective-counts as proposed by Hamling*

Description

Reconstructs the set of pseudo-numbers (or "effective" numbers) of cases and non-cases consistent with the input data (log relative risks). The method was first proposed in 2008 by Hamling.

Usage

```
hamling(y, v, cases, n, type, data)
```

Arguments

<code>y</code>	a vector, defining the (reported) log relative risks.
<code>v</code>	a vector, defining the variances of the reported log relative risks.
<code>cases</code>	a vector, defining the number of cases for each exposure level.
<code>n</code>	a vector, defining the total number of subjects for each exposure level. For incidence-rate data <code>n</code> indicates the amount of person-time within each exposure level.
<code>type</code>	a vector (or a character string), specifying the design of the study. Options are <code>cc</code> , <code>ir</code> , and <code>ci</code> , for case-control, incidence-rate, and cumulative incidence data, respectively.
<code>data</code>	an optional data frame (or object coercible by <code>as.data.frame</code> to a data frame) containing the variables in the previous arguments.

Details

The function reconstructs the effective counts corresponding to the multivariable adjusted log relative risks as well as their standard errors. A unique solution is guaranteed by keeping the ratio non-cases to cases and the fraction of unexposed subjects equal to the unadjusted data (Hamling). See the referenced article for a complete description of the algorithm implementation.

Value

A list containing the following

<code>y</code>	mean or standardized mean differences for each treatment level, included the referent one (0 by calculation).
<code>v</code>	variances corresponding to the mean or standardized mean differences for each treatment level, included the referent one (0 by calculation).
<code>S</code>	co(variance) matrix for the non-referent mean or standardized mean differences.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

References

Hamling, J., Lee, P., Weitkunat, R., Ambuhl, M. (2008). Facilitating meta-analyses by deriving relative effect and precision estimates for alternative comparisons from a set of estimates presented by exposure level or disease category. *Statistics in medicine*, 27(7), 954-970.

Orsini, N., Li, R., Wolk, A., Khudyakov, P., Spiegelman, D. (2012). Meta-analysis for linear and nonlinear dose-response relations: examples, an evaluation of approximations, and software. *American journal of epidemiology*, 175(1), 66-73.

See Also

[covar.logrr, gr1](#)

Examples

```
## Loading data
data("alcohol_cvd")

## Obtaining pseudo-counts for the first study (id = 1)
hamling(y = logrr, v = I(se^2), cases = cases, n = n, type = type,
data = subset(alcohol_cvd, id == 1))

## Obtaining pseudo-counts for all study
by(alcohol_cvd, alcohol_cvd$id, function(x)
hamling(y = logrr, v = I(se^2), cases = cases, n = n, type = type, data = x))

## Restructuring the previous results in a matrix
do.call("rbind", by(alcohol_cvd, alcohol_cvd$id, function(x)
  hamling(y = logrr, v = I(se^2), cases = cases, n = n, type = type,
    data = x)))
```

ir_ex

Incidence-rate data on fiber intake and coronary heart disease risk

Description

The dataset reports the summarized dose-response results from incidence-rate data investigating the association between the long-term intake of dietary fiber and risk of coronary heart disease among women, first presented by Wolk et al. (1999)

Format

A data frame with 5 observations on the following 8 variables:

dose	assigned dose levels.
cases	number of cases for each exposure level.
n	total number of subjects for each exposure level.
adjrr	adjusted incidence rate ratios for each exposure level.

lb lower bound for the confidence limits of the adjusted incidence rate ratios.
 ub upper bound for the confidence limits of the adjusted incidence rate ratios.
 logrr natural logarithm of adjusted incidence rate ratios.
 se standard error for the logarithm of the adjusted incidence rate ratios.

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Wolk, A., J. E. Manson, M. J. Stampfer, G. A. Colditz, F. Hu, F. E. Speizer, C. H. Hennekens, and W. C. Willett. 1999. Long-term intake of dietary fiber and decreased risk of coronary heart disease among women. *Journal of the American Medical Association* 281: 1998-2004.

Greenland, S., Longnecker, M. P. (1992). Methods for trend estimation from summarized dose-response data, with applications to meta-analysis. *American journal of epidemiology*, 135(11), 1301-1309.

logLik.dosresmeta *Extract Log-Likelihood from dosresmeta Objects*

Description

This method function returns the log-likelihood for fitted dose-response models represented in objects of class "dosresmeta".

Usage

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

Arguments

object an object of class "dosresmeta"
 ... further arguments passed to or from other methods.

Value

A numeric scalar of class "logLik".

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

See Also

[dosresmeta](#), [dosresmeta-package](#), [logLik](#)

Examples

```

data("alcohol_crc")

## Dose-response model assuming linearity
lin <- dosresmeta(formula = logrr ~ dose, type = type, id = id, se = se,
                  cases = cases, n = peryears, data = alcohol_crc, proc = "1stage")

## Log-likelihood
ll <- logLik(lin)
ll
attributes(ll)

## AIC and BIC
AIC(ll)
BIC(ll)

```

milk_mort

*Eleven prospective studies on the relation between milk consumption
and all-cause mortality*

Description

The dataset reports the summarized dose-response results from eleven prospective studies on the relation between milk consumption and all-cause mortality.

Format

A data frame with 50 observations on the following 12 variables:

id	id of the studies included in the analysis.
author	names of the first author.
year	year of publication.
type	code for study design.
dose	assigned dose levels.
cases	number of cases for each exposure level.
n	total number of subjects (type = "ci") or person-years (type = "ir") for each exposure level.
rr	adjusted risk estimates.
lb	lower bound for the confidence limits of the adjusted risk estimates.
ub	upper bound for the confidence limits of the adjusted risk estimates.
logrr	natural logarithm of the adjusted risk estimates.
se	standard error for the logarithm of the adjusted risk estimates.

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Larsson SC, Crippa A, Orsini N, Wolk A, Michaelsson K. Milk consumption and mortality from all causes, cardiovascular disease, and cancer: a systematic review and meta-analysis. *Nutrients*. 2015 Sep 11;7(9):7749-63.

milk_ov	<i>Nine studies on the relation between milk consumption and ovarian cancer</i>
---------	---

Description

The dataset reports the summarized dose-response results from nine studies on the relation between milk consumption and ovarian cancer.

Format

A data frame with 37 observations on the following 12 variables:

id	id of the studies included in the analysis.
author	names of the first author.
year	year of publication.
type	code for study design.
dose	assigned dose levels.
case	number of cases for each exposure level.
n	total number of subjects (type = "ir" or "cc") or person-years (type = "ir") for each exposure level.
rr	adjusted risk estimates.
lb	lower bound for the confidence limits of the adjusted risk estimates.
ub	upper bound for the confidence limits of the adjusted risk estimates.
logrr	natural logarithm of the adjusted risk estimates.
se	standard error for the logarithm of the adjusted risk estimates.

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Larsson, S. C., N. Orsini, and A. Wolk. 2005. Milk, milk products and lactose intake and ovarian cancer risk: A meta-analysis of epidemiological studies. *International Journal of Cancer* 118: 431-441.

Greenland, S., Longnecker, M. P. (1992). Methods for trend estimation from summarized dose-response data, with applications to meta-analysis. *American journal of epidemiology*, 135(11), 1301-1309.

Description

These functions compute the value of the log-likelihood for random-effects dose-response meta-analysis, in terms of model parameters. They are meant to be used internally and not directly run by the users.

Usage

```
remlprof.fn(par, Xlist, Zlist, ylist, Slist, nalist, q, nall, ctrl)
```

```
remlprof.gr(par, Xlist, ylist, Slist, nalist, p, nall, ctrl)
```

```
mlprof.fn(par, Xlist, Zlist, ylist, Slist, nalist, q, nall, ctrl)
```

```
mlprof.gr(par, Xlist, ylist, Slist, nalist, p, nall, ctrl)
```

```
iter.igls(Psi, Xlist, Zlist, ylist, Slist, nalist, q)
```

Arguments

par	a vector representing the random-effects parameters defining the between-study (co)variance matrix.
Xlist	a m-dimensional list of study-specific design matrices for the fixed-effects part of the model.
Zlist	a m-dimensional list of study-specific design matrices for the random-effects part of the model.
ylist	a m-dimensional list of study-specific of vectors of estimated outcomes.
Slist	a m-dimensional list of within-study (co)variance matrices of estimated outcomes.
nalist	a m-dimensional list of k-dimensional study-specific logical vectors, identifying missing outcomes.
ctrl	list of parameters for controlling the fitting process, usually internally set to default values by <code>dosresmeta.control</code> .
p, q, nall	numeric scalars: number of predictors, number of observations (excluding missing).
Psi	a $p \times p$ matrix representing the current estimate of the between-study (co)variance matrix.

Details

These functions are called internally by the fitting functions `dosresmeta.ml` and `dosresmeta.reml` to perform iterative optimization algorithms for estimating random effects meta-analytical models.

The maximization of the (restricted) likelihood starts with few runs of an iterative generalized least square algorithm implemented in `iter.igls`. This can be regarded as a fast and stable way to get starting values close to the maximum for the Quasi-Newton iterative algorithm, implemented in `optim`. Alternatively, starting values can be provided by the user in the control list (see `mixmeta.control`).

These functions actually specify the profiled version of the (restricted) likelihood, expressed only in terms of random-effects parameters, while the estimate of the fixed-effects coefficients is provided at each iteration by the internal function `glsfit`, based on the current value of the between-study (co)variance matrix. At convergence, the value of this profiled version is identical to the full (restricted) likelihood. This approach is computationally efficient, as it reduces the number of parameters in the optimization routine.

The parameterization of the between-study (co)variance matrix ensures the positive-definiteness of the estimated matrix. A Cholesky decomposition is then performed on the marginal (co)variance matrix in order to re-express the problem as standard least square equations, an approach which speeds up the computation of matrix inverses and determinants. These equations are finally solved through a QR decomposition, which guarantees stability.

Value

`mlprof.fn` and `remlprof.fn` return the value of the (restricted) log-likelihood for a given set of parameters in `par`. `iter.igls` returns an updated estimate of Ψ given its initial value or the value at the previous iteration.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

See Also

`dosresmeta`, `mixmeta.fit`, `dosresmeta.control`, `mlprof.fn`

oc_breast

Twenty-two case-control studies on the relation between oral contraceptives use and breast cancer

Description

The dataset reports the summarized dose-response results from twenty-two case-control studies on the relation between oral contraceptives use and breast cancer

Format

A data frame with 113 observations on the following 14 variables:

id	id of the studies included in the analysis.
author	names of the first author.
year	year of publication.
type	code for study design.
duration	assigned dose levels.
cases	number of cases for each exposure level.
n	total number of subjects (type = "ir" or "cc") or person-years (type = "ir") for each exposure level.
or	adjusted odds ratios.
lb	lower bound for the confidence limits of the adjusted odds ratios.
ub	upper bound for the confidence limits of the adjusted odds ratios.
logor	natural logarithm of the adjusted odds ratios.
se	standard error for the logarithm of the adjusted odds ratios.
menopause	indicator variable for a study that included postmenopausal women (1 = yes).
period	final year of case accrual (surrogate for the changing formulations of oral contraceptives over time).

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Berlin JA, Longnecker MP, Greenland S. Meta-analysis of epidemiologic dose-response data. *Epidemiology*. 1993 May 1:218-28.

predict.dosresmeta *Predicted Values from dosresmeta Models*

Description

This method function computes predictions from fitted dose-response models represented in objects of class "dosresmeta", optionally for a new set of exposure levels. Predictions are optionally accompanied by confidence intervals and/or standard errors for the predictions.

Usage

```
## S3 method for class 'dosresmeta'
predict(
  object,
  newdata,
  xref,
  expo = FALSE,
  xref_vec,
  ci.incl = TRUE,
```

```

    se.incl = FALSE,
    xref_pos = 1,
    delta,
    order = FALSE,
    ci.level = 0.95,
    safe_sort2stage = TRUE,
    ...
)

```

Arguments

object	an object of class dosreseta.
newdata	an optional data frame or matrix in which to look for variables values with which to predict from dose-response models.
xref	an optional scalar to indicate which levels should serve as referent for the predicted relative risks. See details.
expo	logical switch indicating if the prediction should be on the exponential scale.
xref_vec	an optional numeric to indicate the referent (vector) for the predicted relative risks. See details.
ci.incl	logical switch indicating if confidence intervals need to be included.
se.incl	logical switch indicating if standard errors need to be included.
xref_pos	an optional scalar to indicate the position of the referent for the predicted relative risks. See details.
delta	an optional scalar to specify to predict the linear trend related to that increase.
order	logical to indicate if the predictions need to be sorted by exposure levels.
ci.level	a numerical value between 0 and 1, specifying the confidence level for the computation of confidence intervals.
safe_sort2stage	logical value to ensure the correct order of coefficients for prediction. Only needed for 2stage.
...	further arguments passed to or from other methods.

Details

The method function `predict` produces predicted values from `dosresmeta` objects. When more than one study is included in the analysis, estimated predictions are only based on the fixed part of the model.

If `newdata` is omitted, the predictions are based on the data used for the fit. If `xref` is provided, it must be equal to one of the modeled values. If not provided, the minimum modeled referent value will be used as referent for the predicted relative risks

If `newdata` is specified, it should include all the variables used to model the dose-response relation. Again, if specified, `xref` must be equal to one of the value in the `newdata`. If omitted, the minimum value for the `newdata` will be used as referent.

Only for the linear trend it is possible to specify the predicted increase of risk corresponding to an increase equal to `delta` argument.

By default (`order = TRUE`), the predictions are sorted by exposure levels to facilitate understanding and possible graphical presentation of the results.

Value

The results are returned structured in a data frame.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

See Also

[dosresmeta](#), [predict](#)

Examples

```
## Load data and run the linear and quadratic models
data("alcohol_cvd")
lin <- dosresmeta(formula = logrr ~ dose, type = type, id = id,
                 se = se, cases = cases, n = n, data = alcohol_cvd)
quadr <- dosresmeta(formula = logrr ~ dose + I(dose^2), type = type, id = id,
                  se = se, cases = cases, n = n, data = alcohol_cvd)

## Predicted linear trend (on RR scale)
predict(lin, delta = 12, expo = TRUE)

## Predicted modeled data from quadratic model (on RR scale)
predict(quadr, expo = TRUE)

## Plot predicted dose-response relation
with(predict(quadr, order = TRUE, expo = TRUE), {
  plot(dose, pred, log = "y", type = "l",
       xlim = c(0, 45), ylim = c(.4, 2))
  lines(dose, ci.lb, lty = 2)
  lines(dose, ci.ub, lty = 2)
  rug(dose, quiet = TRUE)
})

## Prediction for new values from quadratic model (on RR scale)
newdata <- data.frame(dose = seq(0, 50, 5))
predict(quadr, newdata, expo = TRUE)

## Loading data
data("ari")

mod <- dosresmeta(y ~ dose + I(dose^2), id = id, sd = sd,
                 n = n, data = ari, covariance = "smd")

## Smoothed plot
newdata <- data.frame(dose = seq(0, 30, 1))
with(predict(mod, newdata), {
```

```

plot(dose, pred, type = "l", ylim = c(0, .6))
lines(dose, ci.lb, lty = 2)
lines(dose, ci.ub, lty = 2)
rug(dose, quiet = TRUE)
})

```

```
print.dosresmeta      Summarizing dosresmeta Models
```

Description

Print and summary method functions for dose-response models represented in objects of class "dosresmeta".

Usage

```

## S3 method for class 'dosresmeta'
print(x, digits = 4, ...)

## S3 method for class 'dosresmeta'
summary(object, ci.level = 0.95, ...)

## S3 method for class 'summary.dosresmeta'
print(
  x,
  digits = max(3, getOption("digits") - 3),
  signif.stars = getOption("show.signif.stars"),
  ...
)

```

Arguments

x	an object of class dosresmeta or summary.dosresmeta produced by dosresmeta or summary.dosresmeta , respectively.
digits	an integer specifying the number of digits to which printed results must be rounded.
...	further arguments passed to or from other methods.
object	an object of class dosresmeta produced by dosresmeta .
ci.level	the confidence level used for defining the confidence intervals for the estimates of the (fixed-effects) coefficients.
signif.stars	logical. If TRUE, 'significance stars' are printed for each coefficient.

Details

the `print` method for class `dosresmeta` only returns basic information of the fitted model, namely the call, estimated (fixed-effects) coefficients, dimensions, and fit statistics (log-likelihood, AIC, BIC).

The `summary` method function computes additional statistics and tests, and produces a list object of class `summary.dosresmeta`. The `print` method function for this class, depending on the number of studies included in the analysis, shows additional information, such as tables reporting the estimates for the fixed and random-effects parts of the model, Chi-square test for model significance, Cochran Q test for heterogeneity and I-square.

Value

The `summary` method function for `dosresmeta` objects produces a list of class `"summary.dosresmeta"`. The components of the lists are some of those stored in the related `dosresmeta` object, plus the following:

AIC	the value of the Akaike information criterion for the fitted <code>dosresmeta</code> model, obtained through a call to AIC .
BIC	the value of the Bayesian information criterion for the fitted <code>dosresmeta</code> model, obtained through a call to BIC .
corFixed	the $p \times p$ correlation matrix of the fixed-effects coefficients, obtained from the (co)variance matrix vcov
corRandom	the $p \times p$ correlation matrix of the random effects, obtained from the between-study (co)variance matrix Ψ
qstat	results from the Cochran Q test for heterogeneity.
ci.level	the confidence level used for defining the confidence intervals for the estimates of the fixed-effects coefficients.
chisq	overall test similar to anova.

As usual, the `print` method functions for classes `"dosresmeta"` and `"summary.dosresmeta"` do not return any value.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

See Also

[dosresmeta](#), [summary](#)

Examples

```
## Load data and run the model
data("alcohol_cvd")
model <- dosresmeta(formula = logrr ~ dose + I(dose^2), type = type, id = id,
                    se = se, cases = cases, n = n, data = alcohol_cvd)

## Default print
model
## Specify digits
print(model, digit = 2)
## summary with 90th confidence intervals
summary(model, ci.level = .8)
```

process_bc	<i>Ten studies on the relation between processed meat and bladder cancer</i>
------------	--

Description

The dataset reports the summarized dose-response results from ten studies on the relation between processed meat consumption and bladder cancer.

Format

A data frame with 73 observations on the following 15 variables:

id	id of the studies included in the analysis.
author	names of the first author.
year	year of publication.
type	code for study design.
dose0	original assigned dose levels, with unit of measurement defined in the "unit" column.
dose	assigned dose levels (converted (if needed) in gm/day).
cases	number of cases for each exposure level.
n	total number of subjects (type = "ci" or "cc") or person-years (type = "ir") for each exposure level.
rr	adjusted risk estimates.
lb	lower bound for the confidence limits of the adjusted risk estimates.
ub	upper bound for the confidence limits of the adjusted risk estimates.
logrr	natural logarithm of the adjusted risk estimates.
se	standard error for the logarithm of the adjusted risk estimates.
area	geographical area of the published study.
unit	unit of measurement for red meat consumption (for dose0).

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Crippa A, Larsson SC, Discacciati A, Wolk A, Orsini N. Red and processed meat consumption and risk of bladder cancer: a dose-response meta-analysis of epidemiological studies. *European journal of nutrition*. 2016 Dec 22:1-3.

qttest.dosresmeta *Cochran Q Test of Heterogeneity for dosresmeta Models*

Description

This method function performs a Cochran Q test of (residual) heterogeneity on fitted dose-response meta-analytical models represented in objects of class "doseremeta". It is implemented only for a two-stage approach and will return NULL otherwise.

Usage

```
## S3 method for class 'dosresmeta'
qttest(object, ...)

## S3 method for class 'qttest.dosresmeta'
print(x, digits = 3, ...)
```

Arguments

object	objects of classe "dosresmeta".
...	further arguments passed to or from other methods.
x	an object of class "qttest.dosresmeta".
digits	an integer specifying the number of digits to which printed results must be rounded.

Details

In (multivariate) dose-response meta-analytical models, the test assesses the null hypothesis that the variability in the (multivariate) distribution of the outcomes is explained only in terms of estimation error in each study, measured by the within-study (co)variance matrices stored in the component `Slist` of `dosresmeta` objects. This is equal to test the hypothesis that the between-study (co)variance matrix is a zero matrix, and there is no random deviation in study-specific estimates.

red_bc *Twelve studies on the relation between red meat and bladder cancer*

Description

The dataset reports the summarized dose-response results from twelve studies on the relation between red meat consumption and bladder cancer.

Format

A data frame with 74 observations on the following 15 variables:

id	id of the studies included in the analysis.
author	names of the first author.
year	year of publication.
type	code for study design.
dose0	original assigned dose levels, with unit of measurement defined in the "unit" column.
dose	assigned dose levels (converted (if needed) in gm/day).
cases	number of cases for each exposure level.
n	total number of subjects (type = "ci" or "cc") or person-years (type = "ir") for each exposure level.
rr	adjusted risk estimates.
lb	lower bound for the confidence limits of the adjusted risk estimates.
ub	upper bound for the confidence limits of the adjusted risk estimates.
logrr	natural logarithm of the adjusted risk estimates.
se	standard error for the logarithm of the adjusted risk estimates.
area	geographical area of the published study.
unit	unit of measurement for red meat consumption (for dose0).

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Crippa A, Larsson SC, Discacciati A, Wolk A, Orsini N. Red and processed meat consumption and risk of bladder cancer: a dose-response meta-analysis of epidemiological studies. *European journal of nutrition*. 2016 Dec 22:1-3.

 sim_os

Simulated data for one-stage dose-response meta-analysis

Description

The dataset contains simulated data from 9 case-control studies.

Format

A data frame with 27 observations on the following 11 variables:

xcati	category limits for the continuous exposure.
id	id of the studies.
type	code for study design.
dose	assigned dose levels.
cases	number of cases for each exposure level.

n	total number of subjects for each exposure level.
rr	adjusted risk estimates for each exposure level.
lrr	lower bound for the confidence limits of the adjusted risk estimates.
urr	upper bound for the confidence limits of the adjusted risk estimates.
logrr	natural logarithm of the adjusted risk estimates.
se	standard error for the logarithm of the adjusted risk estimates.

Author(s)

Alessio Crippa, <<alessio.crippa@ki.se>>

References

Larsson, S. C., Orsini, N. (2011). Coffee consumption and risk of stroke: a dose-response meta-analysis of prospective studies. *American journal of epidemiology*, 174(9), 993-1001.

vpc

Variance Partition Components for dosresmeta Objects

Description

Computes the Variance Partition Components for dose-response meta-analysis.

Usage

vpc(object)

Arguments

object an object of class dosresmeta produced by [dosresmeta](#).

Value

A vector containing the variance partition components for each non-referent observation.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

References

Goldstein H, Browne W, Rasbash J. Partitioning variation in multilevel models. *Understanding Statistics: Statistical Issues in Psychology, Education, and the Social Sciences*. 2002 Dec 2;1(4):223-31.

Examples

```
## loading data
data("sim_os")

## Quadratic (one-stage) dose-response model
quadr <- dosresmeta(logrr ~ dose + I(dose^2), id = id, se = se, type = type,
  cases = cases, n = n, data = sim_os, proc = "1stage")

## Plot of the estimated vpc
plot(sim_os$dose[sim_os$se!=0], vpc(quadr), xlab = "dose")
lines(lowess(sim_os$dose[sim_os$se!=0], vpc(quadr)))
```

waldtest

Wald Test for Model Coefficients

Description

Computes a Wald chi-squared test for 1 or more coefficients, given their variance-covariance matrix.

Usage

```
waldtest(Sigma, b, Terms = NULL, L = NULL, H0 = NULL)

## S3 method for class 'waldtest'
print(x, digits = 2, ...)
```

Arguments

Sigma	a var-cov matrix, usually extracted from one of the fitting functions.
b	a vector of coefficients with var-cov matrix Sigma. These coefficients are usually extracted from one of the fitting functions available in R.
Terms	an optional integer vector specifying which coefficients should be jointly tested, using a Wald chi-squared or F test. Its elements correspond to the columns or rows of the var-cov matrix given in Sigma. Default is NULL.
L	an optional matrix conformable to b, such as its product with b gives the linear combinations of the coefficients to be tested. Default is NULL.
H0	a numeric vector giving the null hypothesis for the test. It must be as long as Terms or must have the same number of columns as L. Default to 0 for all the coefficients to be tested.
x	Object of class "waldtest".
digits	number of decimal places for displaying test results. Default to 2.
...	further arguments passed to or from other methods.

Details

The `waldtest` and the method `print.waldtest` are taken from the `aod` package and simplified for ease of use.

Value

An object of class `waldtest`, printed with `print.waldtest`.

Author(s)

Alessio Crippa, <alessio.crippa@ki.se>

See Also

`aod`, [summary.dosresmeta](#)

Examples

```
## Load data and run the model
data("alcohol_cvd")
model <- dosresmeta(formula = logrr ~ dose + I(dose^2), type = type, id = id,
                   se = se, cases = cases, n = n, data = alcohol_cvd)

## Test for significance of the overall dose-response association
waldtest(b = coef(model), Sigma = vcov(model), Terms = 1:nrow(vcov(model)))
```

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