

# Computing Within-Study Covariances, Data Visualization and Missing Data Solutions for Multivariate Meta-Analysis with metavcov

Min Lu<sup>1,\*</sup>

<sup>1</sup>Division of Biostatistics, Department of Public Health Sciences, Miller School of Medicine, University of Miami, Miami, FL, United States

Correspondence\*: Corresponding Author m.lu6@umiami.edu

## **ABSTRACT**

- 3 Multivariate meta-analysis (MMA) is a powerful statistical technique that can provide more reliable and informative results than traditional univariate meta-analysis, which allows for comparisons 4 across outcomes with increased statistical power. However, implementing appropriate statistical methods for MMA can be challenging due to the requirement of various specific tasks in data 6 7 preparation. The metavcov package aims for model preparation, data visualization and missing data solutions to provide tools for different methods that cannot be found in accessible software. It provides sufficient constructs for estimating coefficients from other well-established packages. 9 For model preparation, users can compute both effect sizes of various types and their variance-10 covariance matrices, including correlation coefficients, standardized mean difference, mean 11 difference, log odds ratio, log risk ratio, and risk difference. The package provides a tool to plot the 12 confidence intervals for the primary studies and the overall estimates. When specific effect sizes 13 are missing, single imputation is available in the model preparation stage; a multiple imputation 14 method is also available for pooling the results in a statistically principled manner from models of 15 users' choice. The package is demonstrated in two real data applications and a simulation study 16 17 to assess methods for handling missing data.
- 18 Keywords: multivariate meta-analysis, effect sizes, variance-covariance matrix, multiple imputation, confidence intervals

#### 1 INTRODUCTION

Multivariate meta-analysis (MMA) is a statistical technique of combining multiple effect sizes, either 20 of the same type or different types, from different studies to produce one overall result. It allows for within-study dependence among effect sizes caused by the fact that multiple outcomes are obtained from 21 the same samples in the primary studies. This dependence could increase the Type I error rate and lead to 22 23 inaccurate estimates of study effects (Riley, 2009; Nam et al., 2003; Jackson et al., 2011; Becker, 2000). 24 Although there are many R packages available for univariate meta-analysis, resources for MMA are limited in terms of data preparation and visualization (Michael Dewey, 2021). There are available R packages 25 26 (see Table 1) designed for fitting MMA models, but they assume that the within-study variance-covariance 27 matrices of the effect sizes from all studies are pre-computed by the users. Therefore, these packages may be unattractive in practice. For example, in some MMA application papers, univariate meta-analysis is still 28

46

47

48

49

50

51

52

53

54

55

56

57 58

59

60

61

62

63

64

65

66

67

68

69 70

- 29 adopted even though several effect sizes are extracted from the same study (Watters et al., 2021; Sebri 30 et al., 2021). Conducting statistically principled MMA confronts challenges, including:
- 31 1. It is challenging to compute the covariances among effect sizes for non-statisticians;
- 32 2. It lacks data visualization tools;
- 33 3. It suffers greatly from the missing data problem.

The availability of generalizable, user-friendly software packages facilitates the incorporation of MMA 34 into various fields of science. The package metavcov aims to provide useful tools for conducting MMA 35 in R (R Core Team, 2016) with examples of how it can provide aid for easy, efficient and accurate computer 36 programs (Lu, 2017). It is not designed to replace a parameter estimation package for MMA, such as 37 mixmeta and metaSEM (Gasparrini, 2019; Cheung, 2021; Aloe et al., 2014a,b), but to provide additional 38 specialized tools. It was initially released in 2017 for computing variance-covariance matrices of effect sizes 39 and has attracted growing downloads as shown in Figure 1. Its new version addresses all the above three 40 points. For point 1, formulas and references are provided in the next section for computing covariances. 41 Tutorials are given to guide users to use R functions that can accommodate different types of effect sizes 42 and their variance-covariance matrices for preparing desired input arguments for packages mixmeta and 43 metaSEM as examples. Note that since the diagonal elements of the variance-covariance matrix are the 44 variances of the estimated effect sizes, this package can also be used for preparing univariate meta-analysis. 45

For point 2, the metavcov package introduces a function for confidence interval plots. Although forest plots are used for displaying effect sizes from all studies and their overall estimator in the univariate meta-analysis (Schwarzer et al., 2007; Boyles et al., 2011; Rücker and Schwarzer, 2021; Sedgwick, 2015), they are inappropriate for MMA because forest plots require a symbol on each confidence interval that is proportional to the weight for each study, but the weighting mechanism in MMA is too complex to be visualized. Therefore, for MMA, the tool for displaying sample effect sizes and their overall estimators is a confidence interval plot without displaying weights. Studies with smaller standard errors for the effect sizes would contribute more to the overall estimators, and these effect sizes have narrower confidence intervals. Hence, although a confidence interval plot does not directly reflect weights for each study, it could provide quite sufficient information for the users.

For point 3, missing data problems in meta-analysis are often tackled through methods of omission, single imputation, such as augmenting the missing values with the sample-size-weighted mean or zero, multiple imputation or integrating the missing pattern into the estimation method such as Higgins et al.'s two-stage method or methods employing a Bayesian framework (Sutton et al., 2000; Yuan and Little, 2009; Rubin, 1976; Graham, 2009; Schafer and Graham, 2002; Mavridis and Salanti, 2013; Allison, 2001; Little and Rubin, 2019). Since MMA requires far more statistical records from each study than univariate meta-analysis, it is harder to get a complete list of effect sizes and sample sizes. Missing data are often omitted by default in packages mixmeta and metaSEM. Meanwhile, mixmeta provides the function mixmetaSim to simulate responses that can be potentially used for missing data imputation, and metaSEM supports handling missing covariates using full information maximum likelihood in metaregression. However, these options do not consider or distinguish different types of effect sizes in detail. For example, when calculating the covariance between two odds ratios, we need to know the sample size  $n_{ikt}$  that counts for individuals reporting both outcomes, j and k, in the treatment group t: if  $n_{ikt}$  is missing, one solution could be taking the minimal value between sample size  $n_{it}$  that counts for individuals reporting outcome j and  $n_{kt}$  for outcome k. Although  $n_{jkt}$  may be inaccurately imputed, this solution could be better than removing the two effect sizes. As a model preparation package, metavcov could

- 72 handle missing data problems more carefully by customizing functions for different types of effect sizes
- 73 case by case. Moreover, the package also offers a function for multiple imputations for missing data, a
- 74 compact computer program that is extensible for different estimation methods of users' choice.

#### 75 **1.1 Models**

85

86

87

88

89

90

91

92

94

In general, an MMA specifies the model at within-study and between-study levels (Wei and Higgins, 2013b). For the within-study level, let  $\hat{\theta}_i$  denote a vector of p observed effect sizes in the ith study, which is assumed from a multivariate normal distribution:

$$\hat{\boldsymbol{\theta}}_{i} \sim \text{MVN}(\boldsymbol{\theta}_{i}, \boldsymbol{\Sigma}_{i}) \text{ with } \boldsymbol{\Sigma}_{i} = \begin{bmatrix} s_{i1}^{2} & \rho_{w12}s_{i1}s_{i2} & \cdots & \rho_{w1p}s_{i1}s_{ip} \\ \rho_{w21}s_{i1}s_{i2} & s_{i2}^{2} & \cdots & \rho_{w2p}s_{i2}s_{ip} \\ \vdots & \vdots & \ddots & \vdots \\ \rho_{wp1}s_{i1}s_{ip} & \rho_{wp2}s_{i2}s_{ip} & \cdots & s_{ip}^{2} \end{bmatrix},$$
(1)

where  $\theta_i$  is the vector of underlying true effect sizes for study i and  $\Sigma_i$  is the within-study variance-covariance matrix, which is composed of the sampling variance of each effect size on the diagonal, denoted by  $s_{ij}^2$   $(j=1,\ldots,p)$  for the jth effect size, and the within-study covariance of each pair of effect sizes on the off-diagonal that reflects within-study correlation, denoted by  $\rho_{wst}$  for the sth and tth effect sizes. Here, index i is omitted for  $\rho_{w.}$  for the reason of simplicity. In the next section, subscript i is added for each study, whereas subscript w is omitted for simplicity since the whole section is about within-study covariances. The assumption for  $\theta_i$  is that the sample is from a multivariate normal distribution that centers around the true effect sizes, denoted by  $\theta = (\theta_1, \theta_2, \ldots, \theta_p)^T$ , as

$$m{ heta}_i \sim ext{MVN}(m{ heta}, m{\Omega}) ext{ with } m{\Omega} = egin{bmatrix} au_1^2 & 
ho_{b12} au_1 au_2 & \cdots & 
ho_{b1p} au_1 au_p \ 
ho_{b21} au_1 au_2 & au_2^2 & \cdots & 
ho_{b2p} au_2 au_p \ dots & dots & \ddots & dots \ 
ho_{bp1} au_1 au_p & 
ho_{bp2} au_2 au_p & \cdots & au_p^2 \end{bmatrix},$$

where  $\Omega$  is the between-study variance-covariance matrix, which is composed of between-study variance for each true effect size on the diagonal and between-study covariance for each pair of effect sizes on the off-diagonal that reflects between-study correlations  $\rho_{b..}$ . This model can also be written as  $\hat{\theta}_i \sim \text{MVN}(\theta, \Sigma_i + \Omega)$ . By adding  $\Omega$ , random effects between studies are accommodated. When  $\Omega = 0$ , the model is referred as a fixed effect model. For meta-regression, it is written as  $y_i \sim \text{MVN}(X_i\beta, \Sigma_i + \Omega)$ , where the notation of  $\hat{\theta}_i$  is substituted by  $y_i$  to follow the notation in regression models.

In order to fit a fixed/random effect meta-analysis or meta-regression, we have to calculate  $\hat{\theta}_i$  and  $\hat{\Sigma}_i$  for study  $i=1,\ldots,N$ . In practice,  $\hat{\Sigma}_i$  is computed from formulas involving  $\hat{\theta}_i$  to replace  $\Sigma_i$  in equation (1), which is discussed in the next section. Although most effect sizes and their variances/covariances in this paper refer to the estimated values, we omit the circumflex in their notations like other papers (Olkin, 1976; Wei and Higgins, 2013b; Borenstein et al., 2021) for the sake of simplicity. Alternatively, one could interpret those notations as the sample estimators from each study conforming to the same formulas as the true underlying random variables. For packages mixmeta and metaSEM, we have to prepare (1) a matrix  $\Theta$ , which is an N by p matrix with  $\hat{\theta}_i$  in each row and is contained via the argument data in both packages, and (2) a matrix  $\Xi$  which is an N by p(p+1)/2 matrix that saves all the variances and covariances from  $\hat{\Sigma}_i$  for study i in each row, denoted by  $S_i$ . Note that  $\hat{\Sigma}_i$  is a p by p symmetric matrix with  $p+(p-1)+(p-2)\cdots+1=p(p+1)/2$  unique elements. It is more convenient to store these unique elements

- 96 in a vector  $S_i$ , which is organized as  $S_i=(s_{i1}^2,\rho_{w21}s_{i1}s_{i2},\ldots,\rho_{wp1}s_{i1}s_{ip},s_{i2}^2,\ldots,\rho_{wp2}s_{i2}s_{ip},\ldots,s_{ip}^2)^T$
- 97 from the lower triagonal entries in  $\hat{\Sigma}_i$ .  $\Xi$  is contained in the argument v in the metaSEM package. For the
- 98 mixmeta package,  $\Xi$  is contained in the argument S, and S also accepts an N-dimensional list of  $p \times p$
- 99 matrices where  $\Sigma_i$  is stored.
- 100 This paper describes how to estimate with-study variance-covariance matrix  $\Sigma_i$  in the next section with
- details including missing data solutions, where the notation  $\hat{\theta}$  is replaced according to different types of
- 102 effect sizes, such as r for correlation coefficients and  $\delta$  for standardized mean differences. Furthermore,
- 103 this paper provides a model estimation section with a data visualization example and a section focusing on
- 104 missing data problems with a simulation study. Package summary and future work are given in the end.

# f 2 ESTIMATING THE WITH-STUDY VARIANCE-COVARIANCE MATRIX $f \Sigma$

#### 105 2.1 Correlation Coefficient

Let  $r_{ist}$  denote the sample correlation coefficient that describes the relationship between variables s and t in study i. Following the notation by Olkin (1976), Becker (2009), and Ahn et al. (2015), we have

$$var(r_{ist}) = (1 - \rho_{ist}^2)^2 / n_i$$

for the variance of  $r_{ist}$ , and the covariance between  $r_{ist}$  and  $r_{iuv}$  is

$$cov(r_{ist}, r_{iuv}) = [.5\rho_{ist}\rho_{iuv}(\rho_{isu}^2 + \rho_{isv}^2 + \rho_{itu}^2 + \rho_{itv}^2) + \rho_{isu}\rho_{itv} + \rho_{isv}\rho_{itu} - (\rho_{ist}\rho_{isu}\rho_{isv} + \rho_{its}\rho_{itu}\rho_{itv} + \rho_{ius}\rho_{iut}\rho_{iuv} + \rho_{ivs}\rho_{ivt}\rho_{ivu})]/n_i,$$
(2)

where  $\rho_{i..}$  represents the corresponding population value. In practice,  $\rho_{i..}$  can be substituted by the observed sample correlation  $r_{i..}$  (Ahn et al., 2015), and  $\text{var}(r_{ist})$  and  $\text{cov}(r_{ist}, r_{iuv})$  could be calculated by setting the argument method = "each" in the function r.vcov(). Note that the calculation of  $\text{cov}(r_{ist}, r_{iuv})$  also involes  $r_{iut}, r_{itv}, \ldots$  that could be missing in the real data, which may make it impossible to conduct MMA for  $r_{ist}$  and  $r_{iuv}$ . In this case, the argument method can be set as "average", so that sample-size weighted mean from all available studies can be chosen to replace  $\rho_{i..}$  in equation (2), which was proposed by Cooper et al. (2009). Furthermore, we can transform  $r_{ist}$  into the Fisher's z score as

$$z_{ist} = \frac{1}{2} \ln \left( \frac{1 + r_{ist}}{1 - r_{ist}} \right).$$

When Fisher's z scores are used, variances and covariances can be computed as

$$var(z_{ist}) = 1/(n_i - 3) \text{ and } cov(z_{ist}, z_{iuv}) = cov(r_{ist}, r_{iuv}) / \left[ (1 - \rho_{ist}^2)(1 - \rho_{iuv}^2) \right].$$

- 106 Besides arguments method and n as the sample size, the R function r.vcov() needs another argument
- 107 conflat to input correlation coefficients from studies as an N by p matrix where values of  $r_{ist}$  are saved
- 108 in each row. The computed z scores are saved in the output value ef, which is an N by p matrix in the
- 109 same format of argument corflat shown in Figure 2 in blue. From r.vcov(), the output values ef,
- 110 list.vcov and matrix.vcov are calculated Fisher's z scores and their covariances; the corresponding
- values in the scale of Pearson's correlation coefficients are stored in output values r, list.rvcov and
- 112 marix.rvcov. In the next subsections, the function mix.vcov() can be used for other effect sizes,
- 113 which also provides output values ef, list.vcov and matrix.vcov.

From r.vcov(), the output value list.rvcov is a list of N matrices, in which list.rvcov[[i]] stores  $var(r_{ist})$  and  $cov(r_{ist}, r_{iuv})$  in equation (2) for study i. The following shows the example from Cooper et al. (2009) on page 388 as an illustration.

```
r \leftarrow matrix(c(-0.074, -0.127, 0.324, 0.523, -0.416, -0.414), 1)
n < -142
computvcov <- r.vcov(n = n, corflat = r,
                              name = paste("C", c("st", "su",
                                     "tu", "tv", "uv"), sep = ""),
                              method = "each")
round(computvcov$list.rvcov[[1]], 4)
        Cst
                        Csv
                                 Ctu
                Csu
                                         Ctv
                                                 Cuv
        Cst
             0.0070
                     0.0036 -0.0025 -0.0005
                                              0.0018
                                                       0.0009
        Csu
             0.0036 0.0068 -0.0025 -0.0002
                                              0.0008
                                                       0.0017
        Csv -0.0025 -0.0025 0.0056 0.0001
                                              0.0000 - 0.0003
        Ctu -0.0005 -0.0002
                              0.0001
                                      0.0037 -0.0013 -0.0013
             0.0018
                     0.0008
                              0.0000 - 0.0013
                                              0.0048
                                                       0.0022
        Ctv
             0.0009
                     0.0017 -0.0003 -0.0013
                                              0.0022
                                                       0.0048
        Cuv
```

117 The z transformed correlation coefficients are saved in the output vector ef.

```
round(computvcov$ef, 4)
          Cst
                  Csu
                           Csv
                                  Ctu
                                           Ctv
                                                  Cuv
        1 - 0.0741 - 0.1277 0.3361 0.5805 - 0.4428 - 0.4404
round(computvcov$list.vcov[[1]], 4)
        Cst.
                Csu
                         Csv
                                 Ctu
                                                  Cuv
        Cst
             0.0072
                      0.0037 -0.0029 -0.0008
                                               0.0022
                                                       0.0011
             0.0037 0.0072 -0.0028 -0.0003
        Csu
                                               0.0010
                                                       0.0021
        Csv -0.0029 -0.0028
                             0.0072 0.0001
                                               0.0000 - 0.0004
        Ctu -0.0008 -0.0003
                              0.0001
                                      0.0072 -0.0022 -0.0022
                              0.0000 - 0.0022
        Ctv
             0.0022
                      0.0010
                                               0.0072
                                                       0.0032
        Cuv
             0.0011
                      0.0021 - 0.0004 - 0.0022
                                               0.0032
                                                       0.0072
```

Note that for m outcomes, there are  $p=m\times (m-1)/2$  correlation coefficients. Since the p by p variance-covariance matrix is symmetric, there are  $p+(p-1)+(p-2)\cdots+1=p(p+1)/2$  unique elements. It is more convenient to store these unique elements in a vector so that if we have N studies, we could have an N by p(p+1)/2 matrix that saves all the variances and covariances, which can be obtained from the output value matrix.vcov. The bottom row in Figure 2 is an illustration of how the variances and covariances are arranged in matrix and list formats. Following the above code, we have

For missing values, we could impute a numeric value such as zero via the argument na.impute.

125 By default, we have na.impute = NA without any imputation. Under the default setting of method 126 = "average", the calculation of  $cov(r_{ist}, r_{iuv})$  is still possible even though it involes  $r_{iut}, r_{itv}, \dots$ that could be missing. Besides imputing a specific number via na.impute, we could also impute 127 the sample-size-weighted mean from those studies with complete records by setting the argument na.impute = "average". Basically, na.impute = "average" imputes the mean values 129 for  $r_{ist}, z_{ist}$  and  $cov(r_{ist}, r_{iuv})$ , while method = "average" imputes the mean values only for 130  $cov(r_{ist}, r_{iuv})$ . These two arguments, na.impute = "average" and method = "average", match the mean imputation method and the method of omission illustrated in Section 4.2 for the missing 132 data problem. Note that all the discussion about missing data in Sections 2 and 3 is about missingness in within-study factors. Missingness in between-study factors can only be handled in functions described in Section 4. 135

# 136 2.2 Standardized Mean Difference

For the treatment group, let  $n_{jt}$ ,  $n_{kt}$  and  $n_{jkt}$  denote the numbers of participants who report outcome j, outcome k, and both outcomes j and k. Similarly, denote  $n_{jc}$ ,  $n_{kc}$  and  $n_{jkc}$  for the control group. These notations are used for all the effect sizes for treatment comparison, including standardized mean difference (SMD), mean difference, log odds ratio, log risk ratio, and risk difference. There are two ways to estimate the population SMD, Hedges' g and the sample SMD. Denote the sample mean score on outcome g in the treatment and control groups as g0, respectively, and the standard deviation of the scores as g0, and g0, Hedges (1981) proposed a minimum variance unbiased estimator for the population SMD, which is defined as

$$g_j = \frac{\delta_j}{J(v_j)} \text{ with } J(v_j) = \frac{\Gamma(v_j/2)}{\sqrt{\frac{v_j}{2}\Gamma(\frac{v_j-1}{2})}} \text{ and } v_j = n_{jt} + n_{jc} - 2,$$

where

$$\delta_j = rac{ar{y}_{jt} - ar{y}_{jc}}{s_j^{
m pool}} ext{ with } s_j^{
m pool} = \sqrt{rac{(n_{jt} - 1)s_{jt}^2 + (n_{jc} - 1)s_{jc}^2}{n_{jt} + n_{jc} - 2}}.$$

Wei and Higgins (2013b) derived the covariance between two effect sizes in terms of Hedges' g, denoted by  $g_i$  and  $g_k$ , as follows

$$\begin{aligned} \cos(g_{j},g_{k}) &= \rho \Big( \frac{n_{jkc}}{n_{jc}n_{kc}} + \frac{n_{jkt}}{n_{jt}n_{kt}} \Big) + \frac{k_{jk}}{k_{j}k_{k}} \rho^{2} \delta_{j} \delta_{k} J(v_{j}) J(v_{k}) \sqrt{\Big( \frac{v_{j}}{v_{j} - 2} - \frac{1}{J(v_{j})^{2}} \Big) \Big( \frac{v_{k}}{v_{k} - 2} - \frac{1}{J(v_{k})^{2}} \Big)}, \\ \text{where } k_{k} &= \frac{2n_{kt} + 2n_{kc} - 4}{(n_{kc} + n_{kt} - 2)^{2}}, \\ k_{j} &= \frac{2n_{jt} + 2n_{jc} - 4}{(n_{jc} + n_{jt} - 2)^{2}}, \\ k_{jk} &= \frac{2}{(n_{jc} + n_{jt} - 2)(n_{kc} + n_{kt} - 2)} \Big( \frac{n_{jt}n_{kt}}{n_{jt} + n_{kt} - 1} + \frac{n_{jc}n_{kc}}{n_{jc} + n_{kc} - 1} - 2 \Big), \end{aligned}$$

137 and  $\rho$  is a simplified notation of  $\rho_{wjk}$  in equation (1).

We could use the function smd.vcov() for calculating Hedges' g from SMD, which is stored in the 138 output value ef. The input arguments for  $\delta_i$ ,  $n_{it}$  and  $n_{ic}$  are d, nt and nc which are all  $N \times p$  matrices in 139 the same arrangement as ef in Figure 2. The arguments for  $\rho$ ,  $n_{ikt}$  and  $n_{ikc}$  are r, n\_rt and n\_rc which 140 are all in a list format with  $N p \times p$  matrices. If  $n_{jkt}$  or  $n_{jkc}$  is missing, the function automatically imputes 141 142  $n_{jkt}$  by the minimal value between  $n_{jt}$  and  $n_{kt}$ , and imputes  $n_{jkc}$  by the minimal value between  $n_{jc}$  and  $n_{kc}$ . This imputation method is used for all the effect sizes for treatment comparison, including SMD, mean 143 difference, log odds ratio, log risk ratio, and risk difference. The variances and covariances of Hedges' g are stored in matrix.vcov and list.vcov in the same arrangement shown in the bottom row of 145 Figure 2. 146

The function smd.vcov() also provides the formula in Olkin and Gleser (2009) for the covariance of the sample SMD,  $\delta_j$ , which is defined as

$$cov(\delta_j, \delta_k) = \frac{(n_t + n_c)\rho}{n_t n_c} + \frac{\delta_j \delta_k \rho^2}{2(n_t + n_c)}.$$

The results are stored in the output values matrix.dvcov and list.dvcov in the same formats of matrix.vcov and list.vcov, respectively. To demonstrate the usage of smd.vcov(), the dataset in Geeganage and Bath (2010) is applied using variables  $SMD\_SBP$  and  $SMD\_DBP$ , which measure the systolic blood pressure (SBP, in mHg) and diastolic blood pressure (DBP, in mHg). The correlation between SBP and DBP is not recorded in the paper, so we impute it as 0.71 based on expert knowledge — ideally, different correlation coefficients should be recorded from N different primary studies saved in a list of N correlation matrices.

```
computvcov <- smd.vcov(nt = Geeganage2010[ ,c("nt_SBP", "nt_DBP")],</pre>
                       nc = Geeganage2010[ ,c("nc_SBP", "nc_DBP")],
                       d = Geeganage2010[ ,c("SMD_SBP", "SMD_DBP")],
                       r = r.Gee
                       name = c("SBP", "DBP"))
                      ## Hedge's g
head(computvcov$ef)
                          DBP
        1 -0.075002006 -0.19339306
        2 0.043155405 -0.01610660
        3 - 0.242782681 - 0.31689607
        4 -0.097028863 -0.16608808
        5 -0.004460966 -0.13364520
        6 -0.286780271 0.08887979
head(computvcov$matrix.vcov)
                              ## variances/covariances for Hedge's g
             var SBP
                        cov SBP DBP
                                       var DBP
        [1,] 0.15560955 0.11051462 0.15591453
        [2,] 0.18256182 0.12959901 0.18254277
        [3,] 0.03190808 0.02264927 0.03198210
        [4,] 0.03115906 0.02212545 0.03119080
        [5,] 0.01965510 0.01395547 0.01967717
        [6,] 0.26813782 0.18910349 0.26680797
head(computvcov$matrix.dvcov)
                               ## variances/covariances for SMD
             var_SBP
                         cov_SBP_DBP
                                       var_DBP
        [1,] 0.15565024 0.11056752 0.15618509
        [2,] 0.18257730 0.12959610 0.18254492
        [3,] 0.03200824 0.02271517 0.03215273
        [4,] 0.03117474
                        0.02213897 0.03123674
        [5,] 0.01965512 0.01395583 0.01969852
        [6,] 0.26896403 0.18897441 0.26688733
```

# 154 2.3 Mean Difference and Log Odds Ratio

Sometimes researchers prefer to keep the original scale of mean differences (MD) instead of standardizing them into SMD, such as body mass index (BMI) (Winter et al., 2014; Torloni et al., 2009) or waist circumference (Czernichow et al., 2011; de Hollander et al., 2012). For dichotomous outcomes such as mortality or morbidity, a popular effect size measurement is the log odds ratio (logOR) (Insua et al., 1994; Thompson et al., 1997). Following the notations for SMD, Wei and Higgins (2013b) also derived the covariances for MD and logOR as

$$cov(MD_j, MD_k) = \frac{n_{jkt}}{n_{jt}n_{kt}} \rho s_{jt} s_{kt} + \frac{n_{jkc}}{n_{jc}n_{kc}} \rho s_{jc} s_{kc}$$

and

$$\operatorname{cov}(\operatorname{logOR}_j, \operatorname{logOR}_k) = \frac{\rho n_{jkc}}{n_{jc} n_{2c}} \sqrt{ \Big( \frac{1}{S_{jc}} + \frac{1}{F_{jc}} \Big) \Big( \frac{1}{S_{kc}} + \frac{1}{F_{kc}} \Big)} + \frac{\rho n_{jkt}}{n_{jt} n_{kt}} \sqrt{ \Big( \frac{1}{S_{jt}} + \frac{1}{F_{jt}} \Big) \Big( \frac{1}{S_{kt}} + \frac{1}{F_{kt}} \Big)},$$

where  $S_{jt}$  and  $S_{jc}$  are the numbers of participants with the outcome j event in the treatment and control groups, respectively, and  $F_{jt}$  and  $F_{jc}$  are the respective numbers without the event. Functions md.vcov() and logOR.vcov() can be used to calculate  $cov(MD_j, MD_k)$  and  $cov(logOR_j, logOR_k)$ . Similary to r.vcov() and smd.vcov(), the variance-covariance matrices are stored in the output values matrix.vcov and list.vcov in matrix and list formats, and the calculated log odds ratios are stored in the output value ef. Similar functions in the **metavcov** package include lgRR.vcov() for log risk ratios and rd.vcov() for risk differences. The function mix.vcov() is designed for merging all of these functions whose details are demonstrated in the next subsection.

The covariance between MD and logOR is calculated as

$$\begin{split} \text{cov}(\text{MD}_j, \text{logOR}_k) = & \rho s_{jc} \frac{n_{jkc} \sqrt{n_{kc}}}{n_{jc} n_{kc}} \sqrt{\left(\frac{1}{S_{kc}} + \frac{1}{F_{kc}}\right) \left(\frac{1}{S_{kc}} + \frac{1}{F_{kc}}\right)} \\ & + \rho s_{jt} \frac{n_{jkt} \sqrt{n_{kt}}}{n_{jt} n_{kt}} \sqrt{\left(\frac{1}{S_{kt}} + \frac{1}{F_{kt}}\right) \left(\frac{1}{S_{kt}} + \frac{1}{F_{kt}}\right)}, \end{split}$$

which can be obtained using the function md\_lgor(), whose output values include lgor that returns the computed log odds ratio and v that returns the computed covariance.

## 165 2.4 Combination of Effect Sizes

166 Besides correlation coefficients, SMD, MD and logOR, the metavcov package also includes log risk ratio (logRR) and risk difference (RD). The formulas for calculating their covariances can be found in Table 167 1 in Wei and Higgins (2013b) and the corresponding R functions can be found in Figure 3. Similar to the 168 function md\_lgor() in the previous subsection, we have lgor\_lgrr() for covariance between logOR 169 and logRR, lgor\_rd() for covariance between logOR and RD, md\_lgrr() for covariance between 170 MD and logRR, md\_rd() for covariance between MD and RD, md\_smd() for covariance between MD 171 and SMD, smd\_lgor() for covariance between SMD and logOR, smd\_lgrr() for covariance between 172 SMD and logRR, and smd\_rd() for covariance between SMD and RD. These functions are designed 173 for simple calculations to prepare for the function mix.vcov(), which merges all of these functions by specifying the input argument type with options "MD" for mean difference, "SMD" for standardized mean 175 difference, "logOR" for log odds ratio, "logRR" for log risk ratio, and "RD" for risk difference. Its 176 output values ef, matrix.vcov and list.vcov are the calculated effect sizes and covariances in matrix and list formats. 178

In order to demonstrate the usage of mix.vcov(), the dataset in Geeganage and Bath (2010) is applied again. There are four outcomes, including systolic blood pressure (SBP, in mHg), diastolic blood pressure

(DBP, in mHg), death (D), and death or disability (DD). Mean difference is used to measure the two 182 continuous outcomes SBP and DBP. Risk difference and log odds ratio are chosen to measure the two dichotomous outcomes D and DD. The type of their effect sizes is specified via a vector for argument type 183 in order. This order is applied to all the other arguments. Note that certain arguments are not available for specific outcomes. For example, arguments d, sdt and sdc are designed for effect sizes SMD or MD, 185 which are not available for logOR, logRR or RD. Therefore, we have to impute NAs in arguments d, sdt 186 and sdc for outcomes D and DD. Similarly, we have to impute NAs for st and sc for outcomes SBP and 187 DBP. The correlation coefficients between these outcomes are not recorded in the paper, so we impute 188 them based on expert knowledge — ideally, different correlation coefficients should be recorded from N189 different primary studies saved in a list of N correlation matrices. The example code is as follows. 190

```
data (Geeganage2010)
## correlation coefficients between outcomes are missing in the data
## impute the correlation coefficient list based on expert knowledge
r12 < -0.71
r13 < -0.5
r14 < -0.25
r23 < -0.6
r24 < -0.16
r34 < -0.16
r \leftarrow vecTosm(c(r12, r13, r14, r23, r24, r34))
diag(r) < -1
mix.r <- lapply(1:nrow(Geeganage2010), function(i){r})</pre>
attach (Geeganage2010)
computvcov <- mix.vcov(type = c("MD", "MD", "RD", "lgOR"),</pre>
                        d = cbind(MD_SBP, MD_DBP, NA, NA),
                        sdt = cbind(sdt_SBP, sdt_DBP, NA, NA),
                        sdc = cbind(sdc_SBP, sdc_DBP, NA, NA),
                        nt = cbind(nt_SBP, nt_DBP, nt_DD, nt_D),
                        nc = cbind(nc_SBP, nc_DBP, nc_DD, nc_D),
                        st = cbind(NA, NA, st_DD, st_D),
                        sc = cbind(NA, NA, sc_DD, sc_D),
                        r = mix.r
                        name = c("MD.SBP", "MD.DBP", "RD.DD", "lgOR.D"))
## save different effect sizes in y
y <- computvcov$ef
head(y)
                                        lqOR.D
          MD.SBP
                  MD.DBP
                              RD.DD
           -2.47
                  -3.44
                          0.00000000 - 1.0986123
        1
        2
                  -0.34
            1.61
                          0.18750000
                                       0.5959834
        3
          -8.16
                  -6.44
                          0.02554455
                                       0.5892102
        4
           -3.17
                  -3.41
                          0.04000000
                                       0.4444945
        5
           -0.15
                  -2.39
                         0.01920750
                                       0.1000835
           -9.83
                  1.93 -0.25000000 -0.5108256
```

Frontiers 10

computvcov\$list.vcov[[1]]

```
MD.SBP
                            MD.DBP
                                        RD.DD
                                                   lgOR.D
        MD.SBP 87.9883122 34.8140903 0.92452778 2.27820442
        MD.DBP 34.8140903 27.8514100 0.62070000 0.79071907
                0.9245278
                            0.6207000 0.04062500 0.02741618
        lgOR.D
                2.2782044
                            0.7907191 0.02741618 1.02083333
$# save variances/covariances of all the effect sizes in a matrix S
S <- computvcov$matrix.vcov
S[1, ]
 var MD.SBP cov MD.SBP MD.DBP
                                  cov_MD.SBP_RD.DD
                                                     cov_MD.SBP_lgOR.D
     87.98831
                      34.81409
                                       0.9245278
                                                        2.278204
 var_MD.DBP
               cov_MD.DBP_RD.DD
                                  cov_MD.DBP_lgOR.D
                                                      var_RD.DD
     27.85141
                                        0.7907191
                                                        0.040625
                       0.6207
 cov_RD.DD_lgOR.D
                     var_lgOR.D
     0.02741618
                     1.020833
```

The matrices y and S in the above code can be used as input arguments for packages mixmeta and metaSEM, which is demonstrated in the next section. After computing within-study covariances, the next step is model fitting for estimating the overall effect sizes, potentially with result visualizations (see Figure 4).

### 3 ESTIMATING THE OVERALL EFFECT SIZES

#### 195 3.1 Generalized Least Squares (GLS) Methods

The GLS method (Berkey et al., 1996) enables us to estimate the overall effect size  $\theta$  from the observed  $\hat{\theta}_i$  and  $\Sigma_i$  from all the N studies. It is similar to the more familiar ordinary least squares method, but it allows the data from which parameters are estimated to have unequal population variances and nonzero covariances. Becker and Olkin have shown that the GLS estimators are also maximum likelihood estimators (Becker, 2009). This section demonstrates the GLS procedure in order that the next section could present handling the missing data problem under its framework.

First, let  $T_{Np\times 1}=(\hat{\theta}_{11},\hat{\theta}_{12},\ldots,\hat{\theta}_{1p},\hat{\theta}_{21},\hat{\theta}_{22},\ldots,\hat{\theta}_{2p},\ldots,\hat{\theta}_{i1},\hat{\theta}_{i2},\ldots,\hat{\theta}_{ip},\ldots,\hat{\theta}_{N1},\hat{\theta}_{N2},\ldots,\hat{\theta}_{Np})'$  be a rearrangement of elements in  $\hat{\theta}_i$  from all the N studies. Given an error vector, denoted by  $e_{Np\times 1}$ , the

204 relationship between the population parameter  $\theta = (\theta_1, \theta_2, \dots, \theta_p)'$  and T is

$$\boldsymbol{T}_{Np\times 1} = \boldsymbol{X}_{Np\times p}\boldsymbol{\theta} + \boldsymbol{e}_{Np\times 1} = \begin{bmatrix} 1 & 0 & 0 & \cdots & 0 & 0 \\ 0 & 1 & 0 & \cdots & 0 & 0 \\ \vdots & \vdots & \ddots & \vdots & \vdots & \vdots \\ 0 & 0 & \cdots & 1 & \cdots & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \cdots & 0 & 0 & 1 \\ \vdots & \vdots & \ddots & \vdots & \vdots \\ 0 & 0 & \cdots & 0 & 0 \\ 0 & 1 & 0 & \cdots & 0 & 0 \\ 0 & 1 & 0 & \cdots & 0 & 0 \\ \vdots & \vdots & \ddots & \vdots & \vdots & \vdots \\ 0 & 0 & \cdots & 1 & \cdots & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \cdots & 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} \theta_{1} \\ \theta_{2} \\ \vdots \\ \theta_{j} \\ \vdots \\ \theta_{p} \end{bmatrix} + \begin{bmatrix} e_{11} \\ e_{12} \\ \vdots \\ e_{1j} \\ \vdots \\ e_{Nj} \\ \vdots \\ e_{Nj} \\ \vdots \\ e_{Nj} \\ \vdots \\ e_{Np} \end{bmatrix}$$

205 where X is an  $Np \times p$  matrix created by stacking N p-dimensional identity matrices.

Assuming the errors in e are normally distributed with a zero mean vector  $\mathbf{0}$  and a variance-covariance matrix  $\mathbf{\Psi}$ , which is a blockwise diagonal matrix with  $\mathbf{\Sigma}_i$  in its diagonal:

$$m{\Psi} = egin{bmatrix} m{\Sigma}_1 & 0 & 0 & \cdots & 0 & 0 \ 0 & m{\Sigma}_2 & 0 & \cdots & 0 & 0 \ dots & dots & \ddots & dots & dots \ 0 & 0 & \cdots & m{\Sigma}_i & \cdots & 0 \ dots & dots & dots & dots & \ddots & dots \ 0 & 0 & \cdots & 0 & 0 & m{\Sigma}_N \end{bmatrix}.$$

Note that in a random effect model, the matrix in its diagonal is  $\Sigma_i + \Omega$ .

The GLS estimator of  $\theta$  and its variance  $Var(\hat{\theta})$  are given by

$$\hat{\boldsymbol{\theta}} = (\boldsymbol{X}'\boldsymbol{\Psi}^{-1}\boldsymbol{X})^{-1}\boldsymbol{X}'\boldsymbol{\Psi}^{-1}\boldsymbol{T} \text{ and } \operatorname{Var}(\hat{\boldsymbol{\theta}}) = (\boldsymbol{X}'\boldsymbol{\Psi}^{-1}\boldsymbol{X})^{-1}. \tag{4}$$

A test of homogeneity with the null hypothesis  $H_0$ :  $\theta_1 = \theta_2 = \cdots = \theta_j = \cdots = \theta_p$  can be conducted via the Q statistic (Higgins and Thompson, 2002; Sera et al., 2019):

$$Q = \hat{\boldsymbol{\theta}}' [\boldsymbol{\Psi}^{-1} - \boldsymbol{\Psi}^{-1} \boldsymbol{X} (\boldsymbol{X}' \boldsymbol{\Psi}^{-1} \boldsymbol{X})^{-1} \boldsymbol{X}' \boldsymbol{\Psi}^{-1}] \hat{\boldsymbol{\theta}},$$

which follows a Chi-square distribution with df =  $(N-1) \times p$  degrees of freedom. The Q statistic generates the  $I^2$  statistic,

$$I^2 = \max\{\frac{Q - \mathrm{df}}{Q}, 0\},\$$

which quantifies the amount of heterogeneity as the proportion of total variation related to sampling error. A value of 0% indicates no observed heterogeneity, and larger values show increasing heterogeneity

- 210 (Higgins et al., 2003). We can use the function metafixed() for conducting a fixed-effect MMA, is
- 211 equivalent as setting method = "fixed" in mixmeta() using the mixmeta package. However, the
- 212 zero heterogeneity fixed effect model is almost never appropriate for psychology.
- 213 For random effect models, methods including the maximum likelihood and the restricted maximum
- 214 likelihood methods (Harville, 1977; Sera et al., 2019; Gasparrini et al., 2012), the method of moments
- 215 (Chen et al., 2012; Jackson et al., 2013), and the method of two stages proposed by Liu et al. (2009) can be
- 216 used to estimate  $\Omega$  and  $\theta$ . These methods can be adopted in the mixmeta package by specifying method
- 217 as "ml", "reml", "mm" or "vc" in mixmeta(). The metaSEM package adopts the maximum
- 218 likelihood and the full information maximum likelihood methods (Cheung, 2021) in functions meta()
- 210 Intermood and the run information maximum intermood methods (Cheding, 2021) in functions meta ()
- 219 and metaFIML(), respectively. When the effect sizes of interest are correlation coefficients, we can use
- metaSEM for conducting meta-analytic structural equation modeling (Cheung, 2008, 2009, 2013, 2015).
- 221 A simple example for the metaSEM package is demonstrated as follows using y and S obtained via the
- 222 output values ef and matrix.vcov from the previous code. For the maximum likelihood estimation
- 223 method, we have

```
library(metaSEM)
MMA_RE <- summary(meta(y = y, v = S, data = data.frame(y,S)))</pre>
```

For the restricted maximum likelihood (REML) estimation method, we have

```
library(metaSEM)
MMA_RE <- summary(reml(y = y, v = S, data = data.frame(y,S)))</pre>
```

- 225 The argument data in the above functions is unnecessary. This is to show that functions mixmeta(),
- 226 meta() and reml() have the argument data so that covariates or predictors can be added for meta-
- 227 regression.
- In summary, we can use the function r.vcov() for correlation coefficients and mix.vcov() for
- 229 other effect sizes from the metavcov package to calculate effect sizes and covariances, which are stored
- 230 in output values ef and matrix.vcov. Then we can use ef and matrix.vcov to conduct a random
- 231 effect MMA via mixmeta or metaSEM. Note that regardless of the chosen function, estimating the
- full variance-covariance matrix  $\Omega$  can be difficult unless N is large, because there are many parameters
- 233 involved. Therefore, it is often wise to consider constrained models for the variance-covariance matrix
- 234  $\Omega$  (McShane and Böckenholt, 2022).

#### 235 3.2 Data Visualization

- 236 The new version of metavcov offers a plot function plotCI() for displaying confidence intervals
- 237 of effect sizes from each study and the overall estimators. The difference between a forest plot and a
- 238 confidence interval plot is that a forest plot requires a symbol on each confidence interval that is proportional
- 239 to the weight for each study (Schwarzer et al., 2007; Boyles et al., 2011; Rücker and Schwarzer, 2021;
- 240 Sedgwick, 2015). Because the weighting mechanism in MMA is too complex to be visualized, such a
- 241 proportional symbol is omitted. Although a confidence interval plot does not directly reflect weights for
- 242 each study, it could provide sufficient information for users because effect sizes with narrower confidence
- 243 intervals often contribute more to the overall estimators. Following the code from the previous subsection,
- an example for the function plotCI () is given below.

We could also set obj <- MMA\_RE in the above code where MMA\_RE was sepecified in the previous subsection from a random effect model using the package mixmeta or metaSEM. The result is shown in Figure 5.

#### 4 THE MISSING DATA PROBLEM

We can conveniently specify the predictors or missing values uing the design matrix X in equation (3). First, let X be informally denoted as  $X = (\mathbb{X}(1), \mathbb{X}(2), \dots, \mathbb{X}(i), \dots, \mathbb{X}(N))'$  for simplicity, where  $\mathbb{X}(i)$  is a p-dimensional identidy matrix in equation (3). If we want to fit a meta-regression model (Van Houwelingen et al., 2002) with covariates or predictors  $x_{i1}, x_{i2} \dots$  from each study, we can rewrite  $\mathbb{X}(i)$  as

$$\mathbb{X}(i) = \begin{bmatrix} 1 & 0 & 0 & \cdots & 0 & 0 & x_{i1} & x_{i2} & \dots \\ 0 & 1 & 0 & \cdots & 0 & 0 & x_{i1} & x_{i2} & \dots \\ \vdots & \vdots & \ddots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\ 0 & 0 & \cdots & 1 & \cdots & 0 & x_{i1} & x_{i2} & \dots \\ \vdots & \vdots & \vdots & \vdots & \ddots & \vdots & \vdots & \vdots & \vdots \\ 0 & 0 & \cdots & 0 & 0 & 1 & x_{i1} & x_{i2} & \dots \end{bmatrix},$$

and  $\theta$  as  $(\beta_{01}, \beta_{02}, \dots, \beta_{0p}, \beta_{11}, \beta_{12}, \dots, \beta_{1p}, \beta_{21}, \beta_{22}, \dots, \beta_{2p}, \dots)'$ . In R, we could use mixmeta or metaSEM to conduct meta-regression. Following the code from the previous section, we could use the code below assuming the predictor is the percentage of male participants in each study.

For the missing data problem, if the jth observed effect size is missing in study i, we could simply delete the jth row in  $\mathbb{X}(i)$ . For example, we expect C1, C2, ..., C6 to be observed for all studies in the Craft et al. meta-analysis (2003), where  $\mathbb{X}(i) = I_6$  for X in equation (3). However, if C5 is missing in study i, then

251

254

257 258

259

260

267

we could input

$$\mathbb{X}(i) = \begin{bmatrix} 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 1 \end{bmatrix}$$

in the design matrix X. This method of omission for missing values in MMA is different from other 252 regression functions such as those performed through lm() or qlm() in R. Specifically, partially missing outcomes do not prevent the study from contributing to estimation. Besides this omission procedure, we 253 can also impute missing data by zero or the sample-size-weighted mean of observed effect sizes. Another 255 way is to integrate the missing pattern in the estimation method such as Higgins et al.'s two-stage method (2008) or methods employing a Bayesian framework (Yuan and Little, 2009; Sutton et al., 2000). Although 256 these techniques are not yet available for the current package, they may well become the methods of choice in the future. Single imputation for missing effect sizes is available through the input argument na.impute in functions r.vcov() and mix.vcov(). Another option is multiple imputation.

#### **Multiple Imputation** 4.1

261 Multiple imputation (MI) is a general approach that allows for the uncertainty about the missing data by generating several different plausible imputed datasets and appropriately combining results obtained 262 from each of them (Rubin, 1976; Graham, 2009; Schafer and Graham, 2002; Mavridis and Salanti, 2013; 263

Allison, 2001; Little and Rubin, 2019). There are three basic phases for MI: 264

1. Imputation Phase: The missing data are imputed from simulated values drawn from some distributions. 265 This process is repeated M times. 266

2. Analysis Phase: The same analysis is performed for each of the M complete datasets.

3. Pooling Phase: The M results are pooled to obtain the final result in some fashion. 268

Since MI methods involve recalculating the variance-covariance matrices for the studies with missing values, the new version of metavcov includes a function metami () that can conduct MI automatically. For the imputation phase, this function imports the package mice published by Van Buuren and Groothuis-Oudshoorn (2011) that imputes incomplete multivariate data by chained equations. The mice package is also recommended by the metafor package for univariate meta-analysis Viechtbauer (2021). For the analysis phase, all the functions mentioned in the previous section are accommodated, including metafixed(), mixmeta() and meta(). The pooling phase is performed via Rubin's rules (Van Buuren and Groothuis-Oudshoorn, 2011; Rubin, 1987; Barnard and Rubin, 1999). Let  $\hat{\theta}_{*m}$ be the estimated coefficient from the mth imputed dataset for one of the p dimensions in  $\theta$ , where  $m=1,\ldots,M$ . The pooled coefficient from MI, denoted by  $\bar{\theta}$ , is simply just an arithmetic mean of the individual coefficients estimated from each of the M analyses. We have

$$\bar{\theta} = \frac{\sum_{m=1}^{M} \hat{\theta}_{*m}}{M}.$$

Estimation of the standard error for each variable is a little more complicated. Let  $V_W$  be the withinimputation variance, which is the average of the variance of the estimated coefficient from each imputed

dataset:

$$V_W = \frac{\sum_{m=1}^{M} \operatorname{Var}(\hat{\theta}_{*m})}{M},$$

where  $Var(\hat{\theta}_{*m})$  is the diagonal element of  $Var(\hat{\theta})$  calculated from equation (4) using the imputed dataset. Let  $V_B$  be the between-imputation variance, which is calculated as

$$V_B = \frac{\sum_{m=1}^{M} (\hat{\theta}_{*m} - \bar{\theta})^2}{M - 1}.$$

From  $V_W$  and  $V_B$ , the variance of the pooled coefficients is calculated as

$$\operatorname{Var}(\bar{\theta}) = V_W + V_B + \frac{V_B}{M}.$$

- 269 The above variance is statistically principled since  $V_W$  reflects the sampling variance and  $V_B$  reflects the
- 270 extra variance due to the missing data.
- Examples of metami () are provided as follows for the data from the Craft et al. meta-analysis (2003)
- 272 in the previous section.

```
## prepare a dataset with missing values
Craft2003.mnar <- Craft2003[, c(2, 4:10)]
Craft2003.mnar[sample(which(Craft2003$C4 < 0), 6), "C4"] <- NA
## prepare input arguments for metami()
dat <- Craft2003.mnar
n.name <- "N"
ef.name <- c("C1", "C2", "C3", "C4", "C5", "C6")</pre>
```

- 273 The number of imputations is specified through the argument M. The argument vcov controls the
- function to be used for computing the variance-covariance matrices for the effect sizes, whose options
- 275 are vcov="r.vcov" for correlation coefficients and vcov="mix.vcov" for all the other types of
- 276 effect sizes. For a random effect model, we can specify the argument func as "mixmeta", which allows
- 277 the function mixmeta() from the package mixmeta to be used for MMA. For the argument func =
- 278 "mixmeta", we have to specify formula and method for mixmeta().

- We could also use func = "meta" in the above code which adopts the function meta() from the metaSEM package, for which it is unnecessary to specify arguments formula and method.
- For meta-regression, we can specify the name of the predictors in the argument x.name:

284

294

295

296

297

298

299

300

301

302

303

307

308

309

310

311

312

282 If we specify func = "mixmeta" in the above code, we also have to add p\_male in the argument 283 formula.

# 4.2 A Simulation Study for the Craft et al. Meta-Analysis (2003)

The metavcov package provides several solutions for handling missing data. In order to compare these 285 286 methods and find the influence of M in the MI method, a simulation study is conducted using the settings in the previous section. There are three missing data mechanisms, including missing completely at random 287 (MCAR), missing at random (MAR), and missing not at random (MNAR). MCAR refers to the situation 288 that neither the variables in the dataset nor the unobserved values of the variable itself predict whether 289 a value will be missing; MAR refers to the circumstance that other variables (but not the variable with 290 missing values itself) in the dataset can predict the missingness of a given variable; a variable is said to 291 be MNAR if the value of the unobserved variable itself predicts missingness (Graham, 2009; Schafer and 292 Graham, 2002; Mavridis and Salanti, 2013; Allison, 2001; Little and Rubin, 2019). 293

The code in the previous section simulated a missing data pattern of MNAR in C4, where only negative values were possibly missing. The MNAR scenario is the most challenging of the three. In order to check the performance of different methods, this procedure was replicated 100 times (B=100). For the MI method, the number of imputations M was varied as 10, 20, 50, and 100. Besides MI, methods of omitting the missing values (omission with mean imputed covariances) and single imputation with sample-size-weighted means (mean imputation) were also included. Recall that from equation (2), missingness in C4 could cause problems for the calculation of covariances between two other correlation coefficients, which makes an MMA impossible. Therefore, sample-size-weighted mean is used for imputing missing values in C4 for calculating covariances, which is achieved by specifying method = "average" in r.vcov().

Bias and mean squared error (MSE) were used to evaluate the methods for which the true parameter, denoted by  $\theta^{\rm RE}$ , was defined as the estimated coefficient from the complete dataset using the function mixmeta() from the mixmeta package with its argument method = "reml". Let  $\bar{\theta}_b$  be the estimated parameter using the imputed dataset from realization b. The bias and MSE were estimated by

$$\widehat{\mathrm{Bias}}(\theta^{\mathrm{RE}}) = \frac{\sum_{b=1}^{B} (\bar{\theta}_b - \theta^{\mathrm{RE}})}{B} \text{ and } \widehat{\mathrm{MSE}}(\theta^{\mathrm{RE}}) = \frac{\sum_{b=1}^{B} (\bar{\theta}_b - \theta^{\mathrm{RE}})^2}{B}.$$

In this dataset, we have N=18 studies and the missing percentage in C4 is 33%. The effect sizes were transformed into Fisher's z scores. The R code for this simulation can be found in the supplementary material.

The simulation results are displayed in Figure 6. The results were based on Fisher's z scores. All methods worked well since the values of bias were all roughly smaller than 0.002. For bias, the method of omission provided a smaller bias, but the results were highly variable. Mean imputation and MI methods gave more consistent results. Because smaller values were more likely to be missing, imputation methods tended to impute larger values based on observed data, generating positive bias. The mean imputation method had a higher bias, which caused higher values of MSE. The results showed that MI methods perform the

- best. Interestingly, the number of imputations M does not affect the result much. It seems that M=20 is
- 314 sufficient. Although missingness in C4 could influence the estimation of other effect sizes in terms of both
- 315 bias and MSE, such influences are on a small scale. Overall, the missing value solutions from metavcov
- 316 seem promising. Note that this conclusion is very specific to this dataset in this particular missingness
- 317 pattern. The purpose of this section is to provide code (see supplemental data) for the users to conduct
- 318 simulations for their own data to get some ideas of parameter settings and perhaps gain some confidence.

#### 5 SUMMARY AND FUTURE WORK

- 319 The metavcov package provides useful tools for conducting MMA with examples in R under a
- 320 generalizable, statistically principled analytical framework. It is very flexible in accommodating functions
- 321 for different effect sizes and functions for different coefficient estimation methods. Compared with its
- 322 earlier versions, functions have more consistent output values: all the model preparation functions, such as
- 323 r.vcov and mix.vcov, store the outputs in ef, list.vcov and matrix.vcov. It is very practical
- 324 with functions for data visualization and handling missing values. As well as being statistically principled,
- 325 it is helpful in practice that once the model has been specified, MI can be conducted automatically. Besides
- 326 end-users, developers can easily extend this package to other existing state-of-the-art trust models (Hedges
- 327 et al., 2010; Pustejovsky and Tipton, 2018; Tipton, 2015; Chen et al., 2015, 2017).
- 328 The MI method was examined in an MNAR scenario from a simulation study. The MNAR scenario is
- 329 very realistic for meta-analysis, which is also known as publication bias. Since published papers tend to
- 330 show significant findings or be in favor of positive results, it is possible that imputing the missing effect
- 331 sizes by zero could balance the findings and outperforms the MI method. The current version integrates
- 332 the mice package for MI. Other packages for modeling missing data such as Amelia (Honaker et al.,
- 333 2011) and mi (Gelman and Hill, 2011) may also be of users' choice for future work. Different estimation
- 334 methods for random effect models, such as the method of moments or Bayesian approaches (Wei and
- Higgins, 2013a), should be compared as well for simulation studies. However, due to space limitations,
- 336 they are not demonstrated in this paper. From a theoretical perspective, no work has been done to calculate
- 337 the covariances between correlation coefficients and other types of effect sizes, such as log odds ratio,
- 338 which is also one of our future goals.

#### CONFLICT OF INTEREST STATEMENT

- 339 The authors declare that the research was conducted in the absence of any commercial or financial
- 340 relationships that could be construed as a potential conflict of interest.

#### **AUTHOR CONTRIBUTIONS**

341 The author confirms being the sole contributor of this work and has approved it for publication.

# **FUNDING**

- 342 This study was funded by the Department of Public Health Sciences 2023 Copeland Foundation Project
- 343 Initiative Award, University of Miami.

#### **ACKNOWLEDGMENTS**

- 344 I thank Soyeon Ahn for her mentorship in the Department of Educational and Psychological Studies at the
- 345 University of Miami School of Education and Human Development.

#### SUPPLEMENTAL DATA

346 The R code for this paper was uploaded to the github link of the package.

#### DATA AVAILABILITY STATEMENT

The datasets analyzed for this study can be found in the github link of the package.

#### **REFERENCES**

- 348 Ahn, S., Lu, M., Lefevor, G. T., Fedewa, A. L., and Celimli, S. (2015). Application of meta-analysis in
- sport and exercise science. In An introduction to intermediate and advanced statistical analyses for sport
- and exercise scientists, eds. N. Ntoumanis and N. D. Myers (John Wiley & Sons), chap. 11. 233–253
- 351 Allison, P. D. (2001). *Missing data* (Sage publications)
- 352 Aloe, A. M., Amo, L. C., and Shanahan, M. E. (2014a). Classroom management self-efficacy and burnout:
- A multivariate meta-analysis. *Educational psychology review* 26, 101–126
- 354 Aloe, A. M., Shisler, S. M., Norris, B. D., Nickerson, A. B., and Rinker, T. W. (2014b). A multivariate
- meta-analysis of student misbehavior and teacher burnout. Educational Research Review 12, 30–44
- 356 Barnard, J. and Rubin, D. (1999). Miscellanea. Small-sample degrees of freedom with multiple imputation.
- 357 *Biometrika* 86, 948–955. doi:10.1093/biomet/86.4.948
- 358 Becker, B. J. (2000). Multivariate meta-analysis. Handbook of applied multivariate statistics and
- 359 *mathematical modeling*, 499–525
- 360 Becker, B. J. (2009). Model-based meta-analysis. In The handbook of research synthesis and meta-analysis,
- eds. H. Cooper, L. V. Hedges, and J. C. Valentine (Russell Sage Foundation), chap. 20. 377–395
- 362 Berkey, C., Anderson, J., and Hoaglin, D. (1996). Multiple-outcome meta-analysis of clinical trials.
- 363 Statistics in medicine 15, 537–557
- 364 Borenstein, M., Hedges, L. V., Higgins, J. P., and Rothstein, H. R. (2021). Introduction to meta-analysis
- 365 (John Wiley & Sons)
- 366 Boyles, A. L., Harris, S. F., Rooney, A. A., and Thayer, K. A. (2011). Forest plot viewer: a new graphing
- 367 tool. *Epidemiology* 22, 746–747
- 368 Chen, H., Manning, A. K., and Dupuis, J. (2012). A method of moments estimator for random effect
- multivariate meta-analysis. *Biometrics* 68, 1278–1284
- 370 Chen, Y., Hong, C., and Riley, R. D. (2015). An alternative pseudolikelihood method for multivariate
- 371 random-effects meta-analysis. *Statistics in medicine* 34, 361–380
- 372 Chen, Y., Liu, Y., Chu, H., Ting Lee, M.-L., and Schmid, C. H. (2017). A simple and robust method for
- multivariate meta-analysis of diagnostic test accuracy. Statistics in medicine 36, 105–121
- 374 Cheung, M. (2021). Meta-Analysis using Structural Equation Modeling. R package version 1.2.5.1
- 375 Cheung, M. W.-L. (2008). A model for integrating fixed-, random-, and mixed-effects meta-analyses into
- 376 structural equation modeling. *Psychological methods* 13, 182
- 377 Cheung, M. W.-L. (2009). Constructing approximate confidence intervals for parameters with structural
- equation models. Structural Equation Modeling: A Multidisciplinary Journal 16, 267–294

- Cheung, M. W.-L. (2013). Multivariate meta-analysis as structural equation models. *Structural Equation Modeling: A Multidisciplinary Journal* 20, 429–454
- 381 Cheung, M. W.-L. (2015). *Meta-analysis: A structural equation modeling approach* (John Wiley & Sons)
- 382 Cichonska, A., Rousu, J., Marttinen, P., Kangas, A. J., Soininen, P., Lehtimäki, T., et al. (2016). metacca:
- summary statistics-based multivariate meta-analysis of genome-wide association studies using canonical
- 384 correlation analysis. *Bioinformatics* 32, 1981–1989
- Cooper, H., Hedges, L. V., and Valentine, J. C. (2009). *The handbook of research synthesis and meta-analysis* (Russell Sage Foundation)
- 387 Craft, L. L., Magyar, T. M., Becker, B. J., and Feltz, D. L. (2003). The relationship between the competitive
- state anxiety inventory-2 and sport performance: A meta-analysis. *Journal of sport and exercise* psychology 25, 44–65
- 390 Czernichow, S., Kengne, A.-P., Stamatakis, E., Hamer, M., and Batty, G. D. (2011). Body mass index,
- 391 waist circumference and waist-hip ratio: which is the better discriminator of cardiovascular disease
- mortality risk? evidence from an individual-participant meta-analysis of 82 864 participants from nine
- 393 cohort studies. *Obesity reviews* 12, 680–687
- 394 de Hollander, E. L., Bemelmans, W. J., Boshuizen, H. C., Friedrich, N., Wallaschofski, H., Guallar-
- Castillón, P., et al. (2012). The association between waist circumference and risk of mortality considering
- body mass index in 65-to 74-year-olds: a meta-analysis of 29 cohorts involving more than 58 000 elderly
- persons. *International journal of epidemiology* 41, 805–817
- 398 Gasparrini, A. (2019). Multivariate and Univariate Meta-Analysis and Meta-Regression. R package
- 399 version 1.0.3
- 400 Gasparrini, A., Armstrong, B., and Kenward, M. G. (2012). Multivariate meta-analysis for non-linear and
- other multi-parameter associations. *Statistics in medicine* 31, 3821–3839
- 402 Geeganage, C. and Bath, P. M. (2010). Vasoactive drugs for acute stroke. Cochrane Database of Systematic
- 403 Reviews
- 404 Gelman, A. and Hill, J. (2011). Opening windows to the black box. Journal of Statistical Software 40
- Graham, J. W. (2009). Missing data analysis: Making it work in the real world. *Annual review of psychology* 60, 549–576
- 407 Harville, D. A. (1977). Maximum likelihood approaches to variance component estimation and to related
- 408 problems. *Journal of the American statistical association* 72, 320–338
- 409 Hedges, L. V. (1981). Distribution theory for glass's estimator of effect size and related estimators. *journal*
- 410 of Educational Statistics 6, 107–128
- 411 Hedges, L. V., Tipton, E., and Johnson, M. C. (2010). Robust variance estimation in meta-regression with
- dependent effect size estimates. Research synthesis methods 1, 39–65
- 413 Higgins, J. P. and Thompson, S. G. (2002). Quantifying heterogeneity in a meta-analysis. Statistics in
- 414 *medicine* 21, 1539–1558
- 415 Higgins, J. P., Thompson, S. G., Deeks, J. J., and Altman, D. G. (2003). Measuring inconsistency in
- 416 meta-analyses. *Bmj* 327, 557–560
- 417 Higgins, J. P., White, I. R., and Wood, A. M. (2008). Imputation methods for missing outcome data in
- 418 meta-analysis of clinical trials. *Clinical trials* 5, 225–239
- 419 Honaker, J., King, G., and Blackwell, M. (2011). Amelia II: A program for missing data. Journal of
- 420 Statistical Software 45, 1–47. doi:10.18637/jss.v045.i07
- 421 Hong, C., Duan, R., Zeng, L., Hubbard, R. A., Lumley, T., Riley, R. D., et al. (2020). The Galaxy Plot: A
- New Visualization Tool for Bivariate Meta-Analysis Studies. American Journal of Epidemiology 189,
- 423 861–869. doi:10.1093/aje/kwz286

- Insua, J. T., Sacks, H. S., Lau, T.-S., Lau, J., Reitman, D., Pagano, D., et al. (1994). Drug treatment of 424
- hypertension in the elderly: a meta-analysis. Annals of internal medicine 121, 355–362 425
- Jackson, D., Riley, R., and White, I. R. (2011). Multivariate meta-analysis: potential and promise. Statistics 426 in medicine 30, 2481-2498 427
- Jackson, D., White, I. R., and Riley, R. D. (2013). A matrix-based method of moments for fitting the 428
- 429 multivariate random effects model for meta-analysis and meta-regression. Biometrical Journal 55,
- 430 231-245
- Little, R. J. and Rubin, D. B. (2019). Statistical analysis with missing data, vol. 793 (John Wiley & Sons) 431
- 432 Liu, Q., Cook, N. R., Bergström, A., and Hsieh, C.-C. (2009). A two-stage hierarchical regression model
- for meta-analysis of epidemiologic nonlinear dose-response data. Computational Statistics & Data 433
- Analysis 53, 4157–4167 434
- Lu, M. (2017). Variance-Covariance Matrix for Multivariate Meta-Analysis. R package version 1.0.1 435
- Luo, S., Chen, Y., Su, X., and Chu, H. (2014). mmeta: an r package for multivariate meta-analysis. *Journal* 436 of Statistical Software 56, 1–26 437
- Mavridis, D. and Salanti, G. (2013). A practical introduction to multivariate meta-analysis. Statistical 438 methods in medical research 22, 133–158 439
- McShane, B. B. and Böckenholt, U. (2022). Multilevel multivariate meta-analysis made easy: An 440 introduction to mlmvmeta. Behavior Research Methods, 1–20 441
- Michael Dewey (2021). CRAN Task View: Meta-Analysis. King's College London, London, United 442 Kingdom 443
- 444 Nam, I.-S., Mengersen, K., and Garthwaite, P. (2003). Multivariate meta-analysis. Statistics in medicine 22, 2309–2333 445
- 446 Nikoloulopoulos, A. K. (2020). A multinomial quadrivariate d-vine copula mixed model for meta-analysis
- 447 of diagnostic studies in the presence of non-evaluable subjects. Statistical Methods in Medical Research
- 29, 2988–3005. doi:10.1177/0962280220913898. PMID: 32323626 448
- Olkin, I. (1976). Asymptotic distribution of functions of a correlation matrix (Shinko Tsusho), chap. 11. 449 235-251 450
- Olkin, I. and Gleser, L. (2009). Stochastically dependent effect sizes. The handbook of research synthesis 451 452 and meta-analysis, 357–376
- Pustejovsky, J. E. and Tipton, E. (2018). Small-sample methods for cluster-robust variance estimation and 453
- hypothesis testing in fixed effects models. Journal of Business & Economic Statistics 36, 672-683 454
- R Core Team (2016). R: A Language and Environment for Statistical Computing. R Foundation for 455 Statistical Computing, Vienna, Austria. ISBN 3-900051-07-0 456
- Riley, R. D. (2009). Multivariate meta-analysis: the effect of ignoring within-study correlation. *Journal of* 457 the Royal Statistical Society: Series A (Statistics in Society) 172, 789–811 458
- Rubin, D. B. (1976). Inference and missing data. *Biometrika* 63, 581–592 459
- Rubin, D. B. (1987). Multiple imputation for nonresponse in surveys, vol. 81 (John Wiley & Sons) 460
- Rücker, G. and Schwarzer, G. (2021). Beyond the forest plot: The drapery plot. Research synthesis methods 461 462 12, 13–19
- 463 Schafer, J. L. and Graham, J. W. (2002). Missing data: our view of the state of the art. Psychological methods 7, 147 464
- Schwarzer, G. et al. (2007). meta: An r package for meta-analysis. R news 7, 40–45 465
- Sebri, V., Durosini, I., Triberti, S., and Pravettoni, G. (2021). The efficacy of psychological intervention on 466
- body image in breast cancer patients and survivors: A systematic-review and meta-analysis. Frontiers in 467
- Psychology 12, 407. doi:10.3389/fpsyg.2021.611954 468

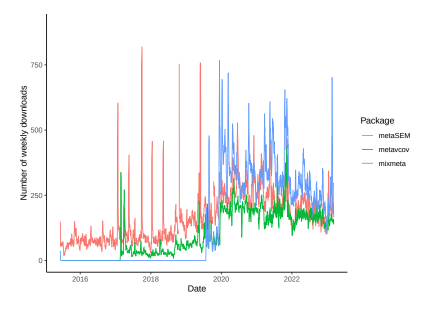
- 469 Sedgwick, P. (2015). How to read a forest plot in a meta-analysis. Bmj 351
- 470 Sera, F., Armstrong, B., Blangiardo, M., and Gasparrini, A. (2019). An extended mixed-effects framework
- for meta-analysis. *Statistics in medicine* 38, 5429–5444
- 472 Sutton, A. J., Abrams, K. R., Jones, D. R., Jones, D. R., Sheldon, T. A., and Song, F. (2000). Methods for
- 473 *meta-analysis in medical research*, vol. 348 (Wiley Chichester)
- 474 Thompson, S. G., Smith, T. C., and Sharp, S. J. (1997). Investigating underlying risk as a source of
- heterogeneity in meta-analysis. *Statistics in medicine* 16, 2741–2758
- 476 Tipton, E. (2015). Small sample adjustments for robust variance estimation with meta-regression.
- 477 Psychological methods 20, 375
- 478 Torloni, M., Betran, A., Horta, B., Nakamura, M., Atallah, A., Moron, A., et al. (2009). Prepregnancy bmi
- and the risk of gestational diabetes: a systematic review of the literature with meta-analysis. *Obesity*
- 480 reviews 10, 194–203
- 481 Van Buuren, S. and Groothuis-Oudshoorn, K. (2011). mice: Multivariate imputation by chained equations
- in r. Journal of statistical software 45, 1–67
- 483 Van Houwelingen, H. C., Arends, L. R., and Stijnen, T. (2002). Advanced methods in meta-analysis:
- 484 multivariate approach and meta-regression. Statistics in medicine 21, 589–624
- 485 Viechtbauer, W. (2010). Conducting meta-analyses in r with the metafor package. *Journal of Statistical*
- 486 *Software* 36, 1–48. doi:10.18637/jss.v036.i03
- 487 [Dataset] Viechtbauer, W. (2021). Multiple imputation with the mice and metafor packages
- 488 Watters, E. R., Aloe, A. M., and Wojciak, A. S. (2021). Examining the associations between childhood
- trauma, resilience, and depression: a multivariate meta-analysis. Trauma, Violence, & Abuse,
- 490 15248380211029397
- 491 Wei, Y. and Higgins, J. P. (2013a). Bayesian multivariate meta-analysis with multiple outcomes. Statistics
- 492 in medicine 32, 2911–2934
- 493 Wei, Y. and Higgins, J. P. (2013b). Estimating within-study covariances in multivariate meta-analysis with
- 494 multiple outcomes. *Statistics in Medicine* 32, 1191–1205
- 495 Winter, J. E., MacInnis, R. J., Wattanapenpaiboon, N., and Nowson, C. A. (2014). Bmi and all-cause
- 496 mortality in older adults: a meta-analysis. *The American journal of clinical nutrition* 99, 875–890
- 497 Yuan, Y. and Little, R. J. (2009). Meta-analysis of studies with missing data. *Biometrics* 65, 487–496

Table 1.	R-packages	for conducting	ng MMA
----------	------------	----------------	--------

Package	Unique features†
metavcov	Preparing within-study variances & covariances; plotting confidence intervals*
mixmeta	Multiple choices for mixed-effect model fitting including maximum likelihood,
	restricted maximum likelihood, method of moments, and variance components
metaSEM	Meta-Analysis using Structural Equation Modeling; plotting model structures
metafor	rma.mv () fits MMA when one outcome is observed at different time points <sup>a</sup>
mmeta	Fitting Bayesian models for binary outcomes <sup>b</sup>
metaCCA	Detecting genetic association with shrinkage for high dimensional outcomes <sup>c</sup>
CopulaREMADA	Fitting copula mixed models for diagnostic test accuracy studies <sup>d</sup>
xmeta	Testing and visualizing publication bias for bivariate meta-analysis <sup>e</sup>

<sup>†</sup> In general, all the listed R-packages can conduct MMA. This table highlights their unique features, rather than major features. They have many features to explore.

<sup>&</sup>lt;sup>a</sup>Viechtbauer (2010); <sup>b</sup>Luo et al. (2014); <sup>c</sup>Cichonska et al. (2016); <sup>d</sup>Nikoloulopoulos (2020); <sup>e</sup>Hong et al. (2020);



**Figure 1.** Number of weekly downloads from CRAN for the three R packages useful for conducting MMA. The package metavcov was initially released in 2017, which is designed for preparing variance-covariance matrices of effect sizes for packages metaSEM and mixmeta that were released in 2015 and 2019, respectively (mixmeta is a new version of the package mvmeta which was initially released in 2011).

<sup>\*</sup>While this paper focuses on demonstrating the utility of metavcov for mixmeta and metaSEM, it can provide similar benefits to other packages as well.

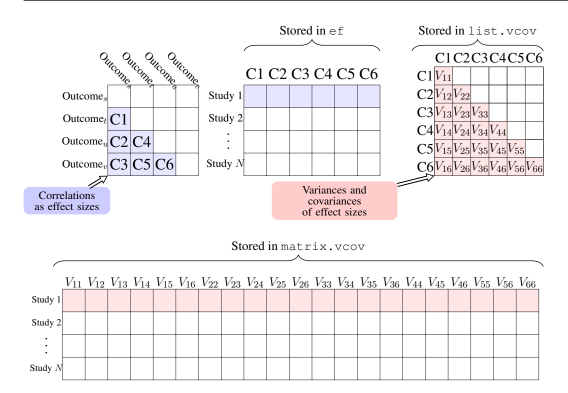


Figure 2. Arrangement of effect sizes and their covariances in matrix and list formats using correlation coefficients as an example. The output value list.vcov is a list of N matrices, in which list.vcov[[i]] represents the matrix  $\Sigma_i$  in equation (1), where the element  $V_{jk}$  in the above figure equals to  $\rho_{wjk}s_{ij}s_{ik}$  in equation (1) and  $V_{jj}$  equals to  $s_{ij}^2$  as the variance of  $\hat{\theta}_{ij}$ . The output value matrix.vcov transforms list.vcov into an  $N \times p(p+1)/2$  matrix. We could use ef and matrix.vcov as input arguments for packages mixmeta or metaSEM to fit an MMA model.

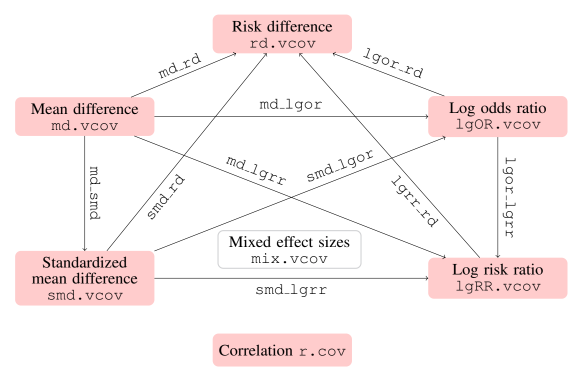


Figure 3. Functions for computing covariances between different types of effect sizes using metavcov. Functions with names connected by dot compute effect sizes of the same type (except mix.vcov) and work with multiple studies, while functions with names connected by underscore compute two effect sizes of different types and only work with one study for simple calculation.

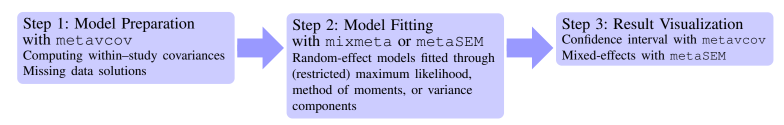
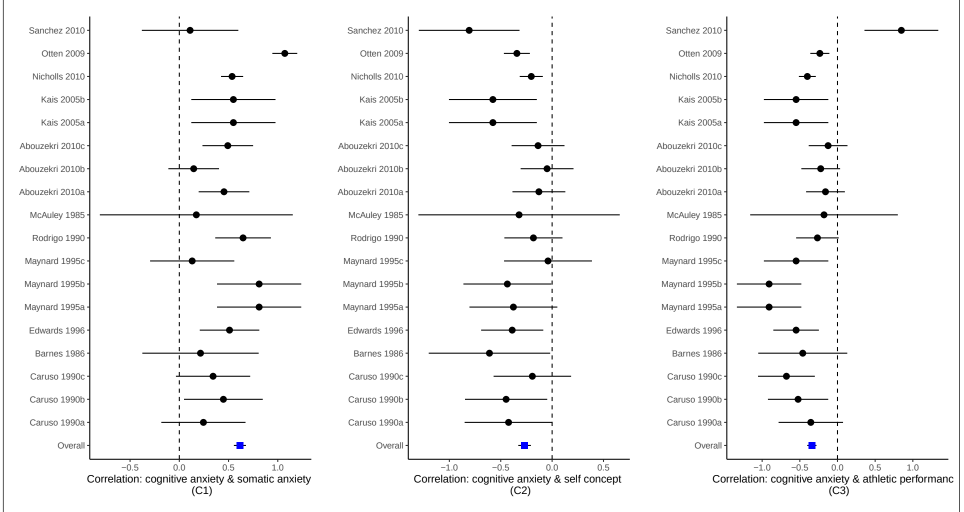


Figure 4. Illustration of the workflow for conducting MMA using the R packages introduced.

Lu



Lu

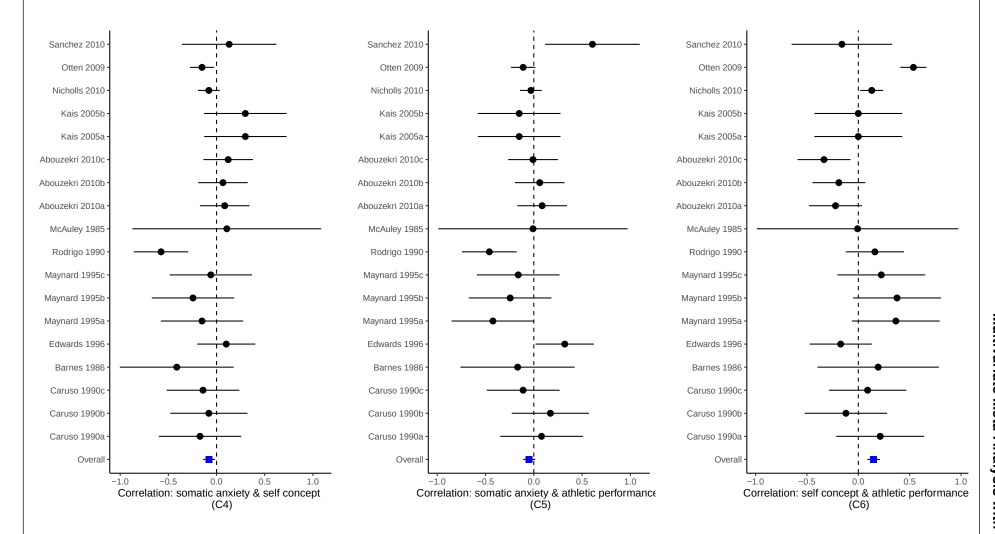
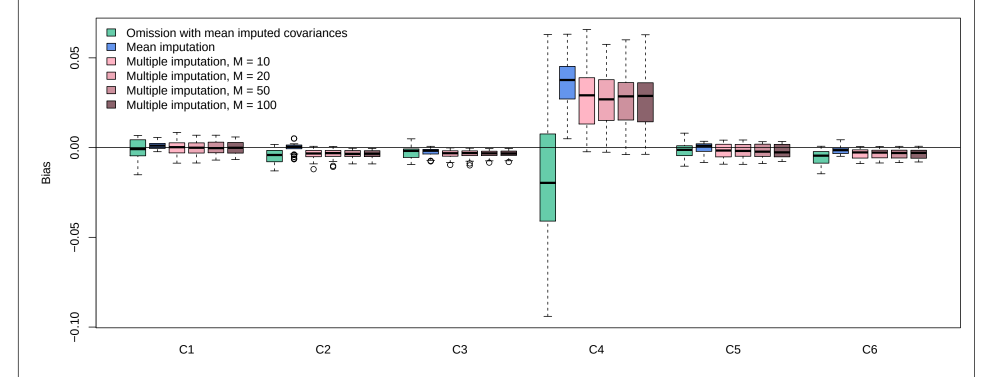
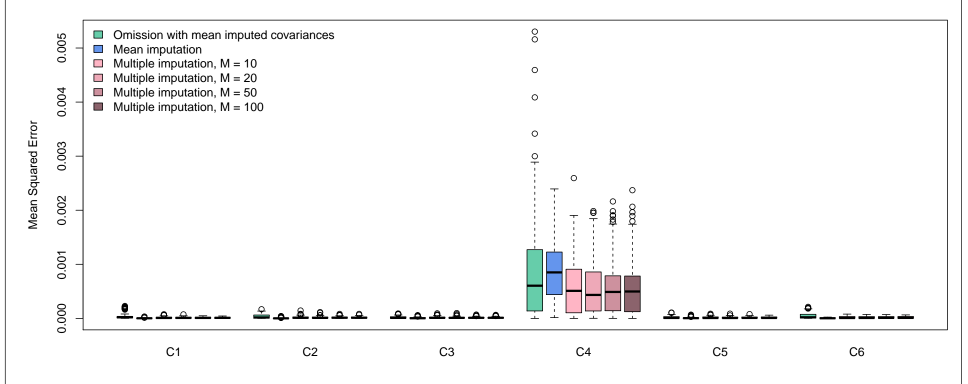


Figure 5. Confidence interval plots for effect sizes from the Craft et al. meta-analysis (2003).

Lu





**Figure 6.** Bias and MSE results from simulation experiments using the data from the Craft et al. meta-analysis (2003). Missing values in C4 were simulated in an MNAR pattern with 33% missing data; there is no missing value in C1, C2, C3, C5 and C6. The multiple imputation method conducted by the function metawicov works better than other methods and M=20 seems sufficient for this specific scenario.