# <span id="page-0-0"></span>Package: mvmeta (via r-universe)

September 6, 2024

Type Package Title Multivariate and Univariate Meta-Analysis and Meta-Regression Version 1.0.3 Date 2019-12-10 Author Antonio Gasparrini Maintainer Antonio Gasparrini <antonio.gasparrini@lshtm.ac.uk> **Depends**  $R (= 2.13.0)$ Imports mixmeta, stats, graphics, grDevices, utils Suggests metafor, meta, rmeta, nlme, MASS, dlnm Description Collection of functions to perform fixed and random-effects multivariate and univariate meta-analysis and meta-regression. URL <http://www.ag-myresearch.com/package-mvmeta> License GPL  $(>= 2)$ LazyData yes Repository https://gasparrini.r-universe.dev RemoteUrl https://github.com/gasparrini/mvmeta

RemoteRef HEAD

RemoteSha 01c767ae87854873a4b762c2d7a270cba1ad0118

## **Contents**



<span id="page-1-0"></span>

<span id="page-1-1"></span>mvmeta-package *Multivariate and Univariate Meta-Analysis and Meta-Regression*

Description

The package mvmeta consists of a collection of functions to perform fixed and random-effects multivariate and univariate meta-analysis and meta-regression in R.

It is now superseded by the package mixmeta, which offers a unified mixed-effects framework to perform various meta-analytical models in R, including non-standard extensions such as multivariate, multilevel, longitudinal, and dose-response models. The package mvmeta will be still maintained, but its development is now discontinued. Improvements and further extensions will be implemented in mixmeta. Users are suggested to switch to the latter.

## Modelling framework

Multivariate meta-analytical models represent an extension of the standard univariate techniques, where estimates of a single effect size, here defined generally as outcome, are pooled across studies. In multivariate meta-analysis, estimates of multiple outcomes are combined while accounting for their correlation. Multivariate meta-regression also models such multivariate distribution in terms of study-level predictors. These statistical tools were originally proposed to model multiple endpoints in clinical trials. Applications and methodological developments are currently proposed also for network meta-analysis (indirect treatment comparison), for the meta-analysis of multi-parameter

## mvmeta-package 3

associations and for meta-analysis of diagnostic studies, among others. See references below for details.

Similarly to univariate methods, fixed-effects models do not assume heterogeneity among studies, and the estimates are conditional on the set of studies collected in the meta-analysis, based on a multivariate weighted average of study-level estimates. Random-effects meta-analysis, instead, allows a degree of heterogeneity among studies, assuming the (true but unobserved) study-specific outcomes as randomly sampled from a multivariate normal distribution of studies. Inference from these models may therefore be extended to other unmeasured studies assumed to belong to the same (usually hypothetical) population.

Estimation and interpretation exploit here the framework of linear mixed models. The fixed part of the model provides an estimate of fixed effects, which represent the population-averaged outcomes and, in the case of meta-regression, are defined by a set of coefficients associated to study-level predictors. The random part of the model describes the deviation from the population averages, estimating the components of a between-study (co)variance matrix. Assuming k outcomes  $y_i$  estimated in each of  $i = 1, \ldots, m$  studies, and related to p study-level predictors  $x_i$ , random-effects multivariate meta-regression models can be generally described with:

$$
\boldsymbol{y}_i \sim \mathrm{N}_k(\boldsymbol{X}_i \boldsymbol{\beta}, \boldsymbol{S}_i + \boldsymbol{\Psi})
$$

Here the outcomes  $y_i$  are assumed to be sampled from a multivariate normal distribution of order k. Their distribution is centred on  $X_i\beta$ , with  $X_i$  as a  $k \times kp$  design matrix and  $\beta$  the vector of fixed-effects coefficients. The marginal  $k \times k$  (co)variance matrix  $\Sigma_i = S_i + \Psi$  is given by the sum of within (assumed known) and between-study (co)variance matrices  $S_i$  and  $\Psi$ , respectively.

Other models are taken as special cases of that above. In multivariate meta-analysis,  $X_i$  becomes an identity matrix with  $p = 1$ , and  $\beta$  reduces to k intercepts, interpreted as the populationaveraged outcomes. For  $k = 1$ , the model reduces to the standard univariate meta-analysis or meta-regression. In fixed-effects meta-analytic models,  $\Psi$  is assumed not to exist, and the variability between studies is due exclusively to the within-study estimation error.

## Estimation methods

The aim is to estimate the coefficients  $\beta$  and, for random-effects models, the components of the between-study (co)variance matrix  $\Psi$ . If the same linear predictor of p terms  $x_i$  measured in each study is specified for all the outcomes (the only option available in the current version), the dimension of  $\beta$  is kp. The parameters for the random part depend on the chosen structure of the between-study (co)variance matrix  $\Psi$ , with  $k(k+1)/2$  parameters for an unstructured form, and a smaller set of parameters for structured (co)variances.

Different estimator are implemented in the package **mymeta**. The estimation options available in the current version are:

- Fixed-effects
- Maximum likelihood (ML)
- Restricted maximum likelihood (REML)
- Method of moments
- Variance components

<span id="page-3-0"></span>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. However, they are based on computationally intensive iterative procedures of optimization, and convergence can be slow for high-dimensional models (with a high number of outcomes). Estimators based on semiparametric alternatives such as the non-iterative method of moments or the iterative variance components approach are generally faster. Further details on estimation methods are given in the related help pages.

## Functions and data included in the package

The main function in the package is [mvmeta](#page-24-1), which performs the various models illustrated above. This function resembles standard regression functions in R, and specifies the model through a regression formula. The function returns a list object of class "mvmeta" (see [mvmetaObject](#page-44-1)).

The estimation is carried out internally through [mvmeta.fit](#page-24-2), a wrapper which prepares the data and calls specific estimation functions for fitting the models. Specifically, mymeta.fixed is applied for fixed-effects models, while estimators for random-effects models are implemented in the functions mymeta.ml and mymeta.reml for (restricted) maximum likelihood, mymeta.mm for the method of moments, and mymeta. vc for variance components. For likelihood-based methods, iterative [optimizations algorithms](#page-20-1) are used for maximizing the (restricted) likelihood, and specific [\(co\)variance structures](#page-42-1) for the between-study random effects are available. Fitting parameter options are set by [mvmeta.control](#page-29-1).

Method functions are available for objects of class "mvmeta" (see [mvmetaObject](#page-44-1) for a complete list). The method [summary](#page-58-1) produces a list of class "summary.mvmeta" for summarizing the fit of the model and providing additional results. The method function [predict](#page-52-1) computes predicted values, optionally for a set of new values of the predictors. [blup](#page-8-1) gives the (empirical) best linear unbiased predictions for the set of studies used for estimation. Other default or specific method functions for regression can be used on objects of class "mvmeta", such as [fitted](#page-0-0) and [residuals](#page-0-0), [logLik](#page-19-1), [AIC](#page-0-0) and [BIC](#page-0-0), among others.

Methods for [model.frame](#page-23-1) and [model.matrix](#page-23-2) are used to extract and construct the model frame and the design matrix of the regression meta-analytical model, respectively. Methods for [na.omit](#page-49-1) and [na.exclude](#page-49-2) help handle correctly missing values.

Simulations can be produced using the function [mvmetaSim](#page-47-1) and the method function [simulate](#page-47-2), which return one or multiple sets of simulated outcomes for a group of studies. The function [inputna](#page-17-1) and [inputcov](#page-15-1) are used internally to augment the missing data values and to input missing correlations, respectively.

The method function [qtest.mvmeta](#page-55-1) (producing an object with class of the same name) performs the (multivariate) Cochran Q test for (residual) heterogeneity, both on the overall multivariate distribution and on each single outcome. The generic method function is [qtest](#page-54-1).

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, mvmeta:::glsfit displays the code of the function glsfit. Also, some comments are added in the original source code.

The package includes the datasets [berkey98](#page-5-1), [fibrinogen](#page-11-1), [hsls](#page-12-1), [hyp](#page-14-1), [p53](#page-50-1) and [smoking](#page-57-1) as data frames, which are used in the examples.

## mvmeta-package 5

## Additional information

A list of changes included in the current and previous versions can be found by typing:

file.show(system.file("ChangeLog",package="mvmeta"))

General information on the development and applications of the **mymeta** package and on the modelling framework of multivariate meta-analysis, together with an updated version of the R scripts for running the examples in published papers, can be found at **[www.ag-myresearch.com](http://www.ag-myresearch.com)**.

## Warnings

This release of the package **mymeta** has been tested with different simulated and real datasets. The functions generally perform well under several scenarios, and comparisons with alternative software implementations show good agreement. However, bugs and bad performance under untested conditions may not be excluded. Please report any error or unexpected behaviour to the e-mail address below.

## Note

The functions included in the package **mymeta** may be applied also to perform standard univariate meta-analysis and meta-regression. However, alternative packages provides a more exhaustive and efficient set of functions. See for example the packages metafor, meta, rmeta and metaLik, among others.

Functions for fitting multivariate models are available in other R packages. The package metafor provides functions to perform fixed or random-effects multivariate models, the latter fitted through maximum or restricted maximum likelihood. A thorough comparison with the implementation offered in **mymeta** is not available yet. The package metaSEM (not in CRAN) offers functions to perfor fixed-effects and likelihood-based random-effects multivariate meta-analysis through structural equation modelling. The package **mytmeta** performs fixed-effects and random-effects multivariate meta-analysis using the same method of moments estimator adopted here, although without allowing for missing outcomes or meta-regression. The package **mada** contains functions for bivariate models used in diagnostic studies.

Use citation("mvmeta") to cite this package.

#### Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

White IR (2009). Multivariate random-effects meta-analysis. *Stata Journal*. 9(1):40–56.

Jackson D, Riley R, White IR (2011). Multivariate meta-analysis: Potential and promise. *Statistics in Medicine*. 30(20);2481–2498.

<span id="page-5-0"></span>White IR (2011). Multivariate random-effects meta-regression: updates to mvmeta. *Stata Journal*. 11(2):255–270.

van Houwelingen HC, Arends LR, et al. (2002). Advanced methods in meta-analysis: multivariate approach and meta-regression. *Statistics in Medicine*. 21(4):589–624.

Lu G, Ades AE (2004). Combination of direct and indirect evidence in mixed treatment comparisons. *Statistics in Medicine*. 23(20):3105–3124.

Nam IS, Mengersen K, et al. (2003). Multivariate meta-analysis. *Statistics in Medicine*. 22(14):2309– 2333.

Arends LR, Voko Z, Stijnen T (2003). Combining multiple outcome measures in a meta-analysis: an application. *Statistics in Medicine*. 22(8):1335–1353.

Ritz J, Demidenko E, Spiegelman G (2008). Multivariate meta-analysis for data consortia, individual patient meta-analysis, and pooling projects. *Journal of Statistical Planning and Inference*. 139(7):1919–1933.

Berkey, CS, Anderson JJ, Hoaglin DC (1996). Multiple-outcome meta-analysis of clinical trials. *Statistics in Medicine*. 15(5):537–547.

Berkey, CS, Hoaglin DC, et al. (1998). Meta-analysis of multiple outcomes by regression with random effects. *Statistics in Medicine*. 17(22):2537–2550.

Jackson D, White IR, Riley RD (2013). A matrix based method of moments for fitting the multivariate random effects model for meta-analysis and meta-regression. *Biometrical Journal*. 55(2):231- 245.

Chen H, Manning AK, Dupuis J (2012). A method of moments estimator for random effect multivariate meta-analysis. *Biometrics*. 68(4):1278-1284.

Cheung MWL, Chan W (2009). A two-stage approach to synthesizing covariance matrices in metaanalytic structural equation modeling. *Structural Equation Modeling*. 16(1):28–53.

Doebler P, Holling H, Bohning D (2012). A mixed model approach to meta-analysis of diagnostic studies with binary test outcome. *Psychological Methods*. 17(3):418–36.

<span id="page-5-1"></span>berkey98 *Five Published Trials on Periodontal Disease*

## Description

The dataset contains the results of 5 published trials comparing surgical and non-surgical treatments for medium-severity periodontal disease, one year after treatment. The 2 estimated outcomes are average improvements (surgical minus non-surgical, in mm) in probing depth (PD) and attachment level (AL).

#### Usage

berkey98

### berkey98 7

## Format

A data frame with 5 observations on the following 7 variables:

pubyear publication year of the trial.

npat number of patients included in the trial.

PD estimated improvement of surgical versus non-surgical treatments in probing depth (mm).

AL estimated improvement of surgical versus non-surgical treatments in attachment level (mm).

var\_PD variance of the estimated outcome for PD.

cov\_PD\_AL covariance of the estimated outcomes for PD and AL.

var\_AL variance of the estimated outcome for AL.

Row names specify the author of the paper reporting the results of each trial.

## Source

Berkey CS, Hoaglin DC, et al. (1998). Meta-analysis of multiple outcomes by regression with random effects. *Statistics in Medicine*. 17:2537–2550.

Berkey C. S., Antczak-Bouckoms A., et al. (1995). Multiple-outcomes meta-analysis of treatments for periodontal disease. *Journal of Dental Research*. 74(4):1030–1039.

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

## Examples

### REPRODUCE THE RESULTS IN BERKEY ET AL. (1998)

```
# INSPECT THE DATA
berkey98
# FIXED-EFFECTS
year <- berkey98$pubyear - 1983
model <- mvmeta(cbind(PD,AL)~year,S=berkey98[5:7],data=berkey98,method="fixed")
print(summary(model),digits=3)
# GLS MODEL (VARIANCE COMPONENTS)
model <- mvmeta(cbind(PD,AL)~year,S=berkey98[5:7],data=berkey98,method="vc",
  control=list(vc.adj=FALSE))
print(summary(model),digits=3)
round(model$Psi,3)
# ML MODEL
model <- mvmeta(cbind(PD,AL)~year,S=berkey98[5:7],data=berkey98,method="ml")
print(summary(model),digits=3)
round(model$Psi,3)
```
## <span id="page-7-0"></span>**Description**

This is a generic function for generating best linear unbiased predictions (BLUPs) from the results of various fitting functions for meta-analytical models. The function invokes particular methods which depend on the [class](#page-0-0) of the first argument. Currently, specific methods exist for several meta-analytical models in various packages: [blup.mixmeta](#page-0-0), [blup.rma.uni](#page-0-0), [blup.mvmeta](#page-8-1), and [blup.dosresmeta](#page-0-0).

#### Usage

blup(object, ...)

#### Arguments



## Details

The generic method function blup calls specific method functions to produces (empirical) best linear unbiased predictions (BLUPs) from model objects.

These predictions are a shrunk version of unit-specific realizations, where unit-specific estimates borrow strength from the assumption of an underlying (potentially multivariate) distribution in a (usually hypothetical) population. The amount of shrinkage depends from the relative size of the within and between-unit covariance matrices.

## Value

The form of the value returned by blup depends on the class of its argument. See the documentation of the particular methods for details of what is produced by that method. Usually, the results consist of point estimates of BLUPs and optionally some measure of their uncertainty.

## Author(s)

Antonio Gasparrini <<antonio.gasparrini@lshtm.ac.uk>> and Francesco Sera <<francesco.sera@lshtm.ac.uk>>

#### References

Verbeke G, Molenberghs G. *Linear Mixed Models for Longitudinal Data*. Springer; 1997.

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

## See Also

Specific methods for various classes: [blup.mixmeta](#page-0-0), [blup.rma.uni](#page-0-0), [blup.mvmeta](#page-8-1), and [blup.dosresmeta](#page-0-0).

<span id="page-8-1"></span><span id="page-8-0"></span>

#### Description

This method function computes (empirical) best linear unbiased predictions from fitted univariate or multivariate meta-analytical models represented in objects of class "mvmeta". Such predictions are optionally accompanied by standard errors, prediction intervals or the entire (co)variance matrix of the predicted outcomes.

#### Usage

```
## S3 method for class 'mvmeta'
blup(object, se=FALSE, pi=FALSE, vcov=FALSE, pi.level=0.95,
  format=c("matrix","list"), aggregate=c("stat","y"), ...)
```
## Arguments



#### Details

The method function blup produces (empirical) best linear unbiased predictions from mvmeta objects. For random-effects models, predictions are given by the sum of the estimated mean outcomes from the fixed part of the model, plus study-specific deviations predicted as random effects given the between-study distribution.

Predicted outcomes from blup are a shrunk version of study-specific realizations, where studyspecific estimates borrow strength from the assumption of an underlying multivariate distribution of outcomes in a (usually hypothetical) population of studies. In practice, the results from blup represent a weighted average between population mean outcomes (estimated by the fixed part of the model) and study-specific estimates. The weights depend from the relative size of the within and between-study covariance matrices reported as components S and Psi in mvmeta objects (see [mvmetaObject](#page-44-1)).

Fixed-effects models do not assume study-specific random effects, and the results of blup for these models are identical to [predict](#page-52-1) with interval="confidence".

<span id="page-9-0"></span>How to handle predictions for studies removed from estimation due to invalid missing pattern is determined by the na.action argument used in [mvmeta](#page-24-1) to produce object. If na.action=na.omit, studies excluded from estimation will not appear, whereas if na. action=na. exclude they will appear, with values set to NA for all the outcomes. This step is performed by [napredict](#page-0-0). See Note below.

In the presence of missing values in the study-specific estimated outcome y of the fitted model, correspondent values of point estimates and covariance terms are set to 0, while the variance terms are set to 1e+10. In this case, in practice, the study-specific estimates do not provide any information (their weight is virtually 0), and the prediction tends to the value returned by [predict](#page-52-1) with interval="prediction", when applied to a new but identical set of predictors. See also Note below.

#### Value

The results may be aggregated in matrices (the default), or returned as lists, depending on the argument format. For multivariate models, the aggregation is ruled by the argument aggregate, and the results may be grouped by statistic or by outcome. If vcov=TRUE, lists are always returned.

## **Note**

The definition of missing in model frames used for estimation in [mvmeta](#page-24-1) is different than that commonly adopted in other regression models such as [lm](#page-0-0) or [glm](#page-0-0). See info on [missing values](#page-49-1) in [mvmeta](#page-24-1).

Differently from [predict](#page-52-1), this method function computes the predicted values in the presence of partially missing outcomes. Interestingly, BLUPs for missing outcomes may be slightly different than predictions returned by [predict](#page-52-1) on a new but identical set of predictors, as the BLUP also depends on the random part of the model. Specifically, the function uses information from the between-study covariance to predict missing outcomes given the observed ones.

## Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## **References**

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

#### See Also

See [predict](#page-52-1) for standard predictions. See [mvmeta-package](#page-1-1) for an overview of the package and modelling framework.

#### <span id="page-10-0"></span>coef.mvmeta 11

## Examples

```
# RUN THE MODEL
model <- mvmeta(cbind(PD,AL)~1,S=berkey98[5:7],data=berkey98)
# ONLY BLUP
blup(model)
# BLUP AND SE
blup(model,se=TRUE)
# SAME AS ABOVE, AGGREGATED BY OUTCOME, WITH PREDICTION INTERVALS
blup(model,se=TRUE,pi=TRUE,aggregate="y")
# WITH VCOV, FORCED TO A LIST
blup(model,se=TRUE,pi=TRUE,vcov=TRUE,aggregate="y")
```
<span id="page-10-1"></span>coef.mvmeta *Extract Coefficients and (Co)Variance Matrix from mvmeta Objects*

## <span id="page-10-2"></span>Description

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

## Usage

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

## Arguments



#### Value

For coef, a vector (default) or matrix with the estimated fixed-effects coefficients. The matrixstructure is used to store the fixed-effects coefficients in mvmeta objects, and is preserved if format="matrix".

For vcov, the (co)variance matrix of the estimated fixed-effects coefficients.

## Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## See Also

See [mvmeta-package](#page-1-1) for an overview of the package and modelling framework.

## Examples

```
# RUN THE MODEL
model <- mvmeta(cbind(PD,AL)~pubyear,S=berkey98[5:7],data=berkey98)
# COEFFICIENTS
model$coef
coef(model)
coef(model,format="matrix")
summary(model)$coef
# (CO)VARIANCE MATRIX
vcov(model)
```
<span id="page-11-1"></span>

fibrinogen *Fibrinogen Studies Collaboration*

## Description

The Fibrinogen Studies Collaboration is a meta-analysis of individual data on 154,012 adults from 31 prospective cohort studies with information on plasma fibrinogen and major disease outcomes. The dataset reports a subset of the results of a first-stage analysis consisting of the log-hazard ratio of coronary heart disease for categories of levels of fibrinogen versus a baseline category.

## Usage

fibrinogen

## Format

A data frame with 31 observations on the following 15 variables:

- cohort: study ID.
- b2, b3, b4, b5: estimated log-hazard ratios for the second to fifth categories versus the baseline category.
- V\_2\_2, V\_3\_3, V\_4\_4, V\_5\_5: variances of the estimated log-hazard ratios.
- $V_2$  2,3,  $V_2$  2,4,  $V_2$  2,5,  $V_3$  3,4,  $V_3$  3,5,  $V_4$  4,5: covariances of the estimated log-hazard ratios.

## Details

The published analysis adopted a fixed-effects model on 10 categories of fibrinogen (Fibrinogen Studies Collaboration 2004, 2005). Here a subset of the results of the first-stage analysis is reported, namely the log-hazard ratio for 4 categories and associated (co)variance terms, ordered as the lower triangular elements of the (co)variance matrix taken by column. Details on the first-stage model and the second-stage meta-analysis are provided in White (2009) and Jackson and colleagues (2010).

<span id="page-11-0"></span>

<span id="page-12-0"></span>hsls and the set of the

## Note

The data provide an example of application of multivariate meta-analysis for multi-parameter association, where a relationship is defined by functions specified by several coefficients. In this case, the coefficients refer to log-hazard ratio for strata of the original variable versus a baseline category. A general overview of the application of multivariate meta-analysis in this setting is provided by Gasparrini and colleagues (2012).

## Source

Fibrinogen Studies Collaboration (2004). Collaborative meta-analysis of prospective studies of plasma fibrinogen and cardiovascular disease. *European Journal of Cardiovascular Prevention and Rehabilitation*. 11:9–17.

Fibrinogen Studies Collaboration (2005). Plasma fibrinogen level and the risk of major cardiovascular diseases and nonvascular mortality: an individual participant meta-analysis. *Journal of the American Medical Association*. 294:1799–1809.

White IR (2009). Multivariate random-effects meta-analysis. *Stata Journal*. 9(1):40–56.

Jackson D, White IR, Thompson SG (2010). Extending DerSimonian and Laird's methodology to perform multivariate random effects meta-analyses. *Statistics in Medicine*. 29(12):1282–1297.

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

## Examples

### REPRODUCE THE RESULTS IN WHITE (2009) AND JACKSON ET AL. (2010)

```
# INSPECT THE DATA
head(fibrinogen)
```

```
# REML MODEL
y <- as.matrix(fibrinogen[2:5])
S <- as.matrix(fibrinogen[6:15])
model <- mvmeta(y,S)
```

```
# SUMMARIZE THE RESULTS
print(summary(model),digits=3)
round(model$Psi,3)
```
#### Description

This is a nationally representative, longitudinal study of more than 21,000 9th graders in 944 schools who will be followed through their secondary and postsecondary years. The data are used for testing whether sex, socioeconomic status and sex by socio-economic status interaction are predictive of the mathematics standardized score in each of the eight race groups.

## Usage

hsls

## Format

A data frame with 8 observations on the following 10 variables:

- race: race group.
- b1, b2, b3: estimated regression coefficients for sex, socio-economic status and sex by socioeconomic status interaction, respectively, on the mathematics standardized score.
- V11, V22, V33: variances of the estimated coefficients.
- V12, V13, V23: covariances of the estimated coefficients.

## Source

Chen H, Manning AK, Dupuis J (2012). A method of moments estimator for random effect multivariate meta-analysis. *Biometrics*. 68(4):1278-1284.

## Examples

```
### REPRODUCE THE RESULTS IN CHEN ET AL. (2012)
# INSPECT THE DATA
hsls
# FIXED-EFFECTS MODEL
S \leftarrow as_matrix(hsls[5:10])model <- mvmeta(cbind(b1,b2,b3),S,data=hsls,method="fixed")
summary(model)
# MM MODEL
model <- mvmeta(cbind(b1,b2,b3),S,data=hsls,method="mm")
summary(model)
model$Psi
```
## <span id="page-14-1"></span><span id="page-14-0"></span>Description

The dataset contains the results of ten studies that assess the effectiveness of hypertension treatment for lowering blood pressure. Each study provides complete data on two treatment effects, the difference in systolic blood pressure (SBP) and diastolic blood pressure (DBP) between the treatment and the control groups, where these differences are adjusted for the participants' baseline blood pressures. The within-study correlations of the two outcomes are known. Some trials are conducted on patients with isolated systolic hypertension (ISH).

## Usage

hyp

## Format

A data frame with 10 observations on the following 7 variables:

- study: study ID.
- sbp, sbp\_se: estimated difference and its standard error in systolic blood pressure.
- dbp, dbp\_se: estimated difference and its standard error in diastolic blood pressure.
- rho: within-study correlation between the estimated differences in systolic and diastolic blood pressure.
- ish: indicator for studies on patients with isolated systolic hypertension.

## Note

The standard errors for the two outcomes are wrongly reported as variances in the original article by Jackson and colleagues (2013).

## Source

Jackson D, White IR, Riley RD (2013). A matrix based method of moments for fitting the multivariate random effects model for meta-analysis and meta-regression. *Biometrical Journal*. 55(2):231– 45.

## Examples

```
### REPRODUCE THE RESULTS IN JACKSON ET AL. (2013)
# INSPECT THE DATA
hyp
# INPUT THE CORRELATION (CAN ALSO BE INPUTTED DIRECTLY, SEE BELOW)
(S <- inputcov(hyp[c("sbp_se","dbp_se")],cor=hyp$rho))
```
#### <span id="page-15-0"></span>16 inputcov and the set of the set

```
# CHECK WITH THE FIRST STUDY
cov2cor(xpndMat(S[1,]))
# META-ANALYSIS, REML MODEL
model <- mvmeta(cbind(sbp,dbp),S=S,data=hyp)
print(summary(model),digits=2)
round(model$Psi,2)
# META-ANALYSIS, REML MODEL (INPUTTING THE CORRELATION DIRECTLY)
model <- mvmeta(cbind(sbp,dbp),S=cbind(sbp_se,dbp_se)^2,data=hyp,
  control=list(Scor=hyp$rho))
print(summary(model),digits=2)
# META-ANALYSIS, MM MODEL
model <- mvmeta(cbind(sbp,dbp), S=S,data=hyp,method="mm")
print(summary(model),digits=2)
round(model$Psi,2)
# META-REGRESSION, REML MODEL
model <- mvmeta(cbind(sbp,dbp)~ish,S=S,data=hyp)
print(summary(model),digits=2)
# META-REGRESSION, MM MODEL
model <- mvmeta(cbind(sbp,dbp)~ish,S=S,data=hyp,method="mm")
```

```
print(summary(model),digits=2)
```
<span id="page-15-1"></span>inputcov *Input (Co)Variance Matrices*

## Description

This function inputs (co)variance matrices of a set of outcomes given the corresponding standard deviation and correlation values.

#### Usage

```
inputcov(sd, cor=NULL)
```
## Arguments



## <span id="page-16-0"></span>inputcov and the contract of t

## Details

Depending the number of outcomes  $k$  and matrices  $m$ , the argument cor is interpreted as:

- if a vector of length 1 (a scalar), the same correlation for all the k outcomes for all the  $m$ matrices;
- if a vector of length m, the same correlation for all the  $k$  outcomes for each of the m matrices;
- if a vector of length  $k(k-1)/2$ , the lower triangular elements (without diagonal, taken by column) of the correlation matrix of the  $k$  outcomes, the same for all the  $m$  matrices;
- if a  $k \times k$  matrix, the correlation matrix for the single matrix (only when  $m=1$ );
- if a  $m \times k(k-1)/2$  matrix, each row represents the lower triangular elements (without diagonal, taken by column) of the correlation matrix of the k outcomes for each of the  $m$  matrices.

## Value

For a single matrix, the (co)variance matrix itself. For multiple matrices, a  $m \times k(k+1)/2$  matrix, where each row represents the vectorized entries of the lower triangle (with diagonal, taken by column) of the related (co)variance matrix (see [vechMat](#page-60-1)).

#### Note

This function is imported from the package **mixmeta**. It is called internally by mymeta for multivariate models to input the correlation(s) when only the within-unit variances are provided through the argument S. In this case, the correlation values are set through the argument Scor in the control list (see [mvmeta.control](#page-29-1)).

## Author(s)

Antonio Gasparrini <<antonio.gasparrini@lshtm.ac.uk>>

## See Also

See [xpndMat](#page-60-2). See [mvmeta.control](#page-29-1).

## Examples

```
# SOME RANDOM SD FOR A SINGLE MATRIX, WITH CONSTANT CORRELATION
(M <- inputcov(runif(4, 0.1, 3), 0.7))
# CHECK CORRELATION
cov2cor(M)
```

```
# NOW WITH A MORE COMPLEX CORRELATION STRUCTURE
(M <- inputcov(runif(3, 0.1, 3), c(0.7,0.2,0.4)))
cov2cor(M)
```

```
# MULTIPLE MATRICES
(V <- matrix(runif(5*3, 0.1, 3), 5, 3,
 dimnames=list(1:5, paste("V", 1:3, sep=""))))
inputcov(V, 0.6)
```

```
# WITH REAL DATA WHEN CORRELATIONS AVAILABLE
hyp
(S <- inputcov(hyp[c("sbp_se","dbp_se")], cor=hyp$rho))
# CHECK FIRST STUDY
cov2cor(xpndMat(S[1,]))
# USED INTERNALLY IN mvmeta
p53
inputcov(sqrt(p53[c("V1","V2")]), 0.5)
model <- mvmeta(cbind(y1,y2), S=cbind(V1,V2), data=p53, control=list(Scor=0.5))
model$S
```
<span id="page-17-1"></span>

inputna *Input Missing Values*

## Description

This function augment data by replacing missing values. It can be used internally in [mvmeta](#page-24-1) through the [control](#page-29-1) list.

## Usage

inputna(y, S, inputvar=10^4)

## Arguments

Assuming a meta-analysis or meta-regression based on  $n$  units and  $k$  outcomes:



## Details

The function augments the data by replacing missing values in the outcomes and the associated (co)variances. Specifically, it replaces missing outcomes and missing covariances (if provided) with 0, and missing variances with the largest observed variance multiplied by inputvar. This value is expected to be very high, by default  $10<sup>4</sup>$ , so that the corresponding observation contributes only negligibly to the final estimate.

<span id="page-17-0"></span>

#### <span id="page-18-0"></span>inputna 19

#### Value

A matrix with the first k column corresponding to the augmented outcomes, and the remaining  $k(k+1)$  $1/2$  or k columns (depending on the availability of the within-study covariances) corresponding to vectorized entries of the lower triangle of the related (co)variance matrices.

#### Note

Data augmentation used to be the approach to deal with missing values in the first implementation of mvmeta. The current algorithms directly account for missing. This function is now imported from the package mixmeta.

Inputting missing values can be useful when two or more outcomes are never observed jointly, and the estimation is entirely based on indirect comparison. This method can be applied in network meta-analysis, also called indirect treatment comparison.

This approach can produce different results than standard methods, especially when the occurrence of missing is substantial. Preliminary analyses indicate that likelihood-based estimation methods do not seem to be affected, while non-iterative estimators such as method of moments and variance components are more sensitive. The user should be careful on the application of missing augmentation.

## Author(s)

Antonio Gasparrini << antonio.gasparrini@lshtm.ac.uk>>

## References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

Jackson D, Riley R, White IR (2011). Multivariate meta-analysis: Potential and promise. *Statistics in Medicine*. 30(20);2481–2498.

White IR (2009). Multivariate random-effects meta-analysis. *Stata Journal*. 9(1):40–56.

White IR (2011). Multivariate random-effects meta-regression: updates to mvmeta. *Stata Journal*. 11(2):255-270.

## See Also

See [inputcov](#page-15-1) for inputting (co)variance matrices.

## Examples

```
# INSPECT THE DATA
head(smoking)
```
# STANDARD APPROACH TO MISSING DATA y <- as.matrix(smoking[11:13]) S <- as.matrix(smoking[14:19])

```
mod1 <- mvmeta(y, S)summary(mod1)
# WITH DATA AUGMENTATION
augdata <- inputna(y, S)
y \leq - augdata[,1:3]
S <- augdata[, -c(1:3)]mod2 <- mvmeta(y, S)
summary(mod2)
# NB: SAME PARAMETER ESTIMATES, BUT WRONG NYUMBER OF OBS
# USED INTERNALLY IN mvmeta
y <- as.matrix(smoking[11:13])
S <- as.matrix(smoking[14:19])
mod3 <- mvmeta(y, S, control=list(inputna=TRUE))
summary(mod3)
# NOW RIGHT NUMBER OF OBS
```
<span id="page-19-1"></span>

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

## Description

This method function returns the log-likelihood for fitted univariate or multivariate meta-analytical models represented in objects of class "mvmeta".

#### Usage

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



#### Value

A numeric scalar of class "logLik" with attributes, providing the (restricted) log likelihood of the model. Attributes correspond to the component df of mvmeta objects, namely the following scalars: nall (number of observations used for estimation, excluding missing values), nobs (equal to nall, minus the number of fixed-effects coefficients for REML models, fixed (number of estimated fixed-effects coefficients), random (number of estimated (co)variance terms).

#### **Note**

This functions is called by [AIC](#page-0-0) and [BIC](#page-0-0) for computing the Akaike and Bayesian information criteria.

#### <span id="page-20-0"></span>mlprof.fn 21

## Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

#### References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

## See Also

See the default method [logLik](#page-0-0). See [mvmeta-package](#page-1-1) for an overview of the package and modelling framework.

## Examples

```
# RUN THE MODEL
model <- mvmeta(cbind(PD,AL)~pubyear,S=berkey98[5:7],data=berkey98)
```

```
# LOG-LIKELIHOOD
ll <- logLik(model)
ll
attributes(ll)
# AIC and BIC
AIC(model)
BIC(model)
```
<span id="page-20-1"></span>mlprof.fn *Likelihood Functions for mvmeta Models*

#### <span id="page-20-2"></span>Description

These functions compute the value of the log-likelihood and the related vectors of first partial derivatives for random-effects multivariate and univariate meta-analysis and meta-regression, in terms of model parameters. They are meant to be used internally and not directly run by the users.

#### Usage

```
mlprof.fn(par, Xlist, ylist, Slist, nalist, k, m, p, nall, bscov, ctrl)
mlprof.gr(par, Xlist, ylist, Slist, nalist, k, m, p, nall, bscov, ctrl)
remlprof.fn(par, Xlist, ylist, Slist, nalist, k, m, p, nall, bscov, ctrl)
remlprof.gr(par, Xlist, ylist, Slist, nalist, k, m, p, nall, bscov, ctrl)
iter.igls(Psi, Xlist, ylist, Slist, nalist, k, m)
```
## <span id="page-21-0"></span>Arguments

Assuming a meta-analysis or meta-regression based on m studies, k outcomes and p predictors:



## Details

These functions are called internally by the fitting functions mymeta.ml and mymeta.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](#page-0-0). Alternatively, starting values can be provided by the user in the control list (see [mvmeta.control](#page-29-1)). The function [optim](#page-0-0) requires the algorithms to compute the value of the (restricted) likelihood and (optionally) the vector of its first partial derivatives, provided by the related likelihood functions.

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 betweenstudy (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, especially for meta-regression models.

The random-effects parameters in par depends on the chosen [structure](#page-42-1) for the between-study (co)variance matrix. The parameterization ensures the positive-definiteness of the estimated matrix. A Cholesky decomposition is then performed on the marginal (co)variance matrix in order to

#### <span id="page-22-0"></span>mlprof.fn 23

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. More details are provided in the references below.

Some parameters of the fitting procedures are determined through [mvmeta.control](#page-29-1). Specifically, the user can obtain the Hessian matrix of the estimated parameters (appropriately transformed, see [mvmetaCovStruct](#page-42-1)) in the optimization function by setting hessian=TRUE, and specific control settings in the optimization process can be defined by the control list argument optim. These values are passed to the [optim](#page-0-0)ization function optim.

## Value

mlprof.fn and remlprof.fn return the value of the (restricted) log-likelihood for a given set of parameters in par. mlprof.gr and remlprof.gr return instead the related vector of first partial derivatives. iter. igls returns an updated estimate of Psi given its initial value or the value at the previous iteration.

#### Note

As stated earlier, these functions are called internally by mymeta.ml and mymeta.reml, and are not meant to be used directly. In particular, their code does not contain any check on the arguments provided, which are expected in specific formats. They are however exported in the namespace and documented for completeness.

#### Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

Goldstein H (1986). Multilevel mixed linear model analysis using iterative generalized least squares. *Biometrika*. 73(1):43.

Lindstrom MJ and Bates DM (1988). Newton-Raphson and EM algorithms for linear mixed-effects models for repeated-measures data. *Journal of the American Statistical Association*. 83(404):1014– 1022.

Pinheiro JC and Bates DM (2000). *Mixed-Effects Models in S and S-PLUS*. New York, Springer Verlag.

## See Also

See [mvmeta.fit](#page-24-2) and [mvmeta.ml](#page-34-1) for additional info on the fitting procedures. See [mvmeta.control](#page-29-1) to determine specific parameters of the fitting procedures. See [mvmetaCovStruct](#page-42-1) for (co)variance structures. See [chol](#page-0-0) and [qr](#page-0-0) for info on the Cholesky and QR decomposition. See [mvmeta-package](#page-1-1) for an overview of the package and modelling framework.

<span id="page-23-1"></span><span id="page-23-0"></span>model.frame.mvmeta *Extract Model Frame and Design Matrix from mvmeta Objects*

## <span id="page-23-2"></span>**Description**

These method functions return the model frame and design matrix for univariate or multivariate meta-analytical models represented in objects of class "mvmeta".

#### Usage

```
## S3 method for class 'mvmeta'
model.frame(formula, ...)
## S3 method for class 'mvmeta'
model.matrix(object, ...)
```
## Arguments

object, formula an object of class "mvmeta".

... further arguments passed to or from other methods.

#### Details

The model frame is produced by [mvmeta](#page-24-1) when fitting the meta-analytical model, and stored in the mvmeta object if argument model=TRUE. Alternatively, the model frame is directly returned by [mvmeta](#page-24-1) with argument method="model.frame". The method function model.frame simply extracts the saved model frame if available, or otherwise evaluates a call to mymeta when method="model.frame".

The method function model.matrix extracts the design matrix from a fitted meta-analytical model. It first extract the model frame by calling model.frame, and then passes the call to the default method.

These methods functions are similar to those provided for regression objects  $\text{lm}$  $\text{lm}$  $\text{lm}$  and  $\text{lm}$ .

## Value

For [model.frame](#page-0-0), a data.frame with special attributes (see the default method model.frame) and the additional class "data.frame.mvmeta".

For model.matrix, the design matrix used to fit the model.

## **Note**

The reason why these specific method functions are made available for class mvmeta, and in particular why a new class "data.frame.mvmeta" has been defined for model frames, lies in the special handling of missing values in multivariate meta-analysis models fitted with [mvmeta](#page-24-1). Methods [na.omit](#page-49-1) and [na.exclude](#page-49-2) for class "data.frame.mvmeta" are useful for properly accounting for missing values when fitting these models.

#### <span id="page-24-0"></span>mvmeta 25

#### Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## See Also

See the default methods [model.frame](#page-0-0) and [model.matrix](#page-0-0). See [na.omit](#page-49-1) and [na.exclude](#page-49-2) on the handling of missing values. See [mvmeta-package](#page-1-1) for an overview of the package and modelling framework.

## Examples

```
# RUN THE MODEL AND SUMMARIZE THE RESULTS
model <- mvmeta(cbind(PD,AL)~pubyear,S=berkey98[5:7],data=berkey98,method="ml")
# MODEL FRAME
model$model
model.frame(model)
update(model,method="model.frame")
class(model.frame(model))
# MODEL MATRIX
model.matrix(model)
```
<span id="page-24-1"></span>mvmeta *Fitting Multivariate and Univariate Meta-Analysis and Meta-Regression Models*

## <span id="page-24-2"></span>Description

The function mvmeta performs fixed and random-effects multivariate and univariate meta-analysis and meta-regression, with various estimation methods. The function mvmeta.fit is a wrapper for actual fitting functions based on different estimation methods, usually called internally. See [mvmeta-package](#page-1-1) for an overview.

#### Usage

```
mvmeta(formula, S, data, subset, method="reml", bscov="unstr", model=TRUE,
 contrasts=NULL, offset, na.action, control=list())
```
mvmeta.fit(X, y, S, offset=NULL, method="reml", bscov="unstr", control=list())

## Arguments

Assuming a meta-analysis or meta-regression based on  $m$  studies,  $k$  outcomes and  $p$  predictors:

formula an object of class ["formula"](#page-0-0) (or one that can be coerced to that class) offering a symbolic description of the linear predictor of the model to be fitted to each outcome. Alternatively, for meta-analysis with no predictor, a single vector (for

<span id="page-25-0"></span>

## Details

The function mvmeta resembles standard regression functions in R (see [lm](#page-0-0) or [glm](#page-0-0)). This function defines the design matrix and the vector (for univariate models) or matrix (for multivariate models)

#### <span id="page-26-0"></span>mvmeta 27

of outcome responses, and calls the wrapper mvmeta.fit to perform the actual fitting. The latter prepares the data and calls specific fitting functions, depending on the chosen method. Functions other than mvmeta are not expected to be called directly for model fitting.

The model is specified through a regression formula. Simple meta-analysis is specified with the formula  $y \sim 1$ , where the left-hand side is a vector (in univariate models) or a matrix (in multivariate models), optionally of form  $\text{cbind}(y1, \ldots, yk)$ , with terms stored in data. Alternatively, matrix or vector objects are allowed, and the formula is retrieved internally adding ~ 1. In meta-regression, other terms are added in the right-hand side of the formula, defining the linear predictor common to all outcomes. Factors, variable transformations and interactions are allowed, following the standard formula specification. Labels are automatically retrieved from the objects in formula. See [formula](#page-0-0) for further details. See  $\text{Im} \text{ or } \text{glm}$  $\text{Im} \text{ or } \text{glm}$  $\text{Im} \text{ or } \text{glm}$  for info on the other arguments.

The within-study (co)variances are provided through the argument S, usually as a matrix. If the correlations are available, each of the m row represents the  $k(k + 1)/2$  vectorized entries of the lower triangle of the related (co)variance matrix, taken by column (see [xpndMat](#page-60-2)). If correlations are not available, each row represents the  $k$  variances, and the correlations are inputted internally through the argument Scor of the control list. See [inputcov](#page-15-1).

Different estimator are available in the package mvmeta and chosen through the argument method. In the current version, the options are:

- method="fixed": [Fixed-effects estimator](#page-32-1)
- method="ml": [Maximum likelihood \(ML\) estimator](#page-34-1)
- method="reml": [Restricted maximum likelihood \(REML\) estimator](#page-34-2)
- method="mm": [Method of moments estimator](#page-37-1)
- method="vc": [Variance components estimator](#page-39-1)

Specific fitting functions are called internally. Likelihood-based methods allow alternative [\(co\)vari](#page-42-1)ance [structures](#page-42-1) for the between-study random effects through the argument bscov. See their help pages for further details on the estimation methods, following the links above.

Missing values are allowed in both sides of formula. In the case of missing predictors (right-hand side of formula), the related study is entirely excluded from estimation. In contrast, a study still contributes to estimation if at least outcome is non-missing. This behaviour is different than in standard regression functions such as  $\text{Im}$  or g[lm](#page-0-0). Before the call to mvmeta. fit, studies matching such missing definition are removed from the the model frame. The missing pattern in S must be consistent with that in y. See further details on handling [missing values](#page-49-1) in mvmeta.

The fitting procedure can be controlled through the additional terms specified in control, which are passed to the function [mvmeta.control](#page-29-1).

#### Value

The mvmeta function typically returns a list object of class "mvmeta" representing the meta-analytical model fit, as described in mymetaObject. When method="data.frame", the model is not fitted and the model frame is returned, namely a data frame with special attributes (see the default method [model.frame](#page-0-0)) and, in this case, the additional class "data.frame.mvmeta".

The wrapper function mvmeta. fit is usually called internally in mvmeta, and returns an intermediate list object with some of the components expected in the "mvmeta" class.

Several method functions for regression objects are available, either default or specifically-written for the "mvmeta" class. See [mvmetaObject](#page-44-1) for a complete list.

<span id="page-27-0"></span>In the current version, the same linear predictor specified in formula is set for all the outcomes.

#### Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

Jackson D, Riley R, White IR (2011). Multivariate meta-analysis: Potential and promise. *Statistics in Medicine*. 30(20);2481–2498.

White IR (2009). Multivariate random-effects meta-analysis. *Stata Journal*. 9(1):40–56.

White IR (2011). Multivariate random-effects meta-regression: updates to mvmeta. *Stata Journal*. 11(2):255-270.

Berkey, CS, Hoaglin DC, et al. (1998). Meta-analysis of multiple outcomes by regression with random effects. *Statistics in Medicine*. 17(22):2537–2550.

## See Also

See additional info on the estimation procedures at the related page of the fitting functions. See alternative [\(co\)variance structures](#page-42-1) for likelihood-based estimation methods. See handling of [missing values](#page-49-1) in mvmeta. See [lm](#page-0-0) or [glm](#page-0-0) for standard regression functions. See [mvmeta-package](#page-1-1) for an overview of this modelling framework.

#### Examples

### BIVARIATE META-ANALYSIS, ESTIMATED THROUGH REML

# RUN THE MODEL model <- mvmeta(cbind(PD,AL),S=berkey98[5:7],data=berkey98)

# SUMMARIZE THE RESULTS summary(model)

```
# RESIDUALS AND FITTED VALUES
residuals(model)
fitted(model)
```

```
# LOG-LIKELIHOOD AND AIC VALUE
logLik(model)
AIC(model)
```
### BIVARIATE META-REGRESSION, ESTIMATED THROUGH METHOD OF MOMENTS

#### mvmeta 29

```
# RUN THE MODEL AND SUMMARIZE THE RESULTS
model <- mvmeta(cbind(PD,AL)~pubyear,S=berkey98[5:7],data=berkey98,method="mm")
summary(model)
# BLUP ESTIMATES AND 90% PREDICTION INTERVALS, AGGREGATED BY OUTCOME
blup(model,pi=TRUE,aggregate="y",pi.level=0.90)
# COCHRAN Q TEST FOR RESIDUAL HETEROGENEITY
qtest(model)
# PREDICTED AVERAGED OUTOCOMES AND STANDARD ERRORS FROM YEAR 1985 TO 1989
newdata <- data.frame(pubyear=1985:1989)
predict(model,newdata,se=TRUE)
# MODEL FRAME AND MODEL MATRIX
model.frame(model)
model.matrix(model)
### UNIVARIATE META-REGRESSION, FIXED-EFFECTS MODEL
# RUN THE MODEL
model <- mvmeta(sbp~ish,S=sbp_se^2,data=hyp,method="fixed")
summary(model)
# RESIDUALS AND FITTED VALUES
residuals(model)
fitted(model)
# COCHRAN Q TEST FOR RESIDUAL HETEROGENEITY
qtest(model)
### MULTIVARIATE META-ANALYSIS WITH MORE THAN 2 OUTCOMES
# RUN THE MODEL
y <- as.matrix(fibrinogen[2:5])
S <- as.matrix(fibrinogen[6:15])
model \leq mwmeta(y, S)summary(model)
### IN THE PRESENCE OF MISSING VALUES
# RUN THE MODEL
y <- as.matrix(smoking[11:13])
S <- as.matrix(smoking[14:19])
model <- mvmeta(y,S)
summary(model)
```
# MODEL FRAME: SEE help(na.omit.data.frame.mvmeta) FOR MORE EXAMPLES model.frame(model)

```
### WHEN WITHIN-STUDY COVIARIANCES ARE NOT AVAILABLE AND/OR NEED TO BE INPUTTED
# GENERATE S
(S <- inputcov(hyp[c("sbp_se","dbp_se")],cor=hyp$rho))
# RUN THE MODEL
model <- mvmeta(cbind(sbp,dbp), S=S,data=hyp)
# INPUTTING THE CORRELATION DIRECTLY IN THE MODEL
model <- mvmeta(cbind(y1,y2),cbind(V1,V2),data=p53,control=list(Scor=0.95))
summary(model)
# SEE help(hyp) AND help(p53) FOR MORE EXAMPLES
### STRUCTURING THE BETWEEN-STUDY (CO)VARIANCE
# DIAGONAL
S \leftarrow as_matrix(hsls[5:10])model <- mvmeta(cbind(b1,b2,b3),S,data=hsls,bscov="diag")
summary(model)
model$Psi
# COMPOUND SYMMETRY
model <- mvmeta(cbind(b1,b2,b3),S,data=hsls,bscov="cs")
summary(model)
model$Psi
# SEE help(mvmetaCovStruct) FOR DETAILS AND ADDITIONAL EXAMPLES
### USE OF THE CONTROL LIST
# PRINT THE ITERATIONS AND CHANGE THE DEFAULT FOR STARTING VALUES
y <- as.matrix(smoking[11:13])
S <- as.matrix(smoking[14:19])
model <- mvmeta(y,S,control=list(showiter=TRUE,igls.iter=20))
# SEE help(mvmeta.control) FOR FURTHER DETAILS
```
<span id="page-29-1"></span>mvmeta.control *Ancillary Parameters for Controlling the Fit in mvmeta Models*

## Description

This internal function sets the parameter options used for fitting meta-analytical models, commonly to pre-specified default values. It is usually internally called by [mvmeta.fit](#page-24-2).

## <span id="page-30-0"></span>mvmeta.control 31

## Usage

```
mvmeta.control(optim=list(), showiter=FALSE, maxiter=100, initPsi=NULL,
 Psifix=NULL,Psicor=0, Scor=0, inputna=FALSE, inputvar=10^4, igls.iter=10,
 hessian=FALSE, vc.adj=TRUE,reltol=sqrt(.Machine$double.eps),
  set.negeigen=sqrt(.Machine$double.eps))
```
## Arguments



<span id="page-31-0"></span>set.negeigen positive value. Value to which negative eigenvalues are to be set in estimators where such method is used to force positive semi-definiteness of the estimated between-study (co)variance matrix.

## **Details**

The control argument of [mvmeta](#page-24-1) is by default passed to [mvmeta.fit](#page-24-2), which uses its elements as arguments of mvmeta.control.

Many arguments refer to specific fitting procedures. Refer to the help page of the related estimator for details.

The function automatically sets non-default values for some control arguments for [optim](#page-0-0), unless explicitly set in the list passed to it. Specifically, the function selects fnscale=-1, maxit=maxiter and reltol=reltol, where the latter two are specified by other arguments of this function.

The function is expected to be extended and/or modified at every release of the package **mvmeta**.

## Value

A list with components named as the arguments.

#### Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

#### References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

## See Also

See [mvmeta](#page-24-1). See also [glm.control](#page-0-0). See the help pages of the related fitting functions for details on each parameter. See [mvmeta-package](#page-1-1) for an overview of this modelling framework.

## Examples

```
# PRINT THE ITERATIONS (SEE ?optim) AND CHANGE THE DEFAULT FOR STARTING VALUES
model <- mvmeta(cbind(PD,AL)~pubyear,S=berkey98[5:7],data=berkey98,
 control=list(showiter=TRUE,igls.iter=20))
# INPUT THE CORRELATION
model \leq mymeta(cbind(y1,y2),S=cbind(V1,V2),data=p53,control=list(Scor=0.5))
```
## <span id="page-32-1"></span><span id="page-32-0"></span>Description

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

#### Usage

```
mvmeta.fixed(Xlist, ylist, Slist, nalist, k, m, p, nall, control, ...)
```
#### Arguments

Assuming a meta-analysis or meta-regression based on  $m$  studies,  $k$  outcomes and  $p$  predictors:



## Details

The estimation involves only the  $kp$  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 mymeta.ml and [mvmeta.reml](#page-34-2), the computation involves Cholesky and and QR decompositions for computational stability and efficiency. The method is described in details in Gasparrini and collaborators (2012) (see references below).

#### <span id="page-33-0"></span>Value

This function returns an intermediate list object, whose components are then processed by mymeta. fit. Other components are added later through [mvmeta](#page-24-1) to finalize an object of class "mvmeta".

## **Note**

As stated earlier, this function is called internally by [mvmeta.fit](#page-24-2), and is not meant to be used directly. In particular, its code does not contain any check on the arguments provided, which are expected in specific formats. The function is however exported in the namespace and documented for completeness.

The arguments above are prepared by mymeta. Fit from its arguments X, y and S. The list structure, although requiring more elaborate coding, is computationally more efficient, as it avoids the specification of sparse block-diagonal matrices, especially for meta-analysis involving a large number of studies.

Some parameters of the fitting procedures are determined by the control argument, with default set by [mvmeta.control](#page-29-1). No missing values are accepted in the fitting functions. See details on [missing values](#page-49-1).

## Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

Berkey, CS, Anderson JJ, Hoaglin DC (1996). Multiple-outcome meta-analysis of clinical trials. *Statistics in Medicine*. 15(5):537–547.

Berkey, CS, Hoaglin DC, et al. (1998). Meta-analysis of multiple outcomes by regression with random effects. *Statistics in Medicine*. 17(22):2537–2550.

## See Also

See [mvmeta](#page-24-1) for the general usage of the functions. See [mvmeta.control](#page-29-1) to determine specific parameters of the fitting procedures. Use the triple colon operator (':::') to access the code of the internal functions, such as glsfit. See [mvmeta-package](#page-1-1) for an overview of the package and modelling framework.

## Examples

```
# UNIVARIATE FIXED-EFFECTS MODEL
model <- mvmeta(yC,S=SCC,data=smoking,method="fixed")
summary(model)
```
# MULTIVARIATE FIXED-EFFECTS MODEL

#### <span id="page-34-0"></span>mvmeta.ml 35

```
y <- as.matrix(smoking[11:13])
S <- as.matrix(smoking[14:19])
model <- mvmeta(y,S,method="fixed")
summary(model)
# MULTIVARIATE FIXED-EFFECTS MODEL: REPLICATE THE RESULTS IN BERKEY ET AL. 1998
model <- mvmeta(cbind(PD,AL)~I(pubyear-1983),S=berkey98[5:7],
  data=berkey98,method="fixed")
summary(model)
```

```
mvmeta.ml ML and REML Estimators for mvmeta Models
```
## <span id="page-34-2"></span>Description

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

## Usage

```
mvmeta.ml(Xlist, ylist, Slist, nalist, k, m, p, nall, bscov, control, ...)
mvmeta.reml(Xlist, ylist, Slist, nalist, k, m, p, nall, bscov, control, ...)
```
## Arguments

Assuming a meta-analysis or meta-regression based on  $m$  studies,  $k$  outcomes and  $p$  predictors:



## <span id="page-35-0"></span>Details

The estimation involves  $kp$  fixed-effects coefficients and a number of random-effects parameters defining the between-study (co)variance matrix, depending on the chosen [\(co\)variance structure](#page-42-1). Up to  $k(k + 1)/2$  parameters are needed for an unstructured form, while a smaller set are required for structed matrices

The hybrid estimation procedure is based first on few runs of iterative generalized least square algorithm and then quasi-Newton iterations, using specific [likelihood functions](#page-20-1), until convergence. The estimation algorithm adopts a profiled (or concentrated) approach, that is expressed only in terms of the random-effects parameters. Cholesky and and QR decompositions are used for computational stability and efficiency, and for assuring the positive-definiteness of the estimated between-study (co)variance matrix. See the help page for the [likelihood functions](#page-20-1) for further details. The method is described in details in Gasparrini and collaborators (2012) (see references below).

## Value

These functions return an intermediate list object, whose components are then processed by mymeta. fit. Other components are added later through [mvmeta](#page-24-1) to finalize an object of class "mvmeta".

#### Note

As stated earlier, these functions are called internally by [mvmeta.fit](#page-24-2), and are not meant to be used directly. In particular, their code does not contain any check on the arguments provided, which are expected in specific formats. The functions are not exported in the namespace, and only documented for completeness.

The arguments above are prepared by mymeta.  $fit$  from its arguments X,  $\gamma$  and S. The list structure, although requiring more elaborate coding, is computationally more efficient, as it avoids the specification of sparse block-diagonal matrices, especially for meta-analysis involving a large number of studies.

Some parameters of the fitting procedures are determined by the control argument, with default set by [mvmeta.control](#page-29-1). No missing values are accepted in the fitting functions. See details on [missing values](#page-49-1).

## Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

Pinheiro JC and Bates DM (2000). *Mixed-Effects Models in S and S-PLUS*. New York, Springer Verlag.

<span id="page-36-0"></span>Lindstrom MJ and Bates DM (1988). Newton-Raphson and EM algorithms for linear mixed-effects models for repeated-measures data. *Journal of the American Statistical Association*. 83(404):1014– 1022.

White IR (2009). Multivariate random-effects meta-analysis. *Stata Journal*. 9(1):40–56.

White IR (2011). Multivariate random-effects meta-regression: updates to mvmeta. *Stata Journal*. 11(2):255-270.

Goldstein H (1986). Multilevel mixed linear model analysis using iterative generalized least squares. *Biometrika*. 73(1):43.

## See Also

See [mvmeta](#page-24-1) for the general usage of the functions. See [mvmeta.control](#page-29-1) to determine specific parameters of the fitting procedures. Use the triple colon operator  $('::::)$  to access the code of the internal functions, such as glsfit. See [mvmeta-package](#page-1-1) for an overview of the package and modelling framework.

## Examples

```
# REML ESTIMATOR: UNIVARIATE MODEL
model <- mvmeta(yC,S=SCC,data=smoking)
summary(model)
# REML ESTIMATOR: REPRODUCE THE RESULTS IN WHITE (2011)
y <- as.matrix(smoking[11:13])
S <- as.matrix(smoking[14:19])
model <- mvmeta(y,S)
summary(model)
# ML ESTIMATOR: REPRODUCE THE RESULTS IN BERKEY ET AL. (1998)
year <- berkey98$pubyear - 1983
model <- mvmeta(cbind(PD,AL)~year,S=berkey98[5:7],data=berkey98,method="ml")
print(summary(model),digits=3)
round(model$Psi,3)
# UNSTRUCTURED AND STRUCTURED BETWEEN-STUDY (CO)VARIANCE
y <- as.matrix(fibrinogen[2:5])
S <- as.matrix(fibrinogen[6:15])
model <- mvmeta(y,S)
summary(model)
model <- mvmeta(y,S,bscov="diag")
summary(model)
model <- mvmeta(y,S,bscov="hcs")
summary(model)
```
# SEE help(mvmetaCovStruct) for additional info and examples

#### <span id="page-37-1"></span><span id="page-37-0"></span>**Description**

This function implements a method of moments estimator for multivariate and univariate randomeffects meta-analysis and meta-regression. It is meant to be used internally and not directly run by the users.

#### Usage

```
mvmeta.mm(Xlist, ylist, Slist, nalist, k, m, p, nall, control, ...)
```
## Arguments

Assuming a meta-analysis or meta-regression based on  $m$  studies,  $k$  outcomes and  $p$  predictors:



#### Details

The estimation involves kp fixed-effects coefficients and  $k(k + 1)/2$  random-effects parameters, corresponding to the lower triangular entries of the between-study (co)variance matrix.

The approach implemented here represents the multivariate extension of the traditional estimator proposed by DerSimonian and Laird (1986), and simplifies to the standard method in the univariate case. This non-iterative routine forces the positive semi-definiteness of the estimated between-study (co)variance matrix by setting its negative eigenvalues to zero.

The specific method of moment estimator used here is described in Jackson and collaborators (2013), and represents a generalization of that developed by Chen and collaborators (2012). However, this general version is computationally more intensive, and may turn out to be slow when applied to meta-analysis of a relatively high number of studies. An alternative and computationally

#### <span id="page-38-0"></span>mvmeta.mm 39

faster method of moment estimator was previously proposed by Jackson and collaborators (2010), although it is not invariant to reparameterization. This latter estimator is not implemented yet in mvmeta. See references below.

#### Value

This function returns an intermediate list object, whose components are then processed by mymeta. fit. Other components are added later through [mvmeta](#page-24-1) to finalize an object of class "mvmeta".

## Note

As stated earlier, this function is called internally by [mvmeta.fit](#page-24-2), and is not meant to be used directly. In particular, its code does not contain any check on the arguments provided, which are expected in specific formats. The function is however exported in the namespace and documented for completeness.

The arguments above are prepared by mymeta. fit from its arguments X, y and S. The list structure, although requiring more elaborate coding, is computationally more efficient, as it avoids the specification of sparse block-diagonal matrices, especially for meta-analysis involving a large number of studies.

Some parameters of the fitting procedures are determined by the control argument, with default set by [mvmeta.control](#page-29-1). No missing values are accepted in the fitting functions. See details on [missing values](#page-49-1).

## Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

Jackson D, White IR, Riley RD (2013). A matrix based method of moments for fitting the multivariate random effects model for meta-analysis and meta-regression. *Biometrical Journal*. 55(2):231- 245.

Chen H, Manning AK, Dupuis J (2012). A method of moments estimator for random effect multivariate meta-analysis. *Biometrics*. 68(4):1278-1284.

Jackson D, White IR, Thompson SG (2010). Extending DerSimonian and Laird's methodology to perform multivariate random effects meta-analyses. *Statistics in Medicine*. 29(12):1282–1297.

DerSimonian R, Laird N (1986). Meta-analysis in clinical trials. *Controlled Clinical Trials*. 7(3):177- 188.

## See Also

See [mvmeta](#page-24-1) for the general usage of the functions. See [mvmeta.control](#page-29-1) to determine specific parameters of the fitting procedures. Use the triple colon operator  $(':::)$  to access the code of the internal functions, such as fbtr. See [mvmeta-package](#page-1-1) for an overview of the package and modelling framework.

## Examples

```
# MM ESTIMATOR: UNIVARIATE MODEL
model <- mvmeta(PD~pubyear,S=berkey98[,5],data=berkey98,method="mm")
summary(model)
# MM ESTIMATOR: REPRODUCE THE RESULTS IN CHEN ET AL. (2012)
S \leftarrow as_matrix(hsls[5:10])model <- mvmeta(cbind(b1,b2,b3),S,data=hsls,method="mm")
summary(model)
# MM ESTIMATOR: REPRODUCE THE RESULTS IN JACKSON ET AL. (2013)
S <- inputcov(hyp[c("sbp_se","dbp_se")],cor=hyp$rho)
model <- mvmeta(cbind(sbp,dbp),S=S,data=hyp,method="mm")
summary(model)
```
mvmeta.vc *Variance Components Estimator for mvmeta Models*

## Description

This function implements a variance components estimator for multivariate and univariate randomeffects meta-analysis and meta-regression. It is meant to be used internally and not directly run by the users.

## Usage

```
mvmeta.vc(Xlist, ylist, Slist, nalist, k, m, p, nall, control, ...)
```
## Arguments

Assuming a meta-analysis or meta-regression based on  $m$  studies,  $k$  outcomes and  $p$  predictors:



<span id="page-39-0"></span>

#### <span id="page-40-0"></span>mvmeta.vc 41



## Details

The estimation involves kp fixed-effects coefficients and  $k(k + 1)/2$  random-effects parameters, corresponding to the lower triangular entries of the between-study (co)variance matrix.

The procedure is based on the estimate of the between-study (co)variance as the difference between the marginal (co)variance and the average within-study (co)variance. This in turn requires the estimate of the marginal (co)variance, obtained by the residuals of the fitted model. The procedure is iterative, with the current estimate of the between-study (co)variance plugged into a generalized least square (GLS) routine. Starting values are provided by a fixed-effects estimator (see mymeta.fixed). The algorithm is fast and generally converges with few iterations.

Similar versions of this estimator has been previously proposed. Berkey and collaborators (1998) simply called it GLS method, and a non-iterative approach was proposed by Ritz and collaborators (2008), referred to as MVEE3. A non-iterative version for univariate models is discussed in Sidik and Jonkman (2007). The results from Berkey and collaborators (1998) are reproduced in the example below.

In the original approach, the estimate of the marginal (co)variance is obtained from the sum of the residual components using a denominator equal to m-p. Following the development proposed by Kauermann and Carroll (2001) and Fay and Graubard (2001) in the context of sandwich (co)variance estimators, then discussed by Lu and collaborators (2007), an adjusted denominator can be computed as a quantity derived from the hat matrix. This method is expected to perform better in the presence of missing values and small data sets. This alternative adjustment is chosen by default by setting vc.adj=TRUE in the control argument.

The variance component estimator is not bounded to provide a positive semi-definite between-study (co)variance matrix, as shown in the simulation study by Liu and colleagues (2009). Here positive semi-definiteness is forced by setting the negative eigenvalues of the estimated matrix to zero at each iteration. Little is known about the impact of such constraint.

#### Value

This function returns an intermediate list object, whose components are then processed by mymeta. fit. Other components are added later through [mvmeta](#page-24-1) to finalize an object of class "mvmeta".

## **Note**

As stated earlier, this function is called internally by [mvmeta.fit](#page-24-2), and is not meant to be used directly. In particular, its code does not contain any check on the arguments provided, which are expected in specific formats. The function is however exported in the namespace and documented for completeness.

<span id="page-41-0"></span>The arguments above are prepared by mymeta. Fit from its arguments X, y and S. The list structure, although requiring more elaborate coding, is computationally more efficient, as it avoids the specification of sparse block-diagonal matrices, especially for meta-analysis involving a large number of studies.

Some parameters of the fitting procedures are determined by the control argument, with default set by [mvmeta.control](#page-29-1). No missing values are accepted in the fitting functions. See details on [missing values](#page-49-1).

## Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

Ritz J, Demidenko E, Spiegelman G (2008). Multivariate meta-analysis for data consortia, individual patient meta-analysis, and pooling projects. *Journal of Statistical Planning and Inference*. 139(7):1919–1933.

Berkey, CS, Hoaglin DC, et al. (1998). Meta-analysis of multiple outcomes by regression with random effects. *Statistics in Medicine*. 17(22):2537–2550.

Liu Q, Cook NR, Bergstrom A, Hsieh CC (2009). A two-stage hierarchical regression model for meta-analysis of epidemiologic nonlinear dose-response data. *Computational Statistics and Data Analysis*. 53(12):4157–4167

Sidik K, Jonkman JN (2007). A comparison of heterogeneity variance estimators in combining results of studies. *Statistics in Medicine*. 26(9):1964–81.

## See Also

See [mvmeta](#page-24-1) for the general usage of the functions. See [mvmeta.control](#page-29-1) to determine specific parameters of the fitting procedures. Use the triple colon operator  $(':::)$  to access the code of the internal functions, such as sumlist. See [mvmeta-package](#page-1-1) for an overview of the package and modelling framework.

#### Examples

```
# VC ESTIMATOR: UNIVARIATE MODEL
model <- mvmeta(PD~pubyear,S=berkey98[,5],data=berkey98,method="vc")
summary(model)
# VC ESTIMATOR: MULTIVARIATE MODEL
model <- mvmeta(cbind(PD,AL)~pubyear,S=berkey98[5:7],data=berkey98,method="vc")
summary(model)
# VC ESTIMATOR: NON-ITERATIVE VERSION
```
## <span id="page-42-0"></span>mvmetaCovStruct 43

```
model <- mvmeta(cbind(PD,AL)~pubyear,S=berkey98[5:7],data=berkey98,method="vc",
 control=list(maxiter=1))
summary(model)
# VARIANCE COMPONENTS ESTIMATOR: REPLICATE THE RESULTS IN BERKEY ET AL. (1998)
model <- mvmeta(cbind(PD,AL)~I(pubyear-1983),S=berkey98[5:7],
 data=berkey98,method="vc",control=list(vc.adj=FALSE))
summary(model)
```
<span id="page-42-1"></span>mvmetaCovStruct *Covariance Structures for mvmeta Models*

## Description

Alternative covariance structures for the between-study (co)variance matrix of random effects in multivariate meta-analysis or meta-regression, usually defined through the argument bscov of the function [mvmeta](#page-24-1).

## **Options**

Assuming a meta-analysis or meta-regression based on  $k$  outcomes:

- unstr: an unstructured form for a general positive-definite matrix. The matrix is represented by  $k(k+1)/2$  unrestricted parameters defined as the upper triangular entries of its Cholesky decomposition.
- diag: a diagonal positive-definite matrix. The matrix is represented by  $k$  unrestricted parameters defined as the logarithm of the diagonal values.
- id: a multiple of the identity positive-definite matrix. The matrix is represented by a single unrestricted parameter defined as the logarithm of the diagonal value.
- cs: a positive-definite matrix with compound symmetry structure. The matrix is represented by 2 unrestricted parameters defined as the logarithm of the diagonal value and the transformed correlation. The latter is parameterized so to obtain a correlation value between  $-1/(k-1)$ and 1, in order to ensure positive-definiteness.
- hcs: a positive-definite matrix with heterogeneous compound symmetry structure. The matrix is represented by  $k+1$  unrestricted parameters defined as the logarithm of the diagonal values and the transformed correlation. The latter is parameterized so to obtain a correlation value between  $-1/(k-1)$  and 1, in order to ensure positive-definiteness.
- ar1: a positive-definite matrix with autoregressive structure of first order. The matrix is represented by  $k+1$  unrestricted parameters defined as the logarithm of the diagonal value and the logistic transformed correlation. The latter is parameterized so to obtain a correlation value between -1 and 1.
- prop: a positive-definite matrix proportional to that provided by the user through the argument Psifix in the control list (see [mvmeta.control](#page-29-1)). The matrix is represented by 1 unrestricted parameter defined as the logarithm of the multiplier.
- <span id="page-43-0"></span>• cor: a positive-definite matrix with correlation structure provided by the user through the argument Psicor in the control list (see mymeta.control). The matrix is represented by  $k$ unrestricted parameters defined as the logarithm of the diagonal values.
- fixed: a known matrix provided by the user through the argument Psifix in the control list (see [mvmeta.control](#page-29-1)). The matrix is known and no parameters are needed to represent it.

#### Details

The structuring of the between-study (co)variance matrix of random effects is only available for models estimated through (restricted) maximum likelihood.

The unrestricted parameters defining the between-study matrix are estimated in the iterative optimization algorithm. The starting values are usually obtained by a iterative generalized least square algorithm (see the specific [likelihood functions](#page-20-1)). The algorithm computes an unstructured matrix. For structured forms, the starting values for variances and/or correlations are taken as the average of the related elements. Although rarely needed and not recommeded, the user can provided a starting value of the (co)variance matrix, from which the parameters are derived (see [mvmeta.control](#page-29-1)).

## **Note**

The choice of structures can affect the performance of the optimization procedure, determining forms of likelihood surfaces which induce convergence to local maxima. In particular, structures such as multiple of identity of proportional to a fixed matrix are based on strong assumptions and should be used with caution.

## Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Pinheiro JC and Bates DM (2000). *Mixed-Effects Models in S and S-PLUS*. New York, Springer Verlag.

#### See Also

See mymeta. See [lm](#page-0-0) or [glm](#page-0-0) for standard regression functions. See mymeta-package for an overview of this modelling framework.

## Examples

```
# UNSTRUCTURED AND STRUCTURED BETWEEN-STUDY (CO)VARIANCE
y <- as.matrix(fibrinogen[2:5])
S <- as.matrix(fibrinogen[6:15])
model <- mvmeta(y,S)
summary(model)
model$Psi
```
## <span id="page-44-0"></span>mvmetaObject 45

```
# DIAGONAL
model <- mvmeta(y,S,bscov="diag")
summary(model)
model$Psi
# HETEROGENEOUS COMPOUND SYMMETRY
model <- mvmeta(y,S,bscov="hcs")
summary(model)
model$Psi
# PROPORTIONAL
y <- as.matrix(smoking[11:13])
S <- as.matrix(smoking[14:19])
model <- mvmeta(y,S,bscov="prop",control=list(Psifix=diag(3)+1))
summary(model)
model$Psi
# CORRELATION
model <- mvmeta(y,S,bscov="cor",control=list(Psicor=0.2))
summary(model)
model$Psi
```
<span id="page-44-1"></span>mvmetaObject *mvmeta Objects*

## **Description**

An object returned by the [mvmeta](#page-24-1) function, inheriting from class "mvmeta", and representing a fitted univariate or multivariate meta-analytical model.

## Value

Objects of class "mvmeta" are lists with defined components. Dimensions of such components may refer to  $k$  outcome parameters,  $p$  predictors and  $m$  studies used for fitting the model (the latter can be different from those originally selected due to missing). The following components needs to be included in a legitimate mvmeta object:



<span id="page-45-0"></span>

#### <span id="page-46-0"></span>mvmetaObject 47

#### Methods

A number of methods functions are available for mvmeta objects, most of them common to other regression functions.

Specifically-written method functions are defined for [predict](#page-52-1) (standard predictions) and [blup](#page-8-1) (best linear unbiased predictions). The method function [simulate](#page-47-2) produces simulated outcomes from a fitted model, while [qtest](#page-55-1) performs the Cochran Q test for heterogeneity. Other methods have been produced for [summary](#page-58-1), [logLik](#page-19-1), [coef](#page-10-1), and [vcov](#page-10-2).

Specific methods are also available for model. frame and [model.matrix](#page-23-2). In particular, the former produces the model frame (a data frame with special attributes storing the variables used for fitting) with the additional class "data.frame.mvmeta". Methods [na.omit](#page-49-1) and [na.exclude](#page-49-2) for this class are useful for the handling of missing values in mvmeta objects.

Printing functions for the objects of classes defined above are also provided. anova methods for performing tests in mvmeta objects are in development.

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 "mvmeta", such as [fitted](#page-0-0), [residuals](#page-0-0), [AIC](#page-0-0), [BIC](#page-0-0) and [update](#page-0-0), among others.

## Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

#### References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

## See Also

See mymeta. See [lm](#page-0-0) or [glm](#page-0-0) for standard regression functions. See mymeta-package for an overview of this modelling framework.

#### Examples

```
# RUN THE MODEL
model <- mvmeta(cbind(PD,AL)~pubyear,S=berkey98[5:7],data=berkey98)
```

```
# INSPECT THE OBJECT
names(model)
```
# LABELS model\$lab

# FORMULA model\$formula

## <span id="page-47-0"></span># CONVERGED? model\$converged

## <span id="page-47-1"></span>mvmetaSim *Simulating Responses for mvmeta Models*

## <span id="page-47-2"></span>Description

These functions simulate sets of multivariate or univariate responses for a group of studies, in terms of their mean (expected) values and within and between-study (co)variances. These sets of outcomes can be used in meta-analytical models for simulation purposes.

## Usage

mvmetaSim(y, S, Psi, sd, cor, nsim=1, seed=NULL, posdeftol) ## S3 method for class 'mvmeta' simulate(object, nsim=1, seed=NULL, ...)

## Arguments

In order to simulate  $k$  outcomes for  $m$  studies:



#### <span id="page-48-0"></span>mvmetaSim 49

#### Details

The set(s) of responses can be simulated either from a fitted model, using the method function simulate for objects of class "mvmeta", or directly through the function mvmetaSim. In the former case, the fitted values from the model are used as mean (expected) outcomes, together with the within and estimated between-study (co)variance. In the latter option, this information need to be provided by the user in the correct dimensions and forms.

In mvmetaSim, the between-study (co)variance matrix can be inputted directly through Psi, or given in the form  $DRD$ , with a diagonal matrix  $D$  of standard deviations and correlation matrix  $R$ . These values are provided through sd and cor. See [inputcov](#page-15-1) for details.

The functions simulate the responses for each study separately from a marginal multivariate normal distribution with mean equal to the expected values and (co)variance equal to the sum of the within and between-study components. The computation is identical to that implemented in the function mvrnorm of the package MASS, involving a eigen decomposition of the marginal (co)variance matrix. Numerical negative definiteness is allowed with a tolerance specified by posdeftol, and positive semi-definiteness is then forced by truncating the eigenvalues at zero.

## Value

If nsim=1, a matrix or vector of simulated k outcomes for the  $m$  studies. If more simulation sets are required (nsim higher than 1), a list of matrices or vectors.

## Note

Studies with missing values in the fitted values or in the components of the within (co)variances are excluded by simulate. Missing values are instead not accepted in metaSim.

## Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

## See Also

See [simulate](#page-0-0) for the general method function. See [inputcov](#page-15-1) for inputting correlations. See [mvmeta-package](#page-1-1) for an overview of the package and modelling framework.

## Examples

```
# RUN A MODEL
model <- mvmeta(cbind(PD,AL)~pubyear,S=berkey98[5:7],data=berkey98)
# SIMULATE A NEW SET OF OUTCOMES
simulate(model)
```
# SIMULATE FROM SCRATCH: 3 OUTCOMES, 8 STUDIES

```
(y \le - matrix(0, 8, 3)(S <- inputcov(matrix(runif(8*3,0.1,2),8,3,dimnames=list(NULL,
 c("V1","V2","V3"))),cor=c(0,0.5,0.7)))
(Psi <- inputcov(1:3,cor=0.3))
mvmetaSim(y,S,Psi)
# ALTERNATIVELY, DEFINE Psi THROUGH STANDARD DEVIATIONS AND CORRELATION 0.2
mvmetaSim(y,S,sd=1:3,cor=0.3)
# 2 SIMULATION SETS
mvmetaSim(y,S,Psi,nsim=2)
```
<span id="page-49-1"></span>na.omit.data.frame.mvmeta

*Handling Missing Values in mvmeta Models*

## <span id="page-49-2"></span>**Description**

These method functions exclude rows corresponding to studies with invalid missing pattern from model frames of class "data.frame.mvmeta". This guarantees the correct handling of missing values while fitting multivariate and univariate meta-analytical models.

#### Usage

```
## S3 method for class 'data.frame.mvmeta'
na.omit(object, ...)
## S3 method for class 'data.frame.mvmeta'
```
na.exclude(object, ...)

#### Arguments



#### Details

A model frame of class "data.frame.mvmeta" is produced by [mvmeta](#page-24-1). A call to na.omit or na.exclude removes from the model frame the rows corresponding to studies with invalid missing pattern. In addition, a na.action attribute is added to the model frame, namely a numeric vector corresponding to the removed rows and class "omit" or "exclude", respectively. This information is used by [naresid](#page-0-0) and [napredict](#page-0-0) to deal with missing values in functions such as [fitted](#page-0-0), [residuals](#page-0-0), [predict](#page-52-1) and [blup](#page-8-1), among others.

The definition of missing, identifying an invalid missing pattern, is different in multivariate metaanalytical models performed through mymeta if compared to other regression functions such as  $\text{lm}$  $\text{lm}$  $\text{lm}$ or [glm](#page-0-0). Specifically, while a study is removed if at least an observation for one predictor is missing, partially missing outcomes do not prevent the study to contribute to estimation (see [mvmeta](#page-24-1)). Specific methods na.omit and na.exclude for class "data.frame.mvmeta" allow this different definition.

<span id="page-49-0"></span>

## <span id="page-50-0"></span>Value

These functions returns the model frame object with rows corresponding to studies with invalid missing pattern being removed. They also add the related na.action attribute as explained above.

## Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## See Also

See [na.action](#page-0-0), [naresid](#page-0-0) and [napredict](#page-0-0). See [model.frame](#page-23-1). See [mvmeta-package](#page-1-1) for an overview of the package and modelling framework.

## Examples

```
# INPUT MISSING VALUES IN PREDICTOR AND ONE RESPONSE
data <- berkey98
data[2,1] <- data[4,3] <- NA
data
# RUN THE MODEL
model <- mvmeta(cbind(PD, AL)~pubyear, S=data[5:7], data=data, method="ml")
# SUMMARIZE: NOTE THE NUMBER OF STUDIES AND OBSERVATIONS
summary(model)
df.residual(model)
# EXTRACT THE MODEL FRAME WITH na.pass
model.frame(model,na.action="na.pass")
# EXTRACT THE MODEL FRAME WITH na.omit (DEFAULT)
model.frame(model,na.action="na.omit")
# COMPARE WITH DEFAULT METHOD FOR na.omit
frame <- model.frame(model,na.action="na.pass")
na.omit(frame)
class(frame)
class(frame) <- "data.frame"
na.omit(frame)
# WITH na.exclude
residuals(model)
```
<span id="page-50-1"></span>residuals(update(model,na.action="na.exclude"))

## <span id="page-51-0"></span>Description

The dataset includes studies providing evidence about whether the presence of mutant p53 tumour suppressor gene is a prognostic factor for patients presenting with squamous cell carcinoma arising from the oropharynx cavity. Unadjusted estimates of log hazard ratios of mutant p53 to normal p53 for disease-free and overall survival, together with the associated variances, are collected from 6 observational studies.

## Usage

p53

## Format

A data frame with 6 observations on the following 5 variables:

- study: study ID.
- y1, V1: estimate and associated variance of the log hazard ratio for disease-free survival.
- y2, V2: estimate and associated variance of the log hazard ratio for overall survival.

#### Details

Only 3 studies provide estimates for disease-free survival. The within-study correlations are not reported in the original studies but are expected to be highly positively correlated. The original data are described in Tandon and colleagues (2010) and used as an example in Jackson and colleagues (2011).

## **Note**

The data provide an example of application of multivariate meta-analysis when the within-study correlations are not known. These correlations can be inputted directly in the [mvmeta](#page-24-1) function through the control argument. See mymeta.control for details.

#### Source

Jackson D, Riley R, White IR (2011). Multivariate meta-analysis: Potential and promise. *Statistics in Medicine*. 30(20);2481–2498.

Tandon S, Tudur-Smith C, Riley RD, et al. (2010). A systematic review of p53 as a prognostic factor of survival in squamous cell carcinoma of the four main anatomical subsites of the head and neck. *Cancer Epidemiology, Biomarkers and Prevention*. 19(2):574–587.

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

## Examples

```
### REPRODUCE THE RESULTS OF EXAMPLE 3 IN JACKSON ET AL. (2011)
# INSPECT THE DATA
p53
```

```
# REML MODEL WITH INPUTTED CORRELATION EQUAL TO 0.95
model <- mvmeta(cbind(y1,y2),cbind(V1,V2),data=p53,control=list(Scor=0.95))
print(summary(model),digits=2)
```
<span id="page-52-1"></span>predict.mvmeta *Predicted Values from mvmeta Models*

## Description

This method function computes predictions from fitted univariate or multivariate meta-analytical models represented in objects of class "mvmeta", optionally for a new set of predictor values in meta-regression models. Predictions are optionally accompanied by standard errors, confidence intervals or the entire (co)variance matrix of the predicted outcomes.

## Usage

```
## S3 method for class 'mvmeta'
predict(object, newdata, se=FALSE, ci=FALSE, vcov=FALSE,
  interval=c("confidence","prediction"), ci.level=0.95,
  format=c("matrix","list"), aggregate=c("stat","y"), na.action=na.pass, ...)
```
## Arguments



## <span id="page-53-0"></span>Details

The method function predict produces predicted values from mvmeta objects, obtained by evaluating the original call to [mvmeta](#page-24-1) in the frame newdata. For both fixed and random-effects models, estimated predictions are only based on the fixed part of the model, ignoring study-specific deviations, differently from **[blup](#page-8-1)**.

For random-effects models, if interval="confidence" (the default), standard errors, confidence intervals and (co)variance matrix of the predicted values are computed only using the estimated (co)variance matrix of the fixed-effects coefficients. If interval="prediction", the estimated between-study (co)variance matrix, stored in the Psi component of mvmeta objects, is also added. In this case, interpretation of the uncertainty reflects that of a new single study sampled from the same population.

If newdata is omitted, the predictions are based on the data used for the fit. In that case how to handle predictions for studies removed from estimation due to invalid missing pattern is determined by the na.action argument used in mymeta to produce object. If na.action=na.omit, studies excluded from estimation will not appear, whereas if na.action=na.exclude they will appear, with values set to NA for all the outcomes. This step is performed by [napredict](#page-0-0). See Notes.

## Value

The results may be aggregated in matrices (the default), or returned as lists, depending on the argument format. For multivariate models, the aggregation is ruled by the argument aggregate, and the results may be grouped by statistic or by outcome. If vcov=TRUE, lists are always returned.

#### **Note**

The definition of missing in model frames used for estimation in [mvmeta](#page-24-1) is different than that commonly adopted in other regression models such as  $\text{lm}$  $\text{lm}$  $\text{lm}$  or [glm](#page-0-0). See info on [missing values](#page-49-1) in [mvmeta](#page-24-1).

#### Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

#### See Also

See [blup](#page-8-1) for best linear unbiased predictions. See the default method [predict](#page-0-0). See [mvmeta-package](#page-1-1) for an overview of the package and modelling framework.

## <span id="page-54-0"></span>qtest 55

## Examples

```
# RUN THE MODEL
model <- mvmeta(cbind(PD,AL)~pubyear,S=berkey98[5:7],data=berkey98)
# PREDICTED FROM YEAR 1985 TO 1987, WITH LABELS
newdata <- data.frame(pubyear=1985:1987,row.names=1985:1987)
# AVERAGED OUTOCOMES AND SE
predict(model,newdata,se=TRUE)
# SAME AS ABOVE, AGGREGATED BY OUTCOME
predict(model,newdata,se=TRUE,aggregate="y")
# SAME AS ABOVE, WITH PREDICTION INTERVALS
predict(model,newdata,se=TRUE,aggregate="y",interval="prediction")
# WITH VCOV, FORCED TO A LIST
predict(model,newdata,se=TRUE,vcov=TRUE,aggregate="y")
```
<span id="page-54-1"></span>qtest *Cochran Q Test of Heterogeneity*

## Description

This is a generic function to perform a Cochran Q test of (residual) heterogeneity. The function invokes particular [methods](#page-0-0) which depend on the [class](#page-0-0) of the first argument. Currently, specific methods exist for several meta-analytical models in various packages: [qtest.mixmeta](#page-0-0), [qtest.mvmeta](#page-55-1), and [qtest.dosresmeta](#page-0-0).

## Usage

qtest(object, ...)

#### Arguments



## Details

The test assesses the null hypothesis that the variability in the distribution of the outcomes is explained only in terms of within-unit estimation errors. This corresponds to a test on the hypotesis that there is no variation attributable to random-effects terms.

## Value

Returned values depend on the specific class. Usually, the results of the test.

## <span id="page-55-0"></span>Author(s)

Antonio Gasparrini << antonio.gasparrini@lshtm.ac.uk>>

#### References

Cochran WG (1950). The comparison of percentages in matched samples". *Biometrika*. 37(3/4):256– 266.

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

## See Also

Specific methods for various classes: [qtest.mixmeta](#page-0-0), [qtest.mvmeta](#page-55-1), and [qtest.dosresmeta](#page-0-0).

<span id="page-55-1"></span>qtest.mvmeta *Cochran Q Test of Heterogeneity for mvmeta Models*

## Description

This method function performs a Cochran Q test of (residual) heterogeneity on fitted univariate or multivariate meta-analytical models represented in objects of class "mvmeta".

#### Usage

## S3 method for class 'mvmeta' qtest(object, ...) ## S3 method for class 'qtest.mvmeta'

print(x, digits=3, ...)

#### Arguments



#### Details

In multivariate 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 S of mvmeta 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. Tests for single outcome parameters, comparable to estimates from multiple univariate meta-analysis, are also reported. This test reduces to the standard Q test in univariate models.

The function compute the statistics by actually fitting the related fixed-effects model, re-evaluating the call of the model with method changed to "fixed".

## <span id="page-56-0"></span>qtest.mvmeta 57

## Value

A list object of class "qtest.mvmeta" with the following components:



As usual, the print method function for class "qtest.mvmeta" does not return any value.

## Note

Tests on single outcome parameters are performed by extracting the related estimates and variances, but they do not account for the correlation between them, which nevertheless has been considered in estimation. These tests are not therefore comparable with those performed by running a univariate model on each outcome parameter.

## Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

Berkey, CS, Hoaglin DC, et al. (1998). Meta-analysis of multiple outcomes by regression with random effects. *Statistics in Medicine*. 17(22):2537–2550.

Ritz J, Demidenkob E, Spiegelman G (2008). Multivariate meta-analysis for data consortia, individual patient meta-analysis, and pooling projects. *Journal of Statistical Planning and Inference*. 139(7):1919–1933.

#### See Also

See [qtest](#page-54-1) for the generic method function. See [mvmeta-package](#page-1-1) for an overview of the package and modelling framework.

<span id="page-57-0"></span>58 smoking

## Examples

```
# RUN THE MODEL
model <- mvmeta(cbind(PD,AL)~1,S=berkey98[5:7],data=berkey98)
# MULTIVARIATE COCHRAN Q TEST FOR HETEROGENEITY
test <- qtest(model)
print(test,digits=2)
unclass(test)
```
<span id="page-57-1"></span>

smoking *Meta-Analysis of Interventions to Promote Smoking Cessation*

## **Description**

The dataset contains the results of 24 trials comparing four alternative interventions to promote smoking cessation. The trials have different designs, comparing two or three different interventions. The data consist of the number of successes out of the total participants, and the estimated log-odds ratio for arms B, C, and D relative to arm A, as well as the (co)variance matrix of these three estimates.

## Usage

smoking

## Format

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

- study: study ID.
- design: design of the trial, reporting the interventions being compared.
- dA, dB, dC, dD: number of successes for each intervention.
- nA, nB, nC, nD: number of participants for each intervention.
- yB, yC, yD: estimated log-odds ratios for interventions B, C and D versus intervention A.
- SBB, SBC, SBD, SCC, SCD, SDD: variances and co-variances of the estimated log-odds ratios for interventions B, C and D versus intervention A. The order corresponds to the lower triangular elements of the (co)variance matrix taken by column.

## Details

Intervention A is chosen as the reference category. Trials without an arm A (trials 2 and 21-24) are augmented with an arm A with 0.01 individuals and 0.001 successes. Trials containing zero cells (trials 9 and 20) have 1 individual with 0.5 successes added to each intervention. Details on the data augmentation and estimation of (co)variances of the log-odds ratios are provided by White (2011).

#### <span id="page-58-0"></span>**Note**

The data provide an example of application of network meta-analysis, also referred to as indirect mixed-treatment comparison. Additional information using examples based on these data are provided by Lu and Ades (2006), White (2011) and Higgins and colleagues (2012).

## Source

Lu G, Ades AE (2006). Assessing evidence inconsistency in mixed treatment comparisons. *Journal of the American Statistical Association*. 101:447–459.

Higgins JPT, Jackson D, Barrett JK et al (2012). Consistency and inconsistency in network metaanalysis: concepts and models for multi-arm studies. *Research Synthesis Methods*. 3(2):98–110.

White IR (2011). Multivariate random-effects meta-regression. *The Stata Journal*. 11:255–270.

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

## Examples

```
### REPRODUCE THE RESULTS IN WHITE (2011)
# INSPECT THE DATA
head(smoking)
names(smoking)
# UNSTRUCTURED BETWEEN-STUDY (CO)VARIANCE
y <- as.matrix(smoking[11:13])
S <- as.matrix(smoking[14:19])
model <- mvmeta(y,S)
summary(model)
# STRUCTURED BETWEEN-STUDY (CO)VARIANCE (PROPORTIONAL)
model <- mvmeta(y,S,bscov="prop",control=list(Psifix=diag(3)+1))
summary(model)
```
# SEE help(mvmetaCovStruct) for additional info and examples

<span id="page-58-1"></span>summary.mvmeta *Summarizing mvmeta Models*

#### Description

Print and summary method functions for fitted univariate or multivariate meta-analytical models represented in objects of class "mvmeta".

## Usage

```
## S3 method for class 'mvmeta'
summary(object, ci.level=0.95, ...)
## S3 method for class 'summary.mvmeta'
print(x, \text{ digits=4}, \ldots)## S3 method for class 'mvmeta'
print(x, \text{ digits=4}, \ldots)
```
## Arguments



## Details

The print method function for class "mvmeta" only returns basic information on 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.mvmeta". The print method function for this class shows additional information, such as tables reporting the estimates for the fixed and random-effects parts of the model, Cochran Q test for heterogeneity and  $I^2$ .

#### Value

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



<span id="page-59-0"></span>

#### <span id="page-60-0"></span>vechMat 61



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

## Author(s)

Antonio Gasparrini, <antonio.gasparrini@lshtm.ac.uk>

## References

Sera F, Armstrong B, Blangiardo M, Gasparrini A (2019). An extended mixed-effects framework for meta-analysis.*Statistics in Medicine*. 2019;38(29):5429-5444. [Freely available [here](http://www.ag-myresearch.com/2019_sera_statmed.html)].

Gasparrini A, Armstrong B, Kenward MG (2012). Multivariate meta-analysis for non-linear and other multi-parameter associations. *Statistics in Medicine*. 31(29):3821–3839. [Freely available [here](http://www.ag-myresearch.com/2012_gasparrini_statmed.html)].

## See Also

See [mvmeta](#page-24-1) and [mvmetaObject](#page-44-1).

## Examples

```
# RUN THE MODEL
model <- mvmeta(cbind(PD,AL)~pubyear,S=berkey98[5:7],data=berkey98)
# SIMPLE PRINT
model
# DEFINE DIGITS
print(model,digit=2)
```
# SUMMARY WITH 80TH CONFIDENCE INTERVALS summary(model,ci.level=0.80)

<span id="page-60-1"></span>vechMat *Vectorization and Expansion of Symmetric Matrices*

## <span id="page-60-2"></span>Description

The function vechMat transforms a symmetric matrix in a vector containing its lower triangular elements, taken by column. The function xpndMat reverses this transformation.

## Usage

```
vechMat(mat, diag=TRUE)
```
xpndMat(vech)

## Arguments



## Value

A vector for vechMat, a symmetric matrix for xnpdMat.

## Note

These functions are imported from the package mixmeta.

## Author(s)

Antonio Gasparrini << antonio.gasparrini@lshtm.ac.uk>>

## See Also

See functions vech and xpnd in package MCMCpack.

## Examples

```
# GENERATE A POSITIVE-DEFINITE MATRIX, VECTORIZE IT AND THEN RE-EXPAND
(M <- crossprod(matrix(rnorm(9),3)))
(v <- vechMat(M))
xpndMat(v)
# EXTRACT VECTORIZED S, EXPAND TO A LIST, AND RE-VECTORIZE
(S <- as.matrix(berkey98[5:7]))
```

```
(Slist <- lapply(seq(nrow(S)), function(i) xpndMat(S[i,])))
t(sapply(Slist,vechMat))
```
# <span id="page-62-0"></span>Index

∗ datasets berkey98, [6](#page-5-0) fibrinogen, [12](#page-11-0) hsls, [13](#page-12-0) hyp, [15](#page-14-0) p53, [51](#page-50-0) smoking, [58](#page-57-0) ∗ htest qtest, [55](#page-54-0) qtest.mvmeta, [56](#page-55-0) ∗ manip inputcov, [16](#page-15-0) inputna, [18](#page-17-0) na.omit.data.frame.mvmeta, [50](#page-49-0) vechMat, [61](#page-60-0) ∗ methods blup, [8](#page-7-0) blup.mvmeta, [9](#page-8-0) coef.mvmeta, [11](#page-10-0) logLik.mvmeta, [20](#page-19-0) model.frame.mvmeta, [24](#page-23-0) mvmetaSim, [48](#page-47-0) na.omit.data.frame.mvmeta, [50](#page-49-0) predict.mvmeta, [53](#page-52-0) qtest, [55](#page-54-0) qtest.mvmeta, [56](#page-55-0) summary.mvmeta, [59](#page-58-0) ∗ models blup, [8](#page-7-0) blup.mvmeta, [9](#page-8-0) coef.mvmeta, [11](#page-10-0) logLik.mvmeta, [20](#page-19-0) mlprof.fn, [21](#page-20-0) model.frame.mvmeta, [24](#page-23-0) mvmeta, [25](#page-24-0) mvmeta.control, [30](#page-29-0) mymeta.fixed.[33](#page-32-0) mvmeta.ml, [35](#page-34-0) mvmeta.mm, [38](#page-37-0)

mvmeta.vc, [40](#page-39-0) mvmetaCovStruct, [43](#page-42-0) mvmetaObject, [45](#page-44-0) mvmetaSim, [48](#page-47-0) na.omit.data.frame.mvmeta, [50](#page-49-0) predict.mvmeta, [53](#page-52-0) qtest.mvmeta, [56](#page-55-0) summary.mvmeta, [59](#page-58-0) ∗ multivariate blup, [8](#page-7-0) blup.mvmeta, [9](#page-8-0) coef.mvmeta, [11](#page-10-0) logLik.mvmeta, [20](#page-19-0) mlprof.fn, [21](#page-20-0) model.frame.mvmeta, [24](#page-23-0) mvmeta, [25](#page-24-0) mvmeta.control, [30](#page-29-0) mvmeta.fixed, [33](#page-32-0) mvmeta.ml, [35](#page-34-0) mvmeta.mm, [38](#page-37-0) mvmeta.vc, [40](#page-39-0) mvmetaCovStruct, [43](#page-42-0) mvmetaObject, [45](#page-44-0) mvmetaSim, [48](#page-47-0) na.omit.data.frame.mvmeta, [50](#page-49-0) predict.mvmeta, [53](#page-52-0) qtest.mvmeta, [56](#page-55-0) summary.mvmeta, [59](#page-58-0) ∗ package mvmeta-package, [2](#page-1-0) ∗ regression blup, [8](#page-7-0) blup.mvmeta, [9](#page-8-0) coef.mvmeta, [11](#page-10-0) logLik.mvmeta, [20](#page-19-0) mlprof.fn, [21](#page-20-0) model.frame.mvmeta, [24](#page-23-0) mvmeta, [25](#page-24-0) mvmeta.control, [30](#page-29-0)

```
33
    mvmeta.ml, 35
    mvmeta.mm, 38
    mvmeta.vc, 40
    mvmetaCovStruct, 43
    mvmetaObject, 45
    mvmetaSim, 48
    na.omit.data.frame.mvmeta, 50
    predict.mvmeta, 53
    qtest.mvmeta, 56
    59
AIC, 4, 20, 47, 60
as.data.frame, 26
berkey98, 4, 6
BIC, 4, 20, 47, 60
blup, 4, 8, 47, 50, 54
blup.dosresmeta, 8
blup.mixmeta, 8
blup.mvmeta, 8, 9
blup.rma.uni, 8
chol, 23
class, 8, 55
coef, 47
coef.mvmeta, 11
control, 18
fibrinogen, 4, 12
fitted, 4, 47, 50
formula, 25, 27, 46
glm, 10, 26–28, 44, 47, 50, 54
glm.control, 32
hsls, 4, 13
hyp, 4, 15
inputcov, 4, 16, 19, 27, 31, 48, 49
inputna, 4, 18, 31
iter.igls, 31
iter.igls (mlprof.fn), 21
lm, 10, 24, 26–28, 44, 47, 50, 54
logLik, 4, 21, 47
logLik.mvmeta, 20
```
methods, *[55](#page-54-0)* mlprof.fn, [21](#page-20-0) mlprof.gr *(*mlprof.fn*)*, [21](#page-20-0) model.frame, *[4](#page-3-0)*, *[24](#page-23-0)[–27](#page-26-0)*, *[46,](#page-45-0) [47](#page-46-0)*, *[51](#page-50-0)* model.frame.mvmeta, [24](#page-23-0) model.matrix, *[4](#page-3-0)*, *[25,](#page-24-0) [26](#page-25-0)*, *[46,](#page-45-0) [47](#page-46-0)* model.matrix.mvmeta *(*model.frame.mvmeta*)*, [24](#page-23-0) model.offset, *[26](#page-25-0)* mvmeta, *[4](#page-3-0)*, *[10](#page-9-0)*, *[17,](#page-16-0) [18](#page-17-0)*, *[24](#page-23-0)*, [25,](#page-24-0) *[31,](#page-30-0) [32](#page-31-0)*, *[34](#page-33-0)*, *[36,](#page-35-0) [37](#page-36-0)*, *[39](#page-38-0)[–47](#page-46-0)*, *[50](#page-49-0)*, *[52](#page-51-0)*, *[54](#page-53-0)*, *[60,](#page-59-0) [61](#page-60-0)* mvmeta-package, [2](#page-1-0) mvmeta.control, *[4](#page-3-0)*, *[17](#page-16-0)*, *[22,](#page-21-0) [23](#page-22-0)*, *[26,](#page-25-0) [27](#page-26-0)*, [30,](#page-29-0) *[33](#page-32-0)[–44](#page-43-0)*, *[46](#page-45-0)*, *[52](#page-51-0)* mvmeta.fit, *[4](#page-3-0)*, *[23](#page-22-0)*, *[30](#page-29-0)*, *[32](#page-31-0)*, *[34](#page-33-0)*, *[36](#page-35-0)*, *[39](#page-38-0)*, *[41,](#page-40-0) [42](#page-41-0)* mvmeta.fixed, *[4](#page-3-0)*, [33,](#page-32-0) *[41](#page-40-0)* mvmeta.ml, *[4](#page-3-0)*, *[22,](#page-21-0) [23](#page-22-0)*, *[33](#page-32-0)*, [35](#page-34-0) mvmeta.mm, *[4](#page-3-0)*, [38](#page-37-0) mvmeta.reml, *[4](#page-3-0)*, *[22,](#page-21-0) [23](#page-22-0)*, *[33](#page-32-0)* mvmeta.reml *(*mvmeta.ml*)*, [35](#page-34-0) mvmeta.vc, *[4](#page-3-0)*, *[31](#page-30-0)*, [40](#page-39-0) mvmetaCovStruct, *[23](#page-22-0)*, [43](#page-42-0) mvmetaObject, *[4](#page-3-0)*, *[9](#page-8-0)*, *[27](#page-26-0)*, [45,](#page-44-0) *[60,](#page-59-0) [61](#page-60-0)* mvmetaSim, *[4](#page-3-0)*, [48](#page-47-0)

na.action, *[51](#page-50-0)* na.exclude, *[4](#page-3-0)*, *[24](#page-23-0)[–26](#page-25-0)*, *[47](#page-46-0)* na.exclude.data.frame.mvmeta *(*na.omit.data.frame.mvmeta*)*, [50](#page-49-0) na.omit, *[4](#page-3-0)*, *[24](#page-23-0)[–26](#page-25-0)*, *[47](#page-46-0)* na.omit.data.frame.mvmeta, [50](#page-49-0) napredict, *[10](#page-9-0)*, *[50,](#page-49-0) [51](#page-50-0)*, *[54](#page-53-0)* naresid, *[50,](#page-49-0) [51](#page-50-0)*

offset, *[26](#page-25-0)* optim, *[22,](#page-21-0) [23](#page-22-0)*, *[31,](#page-30-0) [32](#page-31-0)* options, *[26](#page-25-0)*

p53, *[4](#page-3-0)*, [51](#page-50-0) predict, *[4](#page-3-0)*, *[9,](#page-8-0) [10](#page-9-0)*, *[47](#page-46-0)*, *[50](#page-49-0)*, *[54](#page-53-0)* predict.mvmeta, [53](#page-52-0) print.mvmeta *(*summary.mvmeta*)*, [59](#page-58-0) print.qtest.mvmeta *(*qtest.mvmeta*)*, [56](#page-55-0) print.summary.mvmeta *(*summary.mvmeta*)*, [59](#page-58-0)

qr, *[23](#page-22-0)* qtest, *[4](#page-3-0)*, *[47](#page-46-0)*, [55,](#page-54-0) *[57](#page-56-0)*, *[61](#page-60-0)* qtest.dosresmeta, *[55,](#page-54-0) [56](#page-55-0)* qtest.mixmeta, *[55,](#page-54-0) [56](#page-55-0)* qtest.mvmeta, *[4](#page-3-0)*, *[55,](#page-54-0) [56](#page-55-0)*, [56](#page-55-0)

#### INDEX 65

remlprof.fn *(*mlprof.fn *)* , [21](#page-20-0) remlprof.gr *(*mlprof.fn *)* , [21](#page-20-0) residuals , *[4](#page-3-0)* , *[47](#page-46-0)* , *[50](#page-49-0)* simulate , *[4](#page-3-0)* , *[47](#page-46-0)* , *[49](#page-48-0)* simulate.mvmeta (mvmetaSim), [48](#page-47-0) smoking , *[4](#page-3-0)* , [58](#page-57-0) structure , *[22](#page-21-0)* summary , *[4](#page-3-0)* , *[47](#page-46-0)* summary.mvmeta, [59](#page-58-0)

terms , *[46](#page-45-0)*

update , *[47](#page-46-0)*

vcov , *[47](#page-46-0)* , *[60](#page-59-0)* vcov.mvmeta *(*coef.mvmeta *)* , [11](#page-10-0) vechMat , *[17](#page-16-0)* , [61](#page-60-0)

xpndMat , *[17](#page-16-0)* , *[27](#page-26-0)* , *[48](#page-47-0)* xpndMat *(*vechMat *)* , [61](#page-60-0)