# Statistical Assessment and Adjustment of Publication Bias in the Cochrane Database of Systematic Reviews

Master Thesis in Biostatistics (STA495)

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# Preface

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# Chapter 1

## Introduction

Studies get more attention and are more likely to be published, read and cited if they contain significant effects. Studies with no evidence for an effect are less likely to get published. This generates a bias called "publication bias", a distorted view of the evidence for an effect. Publication bias has been identified as one of the major concerns in irreproducible research (Bishop, 2019). The issue has been discussed in clinical science by many researchers (Dickersin et al. (1987), Sterne et al. (2001a), Dwan et al. (2013)). Consensus is that publication bias exists in clinical science.

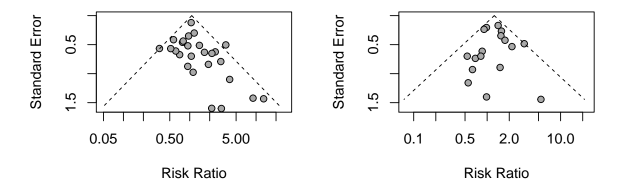
In medical research, clinical trials study the efficacy of therapies and drugs. The gold standard in such intervention studies are randomized controlled trials (RCTs). Results from RCTs influence the treatment of patients in daily clinical practice. The results from multiple intervention studies can be summarized in a meta-analysis to estimate an overall treatment effect (access all studies) (Cochran, 1954). However, publication bias can bias the overall effect estimates from meta-analyses and eventually lead to ineffective treatments that could lead to patient harm, distortion and financial expenses.

There is extensive literature on publication bias in meta-analyses, e.g. (Jones et al. (2013), Turner et al. (2008), Egger et al. (2003), McAuley et al. (2000)). The authors agree on that the exclusion of unpublished results in meta-analyses is responsible for overestimation of treatment effects. Although there are policies that make it mandatory to make all study results publicly accessible (Law, 2007), it is not clear if the situation has improved yet.

There are multiple ways to assess the amount of publication bias. For instance, it is possible to follow studies and assess if they are getting published depending on their findings (Dwan et al. (2013), Decullier et al. (2005), Lee et al. (2008)). One often finds that positive findings (i. e. large effects) are reported and published more often (see also an example in social sciences: Franco et al. (2014)). Another way is to compare results in study registries with results published in journals (e. g. Jones et al. (2013)). Again one finds systematic differences between published and unpublished results.

A third way is to assess the so-called small study effect in a meta-analysis, that is, smaller studies sometimes showing different, often larger treatment effects than large ones (cite sb). Importantly, this method does not assess publication bias directly. But the estimation of small study effects is oftentimes the most efficient way to investigate publication bias in a large number of meta-analyses. However, evidence for small study effects can oftentimes be interpreted as evidence for publication bias (Egger et al., 1997), as this is the most likely cause, even though there may also be other explanations. The underlying rationale is that journal editors are guided by two simple rules (Ioannidis and Trikalinos (2007b), Ioannidis and Thombs (2019)):

- Publish new and trend-setting findings that lead to many citations, increasing the journal impact factor (JIF).
- Publish large trials that are likely to set the standard for a given scientific question, again



**Figure 1.1:** Funnel plots of two meta-analyses: On the left, the improvement in depression syndroms after application of tricyclic antidepressants is compared to placebo. The meta-analysis on the right measures the occurrence of intracranial haemoerrhage by CT after application of any anti-thrombolytic agent. All studies are RCT's.

to increase the JIF.

This then leads to an asymmetry in published studies; small studies show larger effect sizes than larger studies do. A funnel plot(cite sb) allows visual inspection of small study effects by plotting the effects against their standard error. For illustration purposes, one meta-analysis with large funnel plot asymmetry and one with no asymmetry is shown in Figure ??. By means of simple linear regression, one can investigate how much evidence there is for this asymmetry.

The purpose of a funnel plot is to visualise if an how the estimated treatment effects converge to a common "real" treatment effect value (here, the weighted mean treatment effect with weights = inverse variances). This is also shown by the triangle depicted by the dotted line (the spire is at the weighted mean).

We see that while the studies in the meta-analysis on the left are not symmetrically distributed, but rather accumulate at the left side, they seem to be more evenly distributed in the left triangle. From this, we would ultimately conclude that some sort of bias or heterogeneity is distorting the estimate of the overall treatment effect.

The Cochrane Organisation has specialized on systematic reviews of healthcare interventions. Researchers that write a systematic review collect data across studies, review them and try to provide up-to-date information about specific treatment efficacies (Higgins JPT, 2011). By extensive literature scrutinization, they try to circumvent the issue of publication bias. They do not apply funnel plot asymmetry tests to analyse publication bias. Earlier research however suggests that the efforts are only partially successful, and that there still is publication bias within the reviews (Egger et al. (1997), Ioannidis and Trikalinos (2007a), Kicinski et al. (2015), van Aert et al. (2019)). In these publications, Cochrane systematic reviews is analysed with methods to detect publication bias, for example small study effect tests or bayesian hierarchical selection models (Kicinski et al., 2015). They all find moderate to large evidence for publication bias in the Database. Their results will be compared to the results of this study in the last chapter.

## 1.1 Aim of the Study

None of the research so far has estimated the amount and impact of publication bias on metaanalytical findings thoroughly and with the most suitable methods. Also, the results are ironically often presented in the form of dichotomized hypothesis tests, a practice that is partly responsible for publication bias.

The aim of this thesis is to use prevailed methods to detect publication bias, and make use of the full amount of data that the Cochrane Organisation provides. At the end, an approximate, up-to-date estimate of the prevalence of publication bias in the data shall be given. To achieve this, methods to detect and adjust for publication bias in meta-analysis are applied on the data. It is also possible to adjust for publication bias with suitable methods. These methods are applied on the dataset as well, to achieve publication bias adjusted treatment effect estimates. It will be shown if and to what extent treatment effects are overestimated due to publication bias in the Cochrane systematic reviews.

## Chapter 2

## Methods

#### 2.1 Introduction and Notation

The analysis has already been said to be restricted on clinical or health care interventions. The interventions are restricted to comparisons of two treatment groups by some measure of melioration or worsening of health. The difference in this measure between the groups is referred to as the treatment effect. Where it is not particularly mentioned, the term treatment effect refers to any effect measure such as  $\log \operatorname{risk}$  ratio,  $\log \operatorname{hazard}$  ratio, Cohen's d or standardized mean difference, Fisher's z score or Pearsons correlation coefficient.

Let us consider a meta-analysis with n study treatment effects (n > 1, but typically small). A study is indexed by i, and it's treatment effect by  $\theta_i$ . The observed treatment effect is  $\hat{\theta}_i$ . The pooled treatment effect of a meta-analysis will be denoted as  $\theta_M$ , and consequently, the observed pooled treatment effect as  $\hat{\theta}_M$ . Furthermore, each treatment effect is typically measured with some standard error se<sub>i</sub> and an estimate of se<sub>i</sub> is denoted as  $\hat{\text{se}}_i$ . The  $\hat{\text{sign}}$  thus indicates if it is an estimate.

For continuous outcomes, let  $m_t$  be the mean of the treatment group,  $m_c$  the mean of the control group, and equivalently  $\mathrm{sd}_t$  and  $\mathrm{sd}_c$  the corresponding standard deviations.  $n_t$  and  $n_c$  are the total number of participants in the groups. In the case of binary outcomes, let  $e_t$  be the count of events in the treatment arm  $e_c$  the events in the control group. The observed counts in a study i are referred to as  $e_{t,i}$  and analogously  $e_{c,i}$ .

## 2.2 Effect Measures and p-values

#### 2.2.1 Continuous Outcomes

For given  $(m_t, m_c)$ ,  $(\mathrm{sd}_t, \mathrm{sd}_c)$  and  $(n_t, n_c)$ , one can compute mean difference as well as a standardized mean difference (here: Cohen's d) and a standard error thereof. The mean difference can  $\theta$  and its standard error se can be obtained as

$$\theta = m_t - m_c \qquad \text{se} = \sqrt{\operatorname{sd}_t^2/n_t + \operatorname{sd}_c^2/n_t}$$
 (2.1)

The standardized mean difference d and its standard error se can similarly be obtained by

$$se = \sqrt{\frac{(n_t - 1)sd_t^2 + (n_c - 1)sd_c^2}{n_t + n_c - 2}} \qquad g = \frac{m_t - m_c}{se}$$
 (2.2)

Both estimators take into account that the two groups might have unequal variances. A p-value to test the null hypothesis that the mean between group is equal is commonly obtained with the Students t test. The t statistic is obtained, using se and d from (2.2), by

$$t = d/(\text{se }\sqrt{(1/n_t) + (1/n_c)})$$

and the p-value can be obtained with the cumulative Student's t-distribution F with  $n_t + n_c - 2$  degrees of freedom:

$$p = 2(1 - F(|t|))$$

The t-test is known to be not very reliable if combined sample size is small (n < 30), see for example Kasuya (2001).

#### 2.2.2 Binary Outcomes

Two commonly used effect measures for binary outcome data are risk ratios and odds ratios between treatment and control group. The methods presented here can also be found, for example, in (Borenstein *et al.*, 2011, 34). Let  $\theta$  be the natural logarithm of the odds ratio.  $\theta$  and its variance se<sup>2</sup> can be obtained by

$$\hat{\theta} = \log(\frac{e_t \cdot (n_c - e_c)}{e_c \cdot (n_t - e_t)})$$

$$se^2 = 1/e_t + 1/(n_t - e_t) + 1/e_c + 1/(n_c - e_c)$$

Plugging in the observed counts will give the corresponding estimates. The logarithm of the risk ratio  $\theta$  and its variance se<sup>2</sup> is similarly defined as

$$\hat{\theta} = \log(\frac{e_t/n_t}{e_c/n_c}) \tag{2.3}$$

$$se^{2} = 1/e_{t} - 1/n_{t} + 1/e_{c} - 1/e_{t}$$
(2.4)

Using likelihood theory, one could show that the estimators are maximum likelihood estimators and that one can use the asymptotic normal distribution of the maximum likelihood estimator to calculate a *p*-value, *e. g.* (Held and Sabanés Bové, 2014, 98).

Thus, with  $\Phi$  as the cumulative standard normal distribution, we get

$$p = 2 \cdot (1 - \Phi(\left|\hat{\theta}/\hat{\operatorname{se}}\right|),$$

a p-value for the corresponding estimate, which summarizes the evidence against  $\hat{\theta}$  being zero (i. e. the true risk/odds ratio being 1).

Binary and continuous effect measures can be converted into each other, as described in Section 2.6.

#### 2.2.3 Survival Outcomes

Time-to-event data with censoring has to be analyzed by special means. One frequently used method to take into account right-censoring is the Cox proportional hazards regression model (Cox, 1972). Because the method itself is not applied in this thesis, but only the resulting estimates of the parameters are used, the reader is referred to the extensive literature covering this topic (e.g. Cox and Oakes (1984)).

The so-called hazard ratio estimated by Cox regression is the ratio of the instantaneous risk of experiencing the event between two groups. Because it is a maximum likelihood estimator, one

can again use its Wald test statistic to test for equal hazards. Let  $\hat{\theta}$  be an estimate of the log hazard ratio and se an estimate of the standard error of it. As before

$$p = 2 \cdot (1 - \phi(\left|\hat{\theta}/\hat{\operatorname{se}}(\hat{\theta})\right|)$$

will give a p-value for the evidence against the null hypothesis.

## 2.3 Fixed and Random Effects Meta-Analysis

The fixed effects meta-analysis estimator of the pooled treatment effect is a mean of the single treatment effect estimators, weighted by their standard errors (Rosenthal and Rubin, 1982). Let  $w_i = 1/\operatorname{se}_i^2$  be the weights, and  $\theta_M$  be the pooled estimator and  $s_M^2$  its variance. Then

$$\theta_M = \frac{\sum_{i=1}^n w_i \theta_i}{\sum_{i=1}^n w_i} \qquad s_M^2 = \frac{1}{\sum_{i=1}^n w_i}$$
 (2.5)

This estimator minimizes the variance between the effects. An estimate  $\hat{\theta}_M$  can be obtained by plugging in the observed treatment effects and variances  $\hat{\theta}_i$  and  $\hat{\text{se}}_i^2$ . The underlying idea is that we assume  $\theta_i \sim N(\theta_M, \text{se}_i^2)$ ,  $\theta_M$  being the true effect, all  $\theta_i$  being distributed around an equal mean.

The random effects model (Whitehead and Whitehead, 1991) assumes instead that

$$\theta_i \sim N(\mu_i, \mathrm{se}_i^2)$$
  $\mu_i \sim N(\theta_M, \tau^2)$  (2.6)

Marginally, we have  $\theta_i$  being distributed around a common mean  $\theta_M$  with additional variance  $\tau^2$ :

$$\theta_i \mid \mu_i \sim N(\theta_M, \mathrm{se}_i^2 + \tau^2)$$

 $\tau^2$  is often referred to as a population variance or between-study variance, whereas se<sup>2</sup><sub>i</sub> can be interpreted as sampling error. The pooled treatment effect estimate  $\theta_M$  of the random effects model and its variance is obtained by replacing the weights  $w_i$  in equation (2.5) with  $w_i = 1/(\text{se}_i^2 + \tau^2)$ .

The model is superior to the fixed effects model whenever the standard errors of the treatment effects alone are unlikely to fully account for the entire variability observed between studies. Note that as  $\tau^2$  increases, each  $\theta_i$  will eventually get equal weights, irrespective of it's sampling error se<sup>2</sup><sub>i</sub>.

The estimation of  $\tau^2$  has been subject to some debate in the statistical literature. Oftentimes, the method of moment estimator of DerSimonian and Laird (1986) is used. We use the measure of heterogeneity, Q, and divide by C after having subtracted the degrees of freedom n-1:

$$Q = \sum_{i=1}^{n} w_i (y_i - \theta_M)^2 \qquad C = \sum_{i=1}^{n} w_i - \frac{\sum_{i=1}^{n} w_i^2}{\sum_{i=1}^{n} w_i}$$
 (2.7)

$$\tau^2 = \max\left(0, \frac{Q - (n - 1)}{C}\right) \tag{2.8}$$

Again,  $w_i$ ,  $\theta_i$ ... have to be replaced by their estimates in order to get an estimate  $\hat{\tau}^2$ . The Paule-Mandel estimator (Paule and Mandel, 1982) is considered to have most often better properties than the method of moments estimator (e. g. Veroniki et al. (2016)). Since we defined  $w_i = 1/(\mathrm{se}_i^2 + \tau^2)$ , it also holds that

$$w_i \operatorname{Var}(\theta_i) = 1$$
  $\operatorname{Var}(\sqrt{w_i}\theta_i) = 1$ 

For any  $w_i$ , the variance can be estimated and equated to its expected value:

$$se^{2}(w_{i}\theta_{i}) = \frac{\sum_{i=1}^{n} w_{i}(\theta_{i} - \theta_{M})^{2}}{n-1} \qquad \frac{\sum_{i=1}^{n} w_{i}(\theta_{i} - \theta_{M})^{2}}{n-1} = 1$$
 (2.9)

 $\theta_M$  can be estimated using equation (2.5), the only problem is to estimate  $\tau^2$ . It can be obtained through an iterative process, using a newly defined function

$$F(\tau^2) = \sum_{i=1}^{n} w_i (\theta_i - \theta_M)^2 - (n-1)$$

In view of equation (2.9),  $\tau^2$  must be such that  $F(\tau^2) = 0$ . Then, we start with a arbitrary  $\tau^2$  and repeatedly add a term  $\tau_0^2$  to update  $\tau^2$  until  $F(\tau^2 + \tau_0^2)$  is close to zero (using  $\tau^2 + \tau_0^2$  for  $\hat{w}_i, \hat{\theta}_M$ ). Using a truncated Taylor series expansion, one can obtain the partial derivative after  $\tau^2$ , which is a reasonable choice for  $\tau_0^2$ .

The estimation of  $\tau^2$  is accompanied by uncertainty. A common procedure is to test if it is there is significant heterogeneity between the studies (Borenstein *et al.*, 2011, 109). Compute Q, as given in (2.7), based on one of the estimators of  $\tau^2$ . It is assumed that Q follows a central Chi-squared distribution with n-1 degrees of freedom under the null hypothesis of equally distributed effect sizes. Thus, the expected value of Q is n-1, and the excess dispersion is Q-n+1. The p-value against the null hypothesis of equally distributed effect sizes is 1-F(Q), using F as the cumulative distribution function of the Chi-squared distribution with d.f. = n-1. An advantage of the  $\tau^2$  is that it is directly linked to the variability in the data. Additionally, one can use the  $I^2$  statistic to see what portion the between study variance has of the overall variance. It is computed as

$$I^2 = \max\left(0, 1 - \frac{n-1}{Q}\right)$$

Importantly, all proposed methods above assume normally distributed effect sizes and proper estimates se of the true standard error. This assumptions are not met for very small sample sizes and very few event counts. Alternatively, the Mantel-Haenszel method for risk and odds ratios (see e.g. Fleiss et~al. (2013)) could be used in the latter case.

## 2.4 Small Study Effects Tests

The tests that will be presented on the following pages are a common way to detect publication bias. Importantly, they are however not interpretable directly as evidence for publication bias, but this is left for the discussion chapter. The tests are also often referred to as funnel plot asymmetry tests, because of the popularity of a recent test that has been used frequently to test and adjust for publication bias, which goes under the name of trim-and-fill (Duval and Tweedie, 2000). However, it is known for some time that the method has disadvantageous properties, therefore, it will not be discussed here, as well as the funnel plot (the radial plot will be used as an alternative).

A test for small study effects will test in some ways if the size of the estimated treatment effect of a study depends on some measure of its size. An association between study size and treatment effect size can be interpreted as an artifact of publication bias.

#### 2.4.1 Continuous Outcome Tests

For a continuous outcomes that are normally distributed, the sample mean and variance are independent of each other (Schwarzer et al., 2015, 120). Thus, the estimated standard errors of treatment effects can be used as a proxy for study size, as they should in principle not be tied to the effect size.

#### Begg and Mazumdar: Rank Correlation Test

Begg (1988) proposed a rank based test to test the null hypothesis of no correlation between effect size and variance. A standardized effect size  $\theta_i^*$  can be computed as in (2.10).  $se_i^{2*}$  is the variance of  $\theta_i - \theta_M$  as defined in (2.11),  $\theta_M$  being the fixed effects pooled treatment effect ((2.5).

$$\theta_i^{\star} = (\theta_i - \theta_M) / \operatorname{se}_i^{2\star} \tag{2.10}$$

$$se_i^{2\star} = se_i^2 - 1/\sum_{i=1}^n \frac{1}{se_i^2} 1$$
 (2.11)

A rank correlation test based on Kendall's tau is then used. First, the pairs are ordered after their ranks based on  $s^{2\star}$ . Then, for each  $s^{2\star}$  rank, the corresponding ranks based on  $\theta^{2\star}$  that are larger are counted and summed up to u. The number of ranks based on  $\theta^{2\star}$  that are in contrary, smaller, are counted and summed up to l. Then the normalized test statistic Z is given as

$$Z = (u-l)/\sqrt{n(n-1)(2n+5)/18}$$

Thus, large number of concordance between pairs will reflect in large  $\hat{u}$  and small  $\hat{l}$  and thus lead to a large  $\hat{Z}$ . The *p*-value is obtained using the standard normal distribution  $\Phi$ :

$$p = 2 \cdot (1 - \Phi(|Z|))$$

The changes that have to be made int the case of ties are small and can be found in (Begg, 1988, 410).

#### Egger's Test: Weighted Linear Regression Test

First, the concept of simple linear regression is introduced. In short, the model assumes a dependent variable y to be a linear function of another explanatory variable x:

$$y = \beta_0 + \beta_1 x + \epsilon, \qquad \epsilon \sim N(0, \sigma^2) \tag{2.12}$$

 $\epsilon$  is the residual noise term that becomes necessary when n pairs  $(x_i, y_i)$  are given and there is no exact solution. Then it is common to look for the solution that minimizes the squared residuals, the least-squares solution. Formally,

$$\underset{\beta_0,\beta_1}{\operatorname{argmin}} \left( \sum_{i=1}^n y_i - \beta_0 - \beta_1 x_i \right) \tag{2.13}$$

Let **X** be a matrix with the explanatory variables x and **y** a corresponding vector for all y:

$$\mathbf{X} = \begin{bmatrix} 1 & x_{12} \\ 1 & x_{22} \\ 1 & x_{32} \\ \vdots & \vdots \\ 1 & x_{n2} \end{bmatrix} \qquad \mathbf{y} = \begin{bmatrix} y_1 \\ y_2 \\ y_3 \\ \vdots \\ y_n \end{bmatrix}$$

Let  $\beta = (\beta_0, \beta_1)^{\top}$ . It can be shown that

$$\hat{\beta} = (\mathbf{X}^{\top} \mathbf{X})^{-1} \mathbf{X}^{\top} \mathbf{y} \tag{2.14}$$

Is an estimator of  $\beta$  that minimizes the squared residuals. Let  $\hat{\mathbf{y}} = \mathbf{X}\hat{\beta}$  and  $\hat{\mathbf{r}} = \hat{\mathbf{y}} - \mathbf{y}$ . The variance estimates  $\hat{\sigma}^2$  and  $\hat{\mathbf{s}}^2_{\beta}$  are

$$\hat{\sigma}^2 = \frac{1}{n-2} \mathbf{r}^{\mathsf{T}} \hat{\mathbf{r}} \qquad \hat{\mathbf{s}}_{\beta}^2 = \hat{\sigma}^2 (\mathbf{X}^{\mathsf{T}} \mathbf{X})^{-1}$$
 (2.15)

The estimate  $\hat{\beta}_0$  is the intercept and  $\hat{\beta}_1$  the slope of the regression line. Furthermore, in the simple linear regression setting,  $\hat{\beta}_0$  can also be obtained by:

$$\hat{\beta}_0 = \bar{y} - \hat{\beta}_1 \bar{x}$$

 $\bar{x}, \bar{y}$  denoting the sample means of the corresponding values  $x_1, ..., x_n$  and  $y_1, ..., y_n$ . Thus,  $\hat{\beta}_0$  is also called global mean. To test whether there is evidence for the intercept  $\beta_0$  to be unequal to some value  $\beta_{H0}$ , a t-test can be used.

$$p = 2(1 - F(|(\beta_0 - \beta_{H0})/s_{\beta_0}|))$$

where F is the cumulative t distribution with n-2 degrees of freedom. p will give the evidence against the null hypothesis  $\beta_0 = \beta_{H0}$ .

The concept is extendable to weighted linear regression. Weighted linear regression may be used if the residuals  $\mathbf{r}$  have unequal variances, which is equivalent to ascribe different precision to the observed y. The least squares equation (2.13) is extended to

$$\underset{\beta_0,\beta_1}{\operatorname{argmin}} (\sum_{i=1}^n w_i (y_i - \beta_0 - \beta_1 x_i))$$

with positive weights  $w_i$  that penalize large squared residuals for some i more if  $w_i$  is larger compared to other  $w_i$ .

Let **W** be a  $n \times n$  matrix with  $\mathbf{W}_{ii} = w_i$ , the weights on the diagonal and zeros on the off-diagonals. The estimates in (2.14) and (2.15) can still be used if **X** is exchanged with  $\mathbf{X}^* = \mathbf{W}\mathbf{X}$  and  $\mathbf{y}$  with  $\mathbf{y}^* = \mathbf{W}\mathbf{y}$ .

Now it is shown how linear regression can be used to test dependency of effect sizes on study sizes. The simplest application was introduced by Egger et al. (1997). Let  $\theta/s$  be the dependent variable y and 1/s the explanatory variable x. If plotted, this corresponds to a radial or Galbraith plot (Galbraith, 1988). The linear regression equation as introduced before in (2.12) can be written in two ways:

$$\theta/s = \beta_0 + \beta_1/\operatorname{se} + \epsilon,$$
  $\epsilon \sim N(0, \sigma^2)$  (2.16)

Equation (2.16) is often provided due to the correspondence to the radial plot. However, it is equivalent to

$$\theta = \beta_0 + \beta_1 \operatorname{se} + \epsilon,$$

$$\epsilon \sim N(0, w^{-2}\sigma^2) \tag{2.17}$$

with weights  $w = 1/\operatorname{se}^2$ . Thus testing  $\beta_0$  of (2.16) or  $\beta_1$  is equivalent. The corresponding p-value is then used as evidence for a small study effect. Plugging in  $\theta_i/\operatorname{se}_i$ ,  $1/\operatorname{se}_i$  as  $y_i, x_i$  into equation (2.14) and (2.15) will give the estimates for  $\hat{\beta}_0, \hat{\beta}_1, \hat{\operatorname{se}}_{\beta_0}$  and  $\hat{\operatorname{se}}_{\beta_1}$ .

#### Thompson and Sharp's Test: Weighted Linear Regression Random Effects Test

A method proposed in Thompson and Sharp (1999) allows for between study variance  $\tau^2$ , as introduced before in section 2.3. It extends the previously seen linear regression approach with x = 1/se and  $y = \theta/s$  by introducing weights. The effect size  $\theta_i$  is assumed to be distributed as

$$\theta_i \sim N(\beta_0 + \beta_1 \operatorname{se}_i, \operatorname{se}_i^2 + \tau^2) \tag{2.18}$$

 $\tau^2$  is estimated as in equation (2.8) (method of moments). The weights are set as  $w_i = 1/\sqrt{\mathrm{se}_i^2 + \tau^2}$ ). After adjusting for the weights as described in 2.4.1, we can proceed analogous to Egger's test. The *p*-value for  $\beta_0 \neq 0$  reflects the evidence for a small study effect.

#### 2.4.2 Generalized linear model

The introduction of group-sepific random effects allows to analyze grouped or repeated measurements by linear regression (McCullagh and Nelder, 1989). Let i be the group index, and

$$\mathbf{Y}_{ij}|U_i, \epsilon_{ij} = (1, x_i^{\top})\beta + U_i + \epsilon_{ij} \tag{2.19}$$

the marginal distribution of  $\mathbf{Y}_{ij}$  depending on  $U_i, \epsilon_{ij}$ . The random effects  $U_i \sim N(0, g)$  and the residual error term  $\epsilon_{ij} \sim N(0, \tau^2)$  are considered independent. If there are more groups in the data we can extend (2.19) by adding a second random term Q with an index k:

$$\mathbf{Y}_{ij}|U_i, Q_k \epsilon_{ij} = (1, x_i^\top)\beta + U_i + Q_k + \epsilon_{ij}$$
(2.20)

Again,  $U, K, \epsilon$  are distributed independently. The random intercepts model will account for a nested group structure by accordingly indexing the groups.

The conditional distribution in equations (2.19) and (2.20) can be used to construct a conditional likelihood; A bayesian posterior distribution is integrated with respect to the random effects, such that it can be used for restricted or unrestricted maximum likelihood estimation.

#### 2.4.3 Dichotomous Outcomes Tests

The issue with dichotomous outcomes is that effect size and variance of effect size are correlated, which can readily be seen in (2.3) and (2.4). For example, a small number of event counts in one or group will inflate the variance and the effect size. Consequently, the tests above will tend to reject the null-hypothesis too often, *i. e.* report false positives. A number of solutions to this problem are provided in the literature.

#### Peters Test: Weighted Linear Regression Test

Instead of taking the standard error s as explanatory variable x as in Egger's Test, the inverse of the total sample size is used. Additionally, the variances  $se_i^2$  are used as weights. Thus, the subsequent test procedure is identical to Egger's test. Peters test is a small modification of Macaskill's test where the explanatory variable is the sample size instead of its inverse.

#### Harbord's Test: Score based Test

A rank based alternative to Peters test for binary outcomes is Harbord's test (Harbord *et al.*, 2006). It uses a different treatment effect and variance estimate: the score  $\varphi$  of the log-likelihood, evaluated at log odds ratio  $\theta_0 = 0$  and its inverse Fisher information  $s^2$ . Formally,

$$\varphi = e_t - (e_t - e_c)(e_t + (n_t - e_t))/(n_t + n_c)$$
(2.21)

$$s^{2} = \frac{(e_{t} + e_{c})(e_{t} + (n_{t} - e_{t}))(e_{c} + (n_{c} - e_{c}))((n_{t} - e_{t}) + (n_{c} - e_{c}))}{(n_{t} + n_{c})^{2}(n_{t} + n_{c} - 1)}$$
(2.22)

It can be shown that they are both good approximations of the log odds ratio and its variance if the real  $\theta$  is not too far from zero. The standardized estimator  $r_i/v_i$  is also known as Peto odds ratio. The obtained scores and variances can be used in Egger's test as treatment effects and variances.

#### Schwarzer's Test: Rank Correlation Test

Schwarzer et al. (2007) developed a test for the correlation between  $E_t - \mathbb{E}(E_t)$  and the variance of  $E_t$ ,  $E_t$  being a random variable from the non-central hypergeometric distribution, assuming a fixed log odds ratio.

 $\mathbb{E}(E_t)$  and variance of  $E_t$  are then estimated based on  $e_t$ . The standardized cell count deviation

$$(e_t - \mathbb{E}(E_t)) / \sqrt{(\operatorname{se}_i^2)}$$
 (2.23)

and the inverse of  $se_i^2$  is then used as before in Begg and Mazumdar's test.

# Rücker's Test: Using the Variance Stabilizing Transformation for Binomial Random Variables

The correlation between variance and effect size of dichotomous outcome measures can be abolished by the variance stabilizing transformation for binomial random variables. We use that the arcsine function is the variance stabilizing transformation for a proportion. Let

$$\theta_i = \arcsin e_t/n_t - \arcsin e_c/n_c$$
  $\operatorname{se}_i^2 = 1/4n_t + /4n_c$ 

Then one can optionally apply Begg and Mazumdar's rank correlation test or Thompson and Sharp's test using the newly obtained estimates.

#### 2.4.4 Excess Significance Test

Ioannidis and Trikalinos (2007b) developed an exploratory test to detect if the proportion of significant findings is larger than expected. Note that significance is here defined as one-sided significance, with the direction of the test being the same for each study.

We assume that the effects are equally distributed around a true mean effect  $\theta_M$ , which can be estimated by fixed effects Meta-Analysis (2.5). Let O be the number of significant study results out of n studies and  $\alpha$  the significance threshold. Corresponding to the study effect  $\theta_i$ , we can specify the power  $1 - \beta_i$ , the probability to accepting a true result. Let  $z_{\alpha,i}$  be the  $1 - \alpha$  quantile of a normal distribution with standard error  $\hat{se}_i$ . The power of study i can be estimated as:

$$1 - \hat{\beta}_i = F(z_\alpha) \tag{2.24}$$

with F being the cumulative distribution of a normal with mean  $\hat{\theta}_M$  and standard deviation  $\hat{\mathbf{se}}_i$ .

If we assume no bias in  $\theta_i$  and  $\theta_M$ , the expected number of significant study results is then just

$$E = \sum_{i=1}^{n} (1 - \beta_i)$$

E can then be compared to O by constructing a test statistic  $\chi$ :

$$\chi = \left(\frac{(O-E)^2}{E} + \frac{(O-E)^2}{n-E}\right)$$

and consecutively, calculating a p-value for the evidence against the null-hypothesis of O = E with a  $\chi^2$  distribution with one degree of freedom. Alternatively, one can also use a binomial test, which is encouraged when n and O is small. We will get a one sided p-value for excess significance,  $\Pr(X \ge O)$ , by

$$p = \sum_{i=0}^{n} \left( \binom{n}{i} p^{i} (1-p)^{n-i} \right)$$
 (2.25)

with p = E/n and X being a binomial random variable with probability p.

## 2.5 Small Study Effect and Publication Bias Adjustment

There are different approaches to correct for small study effects and publication bias. They can mainly be distinguished by their underlying methods: regression based approaches aim to regress the effect to a study with infinite precision (*i. e.* very small standard error) or to a summary effect, corrected for publication bias. Selection models aim to simulate different, hypothetical selection processes and attain a approximate treatment effect by sensitivity analysis:

#### 2.5.1 Adjustment by Regression

Rücker et al. (2011) use a random effects model to obtain an unbiased estimate. Similarly to regression based tests for small study effects, we have

$$\theta_i = \beta_0 + \beta_1 \sqrt{\operatorname{se}_i^2 + \tau^2} + \epsilon_i \sqrt{v_i + \tau^2}, \epsilon_i \stackrel{\text{iid}}{\sim} N(0, 1)$$
(2.26)

The only difference between Thompson and Sharp's variant is  $x = \sqrt{\sec^2 + \tau^2}$  instead of  $x = \sqrt{\sec^2}$ .  $\beta_1$  represents the bias introduced by small study effects, as illustrated in the following equations:

$$\mathbb{E}((\theta_i - \beta_0)/\sqrt{\mathrm{se}_i^2}) \to \beta_1 \text{ if } \mathrm{se}_i \to \infty$$

$$\mathbb{E}(theta_i) \to \beta_0 + \beta_1 \tau \mathrm{if } \mathrm{se}_i \to 0$$

After estimating  $\tau^2$ , one can estimate  $\beta_0$  and  $\beta_1$  as seen before in the simple linear regression framework. Now we have basically two possible estimates at hand:

- $\beta_0$  the treatment effect without any influence of study precision with standard error  $s_{\beta_0}$
- $\beta_0 + \beta_1 \tau$  the treatment effect of a hypothetical study with infinite precision, corresponding standard error se =  $s_{\beta_0} + s_{\beta_1}$

Simulations in Rücker *et al.* (2011) suggested that the latter estimate is slightly superior with respect to coverage (and mean squared error).

#### 2.5.2 Copas Selection Model

A method proposed in Copas and Shi (2001, 2000); Copas and Malley (2008) assumes that the given sample of treatment effects and standard errors is a selected part of a larger random sample. Selection of studies depends on their effect size and variance. Smaller variance is always accompanied by larger selection probability.

Let  $\theta_i$  be the effect size estimate of study i. Then

$$\theta_i \sim N(\mu_i, \sigma_i^2)\mu_i \sim N(\theta, \tau^2)$$
 (2.27)

which is similar to the random-effects meta-analysis setting.  $\theta$  is the population mean effect,  $\sigma_i^2$  the within study variance and  $\tau^2$  the between study variance. Equations in (2.27) are termed the population model.

The selection model is defined as follows. Suppose a selection of studies with reported ( $\neq$  estimated) standard errors s (likely different from  $\sigma$ ). Only a proportion of the selection will be published, with a defining the overall proportion of published studies and b (assumed to be positive) defining how fast this proportion increases with s becoming smaller. Formally,

$$P(\text{select} \mid s) = \Phi(a + b/s)$$

The equation can be rewritten as

$$z = a + b/s + \delta$$

with  $\delta \sim N(0,1)$ . z is interpreted as the *propensity for selection*. It's sign must be positive in order for the study to be selected. Thus, the larger z, the more likely the study will be selected. Also, a is somewhat like to global retention or selection rate for each study, while b decides about the decline of selection probability with increasing s.

So far, we have, for a study i

$$\theta_i = \mu_i + \sigma_i \epsilon_i$$
$$\mu_i \sim N(\theta, \tau^2)$$
$$z_i = a + b/\operatorname{se}_i + \delta_i$$

where  $(\epsilon_i, \delta_i)$  are standard normal residuals. The two models are coupled by introducing a correlation  $\rho = cor(\theta_i, z_i)$  by defining  $(\epsilon_i, \delta_i)$  as bivariate standard normals. It follows that, if  $\rho_i$  is large and positive and  $z_i > 0$ , then the estimate of a study i that is selected is likely to have positive  $\epsilon_i$  and  $\delta_i$  and thus, the true mean  $\mu$  is likely to be overestimated.

Let  $u_i = a + b/\operatorname{se}_i$ ,  $\lambda(u_i)$  the Mill's ratio  $\phi(u_i)/\Phi(u_i)$  ( $\phi$  is the standard normal density function and  $\Phi$  the cdf) and  $\tilde{\rho}_i = \sigma/\sqrt{(\tau^2 + \sigma_i^2)\rho_i}$ . The probability of a study being selected, given  $\operatorname{se}_i$  and  $\theta_i$ , is

$$P(\text{select} \mid \text{se}_i, \theta_i) = \Phi\left(\frac{u_i + \tilde{\rho}_i((\theta_i - \mu)/\sqrt{(\tau^2 + \sigma_i^2)})}{\sqrt{1 - \tilde{\rho}_i^2}}\right)$$

Which again shows that larger  $se_i$  and  $\theta_i$  lead to a larger selection probability. It can also be shown that the expected value

$$\mathbb{E}(\theta_i \mid \text{se}_i, \text{select}) = \mu + \rho_i \sigma_i \lambda(u_i)$$
 (2.28)

which shows that the expected value for a study is larger for larger  $\sigma$ .

A likelihood for  $\theta_i$ , conditional on z > 0 can be formulated to estimate the parameters of the model. Regarding a and b, there is no way that they can be estimated because the number of missing studies and their effect sizes is not known. Instead, fixed values for a and b have to be chosen. The nuisance parameter  $\sigma_i$  can be replaced by an estimate if the sample size in the studies is large enough. Since

$$Var(\theta_i \mid se_i, z_i > 0) = \sigma_i^2 (1 - c_i^2 \rho_i^2)$$

with  $c^2 = \lambda(u_i)(u_i + \lambda(u_i))$ , we can replace  $\sigma_i^2$  by  $\hat{\sigma}_i^2 = \frac{1}{1 - c_i^2 \rho_i^2}$ . Although one has to evaluate the likelihood for fixed pairs (a, b), one can compare the fit of the model to evaluate which one is more suitable: With equation (2.28), one can obtain fitted values of  $\theta_i$  based on se<sub>i</sub>. Also, for two different pairs (a, b),  $(a^*, b^*)$ ,

$$\mathbb{E}(\theta_i \mid z_i > 0, a^*, b^*) - \mathbb{E}(\theta_i \mid z_i > 0, a, b) \approx c^* + \rho(\lambda(a^*) - \lambda(a)) \operatorname{se}_i$$

and that local departures of  $\theta_i$  can be approximated by adding a linear term in se<sub>i</sub> to the expectation of  $\theta_i$ . Thus, to test a pair (a, b), it is sufficient to test  $\beta \neq 0$  in

$$\theta_i = \theta + \beta \operatorname{se}_i + \sigma_i \epsilon_i$$

with restriction that  $\rho \geq 0$ . To test some pair (a,b) against the scenario with no selection, we set  $a^* = \infty$  (or  $\rho = 0$ ). A likelihood ratio test will give a test statistic to test against H0 = no selection:

$$\chi^2 = 2 \cdot (\max_{\theta, \tau, \beta} \tilde{L}(\theta, \tau, \beta) - \max_{\theta, \tau} \tilde{L}(\theta, \tau, 0))$$
(2.29)

with

$$\tilde{L}(\theta, \tau, \beta) = -\frac{1}{2} \sum_{i=1}^{n} [\log(\tau^2 + \sigma_i^2) + \frac{(\theta_i - \theta - \beta se_i)^2}{(\tau^2 + \sigma_i^2)}]$$

 $\chi^2$  can be used with a  $\chi^2$  distribution with one degree of freedom to obtain a *p*-value. Note that the test is almost equivalent to Egger's small study effect test with  $\tau^2 = 0$ . Thus, although the copas selection model models publication bias, it is dependent on the small study effects to find the most suitable pair (a, b).

If one applies the model to a single meta-analysis, a sensitivity analysis is suggested. One can observe how  $\theta$  and it's confidence intervals change dependent on the underlying selection process. Selection models are in general not recommended for inference (e. g. McShane et al. (2016)). Rücker et al. (2011) have shown how the method can be implemented in a simulation for inference purposes.

A range of values of (a, b) are applied, and the test for residual small study effect as described in equation (2.29) is applied. If all obtained p-values from the test are above a threshold 0.1, this is interpreted as no evidence, and no need for modelling, and the standard, classical random effects meta-analysis is retained. If none of the p-values is above the threshold, a wider range of values for (a, b) is used. When some p-values are above, and some below the threshold, the pair (a, b) with the smallest number of missing studies is retained (that is, the least intense underlying selection model is chosen).

Currently, there is no test to detect miss-specifications in the model itself, the authors themselves have argued that a non-parametric test of the residuals would lack power.

### 2.6 Transformation between Effect Measures

Assuming that binary outcomes result from a dichotomization of originally continuous random variables, in this case, the logistic distribution, a transformation from typical binary effect measures to Cohen's d can be achieved (Borenstein et al., 2011, 47).

Let  $\theta$  be a log odds ratio and s it's standard error. Cohen's d and it's variance  $s_d^2$  is obtained by

$$d = \theta \frac{\sqrt{3}}{\pi} \qquad \qquad s_d^2 = \sec^2 \frac{\sqrt{3}}{\pi}$$

The factor  $\frac{\pi}{\sqrt{3}} = 1.81$  is the standard deviation of the logistic distribution  $L(\mu, \eta)$  with scale parameter  $\eta = 1$ , so we just divide the log odds ratio and it's variance through the standard deviation. The approximation works only well if  $e_t$  and  $e_c$  are not very small, especially in the case of  $s_d^2$ . Plugging in the observed log odds ratio  $\hat{\theta}$  and  $\hat{se}^2$  will give an estimate of Cohen's d. Pearson's correlation can be attained by the formulas (Hedges and Olkin (1985), (Borenstein et al., 2011, 48))

$$r = \frac{d}{\sqrt{d^2 + a}} \qquad \qquad a = (n_c + n_t)^2 / n_c n_t$$

where a is a correction factor if  $n_t \neq n_c$ . The variance of r,  $s_r^2$  is computed by

$$s_r^2 = \frac{a^2 s_d^2}{(d^2 + a)^3}$$

Finally, we can get to a fisher's z-scaled correlation z and it's variance  $s_z^2$  by using

$$z = 0.5 \ln \left( \frac{1+r}{1-r} \right)$$
  $s_z^2 = \frac{1}{n-3}$ 

## Chapter 3

# The Cochrane Dataset

## 3.1 Cochrane Systematic Reviews

The Cochrane Group has specialized on systematic reviews in clinical science. Certain knowledge of standards and principles of the Cochrane Group may help to assess the quality and the properties of the dataset. The following information stems from the Cochrane Handbook for Systematic Reviews (Higgins JPT, 2011).

The definition of a systematic review is that it "attempts to collate all empirical evidence that fits pre-specified eligibility criteria in order to answer a specific research question." Thus, the "key properties of a review are":

- "a clearly stated set of objectives with pre-defined eligibility criteria for studies"
- "an explicit, reproducible methodology"
- "a systematic search that attempts to identify all studies that would meet the eligibility criteria"
- "an assessment of the validity of the findings of the included studies, for example through the assessment of risk of bias"

At the end of a systematic review, "a systematic presentation, and synthesis, of the characteristics and findings of the included studies" is done.

53 Cochrane Review Groups prepare and maintain the reviews within specific areas of health care. A group consists of "researchers, healthcare professionals and people using healthcare services (consumers)".

The groups are supported by Method Groups, Centers and Fields. The Cochrane Method Groups aim to discuss and consult the groups in methodological questions concerning review preparation. The Centers play a main role in training and support of the Groups. The Fields are responsible for broad medical research areas and follow priorities in those areas by advice and control of the groups.

The first step in a review is writing a protocol, specifying the research question, the methods to be used in literature search and analysis and the eligibility criteria of the study. Changes in protocols are possible but have to be documented and the protocol is published in advance of the publication of the full review. The choices of methodology as well as the changes should not be made "on the basis of how they affect the outcome of the research study".

In order to avoid potential conflicts of interests, there is a code of conduct that all entities of the Cochrane Organization have to agree on: conflicts of interest must be disclosed and possibly be forwarded to the Cochrane Center, and participation of review authors in the studies used have to be acknowledged. Additionally, a Steering Group publishes a report of potential conflicts of interests based on information about external funding of Cochrane Groups.

In order for keeping the reviews up-to-date, they are revised in a two-year circle with exceptions. In addition to inclusion of new evidence in a field, the revision and maintenance process may as well includes change in analysis methods. This can reflect some advance in clinical science as for example new information about important subgroups, as well as new methods for conducting a Cochrane Review. However, there are no clear guidelines and the Cochrane Groups are free in the rate and extent of up-dating their reviews.

#### 3.1.1 Methods for Cochrane Reviews

A research question defines the following points: "the types of population (participants), types of interventions (and comparisons), and the types of outcomes that are of interest". From the research question, usually the eligibility criteria follow. Usually, outcomes are not part of eligibility criteria, except for special cases such as adverse effect reviews.

The type of study is an important eligibility criterium. The Cochrane Collaboration focuses "primarily on randomized controlled trials", and also, the methods of study identification in literature search are focused on randomized trials. Furthermore, study characteristics such as blinding of study operators with respect to treatment and cluster-randomizing might be additional eligibility criteria which have to be chosen by the review authors.

After having specified the eligibility criteria, studies have to be collected. The central idea of systematic reviews, and also meta-analyses, is that the collected studies are a random sample of a population of studies, i.e. that they are representative and can be used to assess population properties. Therefore, the search process is crucial, as a selective search result may impose bias on the sample of studies available, making it a non-random sample. For this purpose, the Cochrane Groups are advised to go beyond MEDLINE !!cite!!, because a search restricted to it has been shown to deliver only 30% to 80% of available studies. "Time and budget restraints require the review author to balance the thoroughness of the search with efficiency in use of time and funds and the best way of achieving this balance is to be aware of, and try to minimize, the biases such as publication bias and language bias that can result from restricting searches in different ways." It is important to note that not only studies, but also study reports are occasionally used in the reviews, as they may provide useful information.

There are different sources that are being used to search for studies.

- The Cochrane Central Register of Controlled Trials is a source of reports of controlled trials. "As of January 2008 (Issue 1, 2008), CENTRAL contains nearly 530,000 citations to reports of trials and other studies potentially eligible for inclusion in Cochrane reviews, of which 310,000 trial reports are from MEDLINE, 50,000 additional trial reports are from EMBASE and the remaining 170,000 are from other sources such as other databases and handsearching." It includes citations published in many languages, citations only available in conference proceedings, citations from trials registers and trials results registers.
- MEDLINE. MEDLINE includes over 16 million references to journal articles. 5,200 journals
  publishing in 27 languages are indexed for MEDLINE. PubMed gives access to a free version
  of MEDLINE with up-to-date citations. NLM gateway such as the Health Services Research
  Project, Meeting Abstracts and TOXLINE Subset for toxicology citations allows for search
  in both databases together with additional data from the US National Library of Medicine.
- EMBASE. 4,800 Journals publishing in 30 languages are indexed to EMBASE, which includes more than 11 million records from 1974 onward. EMBASE.com also includes 7 million unique records from MEDLINE (1966 up to date) together with its own records. Additionally, EMBASE Classic allows access to digitized records from 1947 to 1973. EMBASE and MEDLINE each have around 1,800 journals not indexed in the other database.
- Regional or national and subject specific databases can additionally be consulted and

often provide important information. Financial considerations may limit the use of such databases.

- General search engines such as Google Scholar, Intute and Turning Research into Practice (TRIP) database can be used.
- Citation Indexes. The database lists articles published in around 6,000 Journals with articles in which they have been cited and is available online as SciSearch. This form of search is known as cited reference searching.
- Dissertation sources. Dissertations are often listed in MEDLINE or EMBASE but one is advised to also search in specific dissertation sources.
- Grey Literature Databases. Approximately 10% of the results in the Cochrane Database stems from conference abstracts and other grey literature. The Institute for Scientific and Technical Information in France provides access to entries of the previously closed System for Information on Grey Literature database of the European Association for Grey Literature Exploitation). Another source is the Healthcare Management Information Consortium (HMIC) database containing records from the Library and Information Services department of the Department of Health (DH) in England and the King's Fund Information and Library Service. The National Technical Information Service (NTIS) gives access to the results of US and non-US government-sponsored research, as well as technical report for most published results. References from newsletters, magazines and technical and annual reports in behavioral science, psychology and health are provided in the PsycEXTRA database which is linked to PsycINFO database.

#### 3.1.2 Structure and Content

The dataset consists of 6354 systematic reviews from the Cochrane Library with 70662 studies and 744720 results. The results all stem from comparisons of a healthcare interventions or treatments to a reference. The reference may be a placebo control group or a other intervention or treatment. A result can not only be about efficacy of the treatments, but also about safety (adverse effects).

In Table 3.1, two results from a systematic review about barbiturates are shown as they are given in the dataset. As can be seen, further specifications are provided by the variables in the columns.

The comparison.name variable specifies what kind of treatments or interventions are compared, the outcome.name variable how it is compared, and the subgroup.name variable (not indicated in table) if and to what experimental subgroup the result belongs.

The result is of a binary type, and the counts of events in the treatment group are in events1 and of the control group in events2 and the total number of participants are given in columns total1 and total2. As can be seen, events denote here "death at the end of follow-up".

study.name	comparison.name	outcome.name	events1	total1	events2	total2
Bohn 1989	Barbiturate vs no barbiturate	Death at the end of follow-up	11	41	11	41
Ward 1985	Barbiturate vs no barbiturate	Death at the end of follow-up	14	27	13	26

**Table 3.1:** Example of two results as given in the dataset. Events denotes the count of events in the treatment group while Events c the count of events in the group compared to. Further descriptive variables have been ommitted

A complete listing of the variables of a result is given in Table 3.2. Depending on the type of the data, e. g. if it is binary or continuous, some variables are missing for the specific results.

Results are part of studies that are again part of a (systematic) review. This structure of a review is shown in Figure ??.

Variable	Description
id	An id of the review for identification purposes
study.name	Name of the study to which the result belongs
study.year	Year in which the study was published
study.data.source	Source of the study, either "Publ
comparison.name/.nr	Specification of the interventions compared in the study
	and a unique number for the comparison
outcome.name/.nr	Specification by which outcome the interventions are compared
	and a unique number for the outcome
subgroup.name/.nr	Potentially indication of affiliation to subgroups and a
	unique number for the subgroup
outcome.measure	Indication of the quantification method of the effect
	(of one intervention compared to the other).
effect	Measure of the effect given in the quantity denoted by
	outcome measure.
se	Standard error of the measure of the effect,
events1/events2	The counts of patients with an outcome $if$
	measurement/outcome is binary or dichotomous
	2 (1 for treatment group and 2 for control group).
total1/total2	Number of patients in groups.
mean1/mean2	Mean of patient measurements $if$ outcome is continuous.
sd1/sd2	Standard deviation of mean if
	outcome is continuous.

**Table 3.2:** Dataset variable names and descriptions

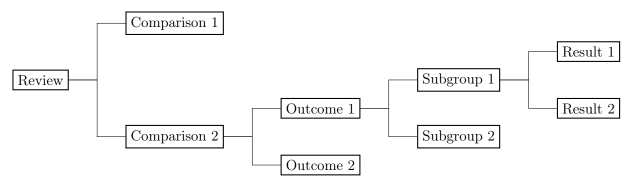


Figure 3.1: Structure of a hypothetical review with two different comparisons

The structure of a review will now be outlined based on an example of the dataset. Let us consider the previously mentioned barbiturate and head injury review. The aim was to "assess the effects of barbiturates in reducing mortality, disability and raised ICP (intra-cranial pressure) in people with acute traumatic brain injury" as well as to "quantify any side effects resulting from the use of barbiturates" The review comprises five studies in total. Three of them compared barbiturate to placebo, one compared barbiturate to Mannitol and one Pentobarbital to Thiopental. The studies have different outcomes, for example, death or death and severe disability at follow up, but also dropout counts or adverse effects (secondary outcomes). We have continuous (e.g. mean body temperature) and binary outcome data (e.g. death/no death). One study split up outcomes for patients with and without haematoma, which would be subgroups. Thus, it is important not to confuse results with studies. A study can contribute multiple results to a systematic review, for example, primary and secondary outcomes and adverse effects.

study.name	comparison.name	outcome.name
Bohn 1989	Barbiturate vs no barbiturate	Death at the end of follow-up
Bohn 1989	Barbiturate vs no barbiturate	Death or severe disability at the end of follow-up
Eisenberg 1988	Barbiturate vs no barbiturate	Death at the end of follow-up
Eisenberg 1988	Barbiturate vs no barbiturate	Uncontrolled ICP during treatment
Eisenberg 1988	Barbiturate vs no barbiturate	Hypotension during treatment
Perez-Barcena 2008	Pentobarbital vs Thiopental	Death at the end of follow-up (6 months)
Perez-Barcena 2008	Pentobarbital vs Thiopental	Death or severe disability at the end of follow-up (6 months)
Perez-Barcena 2008	Pentobarbital vs Thiopental	Uncontrolled ICP during treatment
Perez-Barcena 2008	Pentobarbital vs Thiopental	Hypotension during treatment
Schwartz 1984	Barbiturate vs Mannitol	Death at the end of follow-up (1 year)
Schwartz 1984	Barbiturate vs Mannitol	Death at the end of follow-up (1 year)
Schwartz 1984	Barbiturate vs Mannitol	Uncontrolled ICP during treatment
Ward 1985	Barbiturate vs no barbiturate	Death at the end of follow-up
Ward 1985	Barbiturate vs no barbiturate	Death or severe disability at the end of follow-up
Ward 1985	Barbiturate vs no barbiturate	Mean ICP during treatment
Ward 1985	Barbiturate vs no barbiturate	Mean arterial pressure during treatment
Ward 1985	Barbiturate vs no barbiturate	Hypotension during treatment
Ward 1985	Barbiturate vs no barbiturate	Mean body temperature during treatment

**Table 3.3:** Barbiturate and head injury review. In the columns, study names, comparison and outcome measure of the results are given

Information about missing values in the dataset is given in Table 3.4. The relative amount of missing values is low, except for study years. For continuous outcomes, the cases are counted were neither a mean difference nor means are available. Similarly, the counts of cases where neither standard errors nor standard deviations are available are provided. Study years before 1920 and after 2019 are declared as missing, as well as sample sizes equal to zero.

Neither means nor standard deviations (CONT)	775
Zero participants in one group	15162
Missing study publication year	7942

Table 3.4: Number of missing variables and measurements in the dataset

The studies that are included in the reviews and have been published are most often from the years after 1980 (5% quantile = 1982, 95% quantile = 2014,). The median of the publication years is 2003, the mean 2000.97 and the quartiles are 1996 and 2008. 1075 studies have been published in 2018, none in 2019.

The top treatment effect measure (risk ratio, mean difference, hazard ratio etc.) abundances are summarized in Table 3.5. One can conclude of the table that roughly 31 % of outcomes in the dataset are continuous and the rest being some sort of discrete or binary outcomes, most often binary (more than 65%).

The sample sizes among results vary to some extent. There are 5% of treatment group sample sizes that are smaller than 9, 95% smaller than 510. The first quartile is 23, the median 48, the mean 256.71 and the third quartile 116. The large difference between median and mean is caused by very large groups with over 2,000,000 participants. Analogously, the quantiles of the total sample size are: 5% quantile = 17, first quartile = 44, median = 94, third quartile = 223 and 95% = 983. The mean is 623.15.

There are 519 reviews with five or fewer results. The 5% and 95% quantiles are 4 and 447. The mean and median number of results per review are 13.6 and 8, and the quartiles are 16 and 109. Similarly, the number of reviews with a maximum of two studies included is 1040, the mean study number is 13.6, the median 8 and the interquartile range 4 and 16 and the 95% quantile 45. The discrepancy between mean and median is due to large reviews with a high number of studies and results, most extreme in which is a systematic review about antibiotic prophylaxis for preventing infection after cesarean section, with 95 studies and 1,497 results in total.

Outcome measure	n	Percentage
RR	361902	48.6%
MD	164923	22.1%
OR	76067	10.2%
SMD	70717	9.5%
PETO OR	39710	5.3%
RD _	11068	1.5%
Hazard Ratio	8054	1.1%
Rate Ratio	3724	0.5%
other	8555	1.1%
OR SMD PETO_OR RD Hazard Ratio Rate Ratio	76067 70717 39710 11068 8054 3724	10.2% 9.5% 5.3% 1.5% 1.1% 0.5%

**Table 3.5:** Frequencies of outcome measures among results. n denotes the total number of results with the outcome measure and percentage the percentage of the outcome measure,

For results to be suitable for usage in meta-analysis, they have to be identical with respect to comparison and outcome. The studies in the dataset that have the same comparison, outcome and subgroup can be pooled in a meta-analysis. This distinction is also used by the Cochrane Organization itself, *i. e.* the meta-analyses are identical to the meta-analyses done in the systematic reviews.

The size of a meta-analysis denotes how many results are included in a group. Table 3.6 shows the number of meta-analysis with size  $\geq n$  results.

## Warning: Setting row names on a tibble is deprecated.

n	Number of groups	Cumulative sum of groups
1	143378	268906
2	45459	125528
3	23232	80069
4	14184	56837
5	9493	42653
6	6449	33160
7	4583	26711
8	3412	22128
9	2585	18716
10	2046	16131
11	1524	14085
12	1197	12561
13	1022	11364
14	785	10342
15	9557	9557

Table 3.6: Number and cumulative number of groups with meta-analysis size n.

## 3.2 Data Tidying and Processing

The dataset was scraped from the Cochrane Database of Systematic Reviews at the  $\dots$  . A specialized code was used to convert the xtml. files from the webpage into the data at hand cite... .

#### 3.2.1 Newly Introduced Variables

Some new variables are added to the obtained dataset:

• outcome.measure.merged: Outcome measure specifications given in outcome.measure were standardized whenever they were supposed to denote the same outcome measure.

An example would be the formally different notations for odds ratios: "Odds Ratio", "odds ratio", "OR", which were all denoted as "Odds Ratio".

- lrr and var.lrr: log risk ratio and variance of the log risk ratio for outcome.flag "bin".
- smd and var.smd: Hedges q and the variance of Hedges q for outcome.flag "cont".
- smd.ordl and var.smd.ordl: Cohen's d and its variance as obtained by transformation of a log odds ratio for outcome.flag "bin".
- cor.Pearson and var.cor.Pearson: Pearson correlation coefficient and variance as obtained from the d (for outcome.flag "bin") or g (for outcome.flag "cont") to r transformation.
- z and var.z: Fisher's z score and it's variance obtained from the Pearson correlation r to z transformation.
- pval.single: p-value against the null hypothesis of no treatment effect, derived by a t-test for outcome.flag "cont" or Wald test for outcome.flag "bin".
- events1c and events2c: Correction of events1 and events2 zero event counts or event counts = patient number. When no events occurred, 0.5 was added, and when all patients experienced the event, 0.5 was subtracted. When one of events had zero counts while the other had maximum counts, no adjustment occurred.
- meta.id: Meta-analysis ID variable to uniquely identify any potential meta-analysis in the dataset. Consistent to what has been discussed before, all results that share a common comparison, outcome and subgroup (optional, subgroups not given in any case) may be combined in a meta-analysis.
- se.new: Depending on outcome.flag, se.new is equal to the square-root of var.lrr (outcome.flag = DICH), square-root of var.smd (CONT) or se (IV).

#### 3.2.2 Eligibility criteria for Publication Bias Test and Adjustment

Initially, the analysis is restricted to results with outcome.flag DICH, CONT and IV. binary, continuous and survival outcomes. This corresponds to 734954 out of a total of 744720 results. Ioannidis and Trikalinos (2007a) outlined criteria for application of small study effect tests:

- Sample size: A meta-analysis is comprised of at least ten studies (n = 9916 remaining).
- Study size: The ratio between largest variance of an estimate and smallest variance of an estimate is larger than four (n = 9616 remaining).
- Significance: At least one treatment effect has a p-value below the significance threshold 0.05 (n = 7586 remaining)
- **Heterogeneity**: The  $I^2$  statistic of a given meta-analysis is smaller than 0.5, thus, the proportion of between study variance of the overall variance is smaller than 0.5 (n = 405 remaining).

Additionally, the following criteria have been applied:

• Sensitivity Analyses: When the same results are used multiple times for different metaanalyses, only one is retained. More precisely, if a study hat the same study.name and same effect, it was considered a duplicate, and the smaller meta-analysis of the two was excluded. The intention is to exclude sensitivity analyses which are operated on subsets of the available results. • **Zero events**: In the case of binary outcomes, meta-analyses with zero events in any study and any group are excluded (n = 20 out of meta-analyses with at least ten studies).

The results of this reduction of the dataset are shown in the flow-chart in Figure 3.2. By applying the criteria, we are only able to analyse a subset of the original data. Only the data that had accessible all results data available was used for adjustment. Thus, all meta-analyses with outcome.flag == IV are ommitted in a second step of the analysis (Analysis dataset (1) and (2) in Figure 3.2).

#### **Exclusions due to Computational Errors**

There has been cases where meta-analyses could not be analysed with Copas selection model because the likelihood function could not be optimized based on the settings. It was ommitted in this case.

- Meta-analysis with meta.id = 93868, id = "CD004072", subgroup.id = "CMP-006.08.01". One small meta-analysis (se.new = 1.02) with a risk ratio 0.072, while the remaining risk ratios were all larger than 0.45. (and the fixed effects estimate close to 1). It failed when analysed based on log risk ratios.
- Meta-analysis with meta.id = 144568, id = "CD006109", outcome.id = "CMP-001.06" and outcome.measure "MD". z-score based analysis failed, although funnel plot looks more or less symmetric. Small studies with very similar var.z with z-scores 0.3 and -0.1.
- Meta-analysis with meta.id = 35673, id = "CD001790", outcome.id = "CMP-001.03" and outcome.measure = "PETO.OR". Analysis failed when using Hedges g transformed effects. Large studies have  $g \sim 0$  while one smaller study (se = 0.3) has g = 1 (and another with a little larger se has  $g \sim 0$ ).

Because the number of errors is low, the meta-analyses have only been ommitted for the specific outcome measures, but are kept in the others.

#### The Analysis Dataset

The dataset that was ultimately tested and adjusted for publication bias comprises 1 metaanalyses and 6592 results. The mean number of participants in the treatment group is 206.9 vs 256.7 in the unrestricted dataset and the mean total number of participants 393.2 vs 623.2. The mean publication year is 1999.8575604 vs 2000.9711014 in the unrestricted dataset.

From the meta-analyses with incomplete data (outcome.type == IV), there are 36 with outcome.measure.merged = "Risk Ratio", 26 "Std. Mean Difference", 25 "Hazard Ratio" and 15 "Odds Ratio" (and 25 additional, different outcome measures).

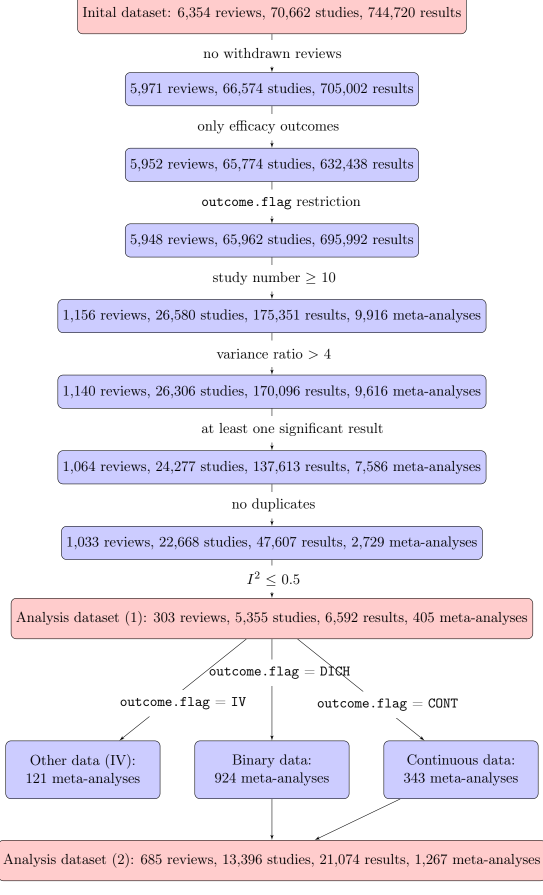
#### 3.2.3 Analysis Procedure

All computations were performed in the R computing environment (R Core Team, 2018). The R packages meta (Schwarzer, 2007) and metafor (Viechtbauer, 2010) were used for meta-analysis, small study effect tests and adjustments. The excess significance test was adapted from van Aert et al. (2019). For data manipulation procedures, the tidyverse packages were used (Wickham, 2017), and for plotting the ggplot2 package (Wickham, 2016).

The methods described in the methods chapter 2 most often apply directly to the algorithms used in meta and metafor. For binary outcomes, log risk ratios were used as treatment effect estimates in the meta-analysis, and for continuous outcomes, Hedge's g. For the other outcomes, the effect and se as provided in the dataset was passed to meta::metagen.

The effect sizes were transformed for adjustment in the way described in section 2.6. The *p*-values for adjusted treatment effect estimates were calculated by the Wald method.

In the case of the one-sided test procedure (small study effect and excess significance tests), the effect side in which bias was expected had to be prespecified. This was solved by comparing the number of significant findings on each side (original effect scale, i.e. for binary outcomes log risk ratios, etc.); the side with more significant findings (two-sided p-value < 0.05) was considered the side of potential bias. If numbers were equal, the side of the fixed effects treatment was used. Details to Copas selection model algorithm and its application can be found in Rücker et al. (2011). In short, two values for a and b in section 2.5.2 were used; a limited range with a between -1.7 and 2, and b between 0.16 and 0.32 was applied first (analog to a most extreme selection process of P(select|small trial with sd = 0.4) = 0.1 and P(select|large trial w. sd = 0.05) = 0.9). If the most extreme selection process is unable to pass the significance test of no small study effect (p-value > 0.1), then a wider range was applied (a between -5.4 and 2 and b between 0 and 0.32). If there is still no non-significant small study effect, the result was NA.



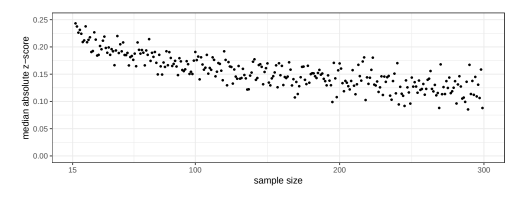
**Figure 3.2:** Flow-chart of the exclusion of meta-analyses for the final analysis. The exclusion criteria are given at the right of the arrows.

# Chapter 4

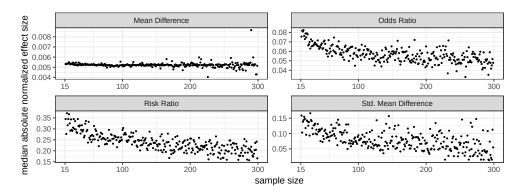
## Results

Using Fisher's z transformation, we can transform all effect sizes of the dataset on the same scale such that they are comparable to each other. Thus, as a first step, an exploratory plot of the median effect size and it's dependence on the study sample size can be shown.

The absolute value of the median z-score for a given sample size of a trial is shown in Figure 4.1. It is clearly visible that the absolute value decreases with increasing sample size, i.e. that the effect size is becoming smaller. The trend flattens off after  $\sim$  sample size = 400 (not shown). The pattern is similar in Figure 4.2, where the original effect size measures "Odds Ratio", "Risk Ratio", "Mean Difference" and "Std. Mean Difference" are used (the most common measurs in the dataset). Here, the effect sizes are scaled (mean subtraction and divison through standard error) with respect to all other effects of the same measure.



**Figure 4.1:** Median of the absolute z-score plotted against the total sample size.



**Figure 4.2:** Median of the absolute value of the normalized, original effect size plotted against the total sample size.

#### 4.1 Publication Bias Test Results

The meta-analyses fulfilling the criteria from chapter 3, section 3.2, are analysed with one-sided small publication bias tests and excess significance tests. The direction in which bias is expected is the one on which more significant results are (p-value < 0.05, two-sided). The tests are applied on the original effect size measures, since the journal editors and the researchers also base their decisions on them. Different tests are applied depending on the outcome being binary or continuous.

Multiple tests are applied on the same data to compare their results. A histogram of *p*-values for each test and all meta-analyses will summarize the overall evidence against the null-hypothesis of no publication bias, as displayed in Figure 4.3. Skewedness indicates evidence for publication bias

The abbreviations in Figure 4.3 are shortly explained with references to chapter 2:

"Excess significance" denotes the excess of significant p-values testing method from Ioannidis and Trikalinos (2007b), see 2.4.4. For continuous and IV outcomes, the names refer to:

- Egger's test, weighted linear regression test described in section 2.4.1
- Thompson and Sharp's test, weighted linear regression test adjusted for between-study heterogeneity, section 2.4.1
- Begg and Mazumdar's test, rank test described in section 2.4.1

For binary outcomes, the names refer to:

- Harbord's test, likelihood score based test (section 2.4.3)
- Peter's test, weighted linear regression with inverse sample size as explanatory variable described in section 2.4.3
- Rücker's test, test based on the arcsine transformation of proportions, in combination with Thompson and Sharp's regression test (section 2.4.3)
- Schwarzer's test, rank based test using the expected event counts computed with the hypergeometric distribution (section 2.4.3)

The histograms in Figure 4.3 show that the tests are not alway in agreement if publication bias is present in the data. The p-values of excess significance and Schwarzer's test and the p-values for IV outcomes are rather uniformly distributed, or even skewed to the right. Notably, excess significance test, Schwarzer's test and rank tests have been shown to lack power. The tests that are more suitable (regression based tests in general) all have proportions of significant p-values (p > 0.1) clearly above 10 % which would be the expected false positive rate.

In Figure 4.3, the meta-analyses with an estimated  $I^2$  of zero are depicted, because some methods are known to only be suitable when no heterogeneity is present (excess significance test and also Egger's test). Others are specially constructed to adjust for between study heterogeneity (Thompson and Sharp's test and Rücker's test). In continuous outcomes, when Thompson and Sharp's test is applied, the evidence for publication bias decreases, as can be seen in the smaller proportion of blue on the left corner in Figure 4.3. However, this is also due to application to meta-analyses with  $I^2$  equal zero, in which case the method is known to lack power. Similarly, the evidence decreases somewhat when Rücker's test is extended by Thompson and Sharp's method to account for heterogeneity. The moderate decrease indicates however that the previous restriction to meta-analyses with  $I^2 < 0.5$  is sufficient to remove meta-analyses with large heterogeneity and that the test results are not heavily influenced by unaccounted between study heterogeneity.

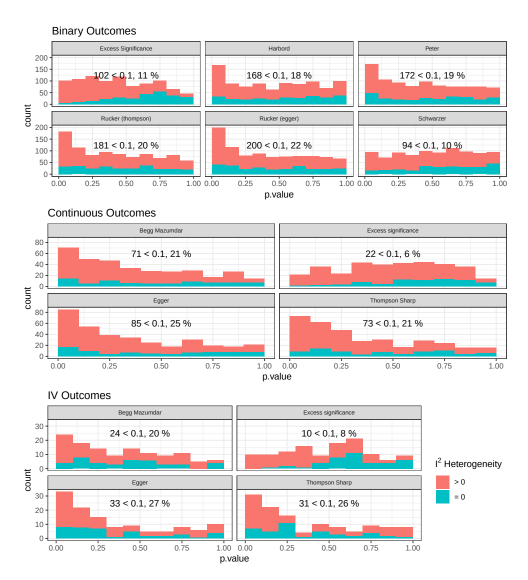
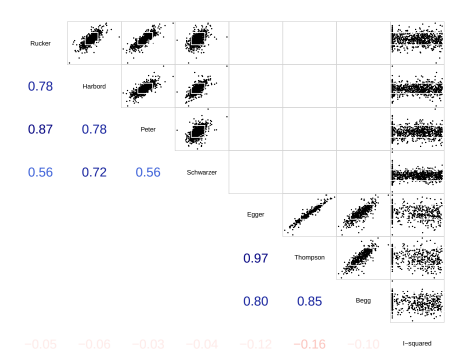


Figure 4.3: Histogram of one-sided p-values for small study effect in direction of larger effect sizes. The testing method is indicated in the header, bin width is equal to 0.1. The proportion of meta-analyses with signifiant publication bias based on the threshold of 0.1 is displayed inside the figures.

The p-values of tests can be summarized by computing their harmonic mean. In the case of binary tests, the p-values of Rücker's, Peters, Harbord's, Schwarzer's and excess significance tests are used, in the case of continuous and outcome.flag = IV outcomes, Egger's, Thompson and Sharp's, Begg and Mazumdar's and excess significance tests are used. This lead's to an overall of 20.3% significant results ( $p_{\text{harmonic}} < 0.1$ , 19.7% outcome.flag = DICH, 20.1% outcome.flag = CONT, 25.6% outcome.flag = IV).

#### 4.1.1 Publication Bias Test Consistency

It cannot be seen in Figure 4.3 if the tests used are finding small study effects for the same metaanalyses. A simple method to check the consistency of test results is to compare scatterplots and empirical Spearman correlations between the test statistics. This is done in Figure 4.4. Here, there is no separation between IV and continuous outcomes. The upper left rectangle is displaying binary outcome results and the lower right continuous and IV outcomes results. Also, the  $I^2$  statistic is included. Since no approximately normally distributed test statistic is used for



**Figure 4.4:** Pairs-plot for test statistics of small study effect and excess significance. The lower panel gives the Spearman correlations for the different test statistics, and the upper panel displays a scatterplot. The colors indicate magnitude and direction of the correlation coefficients. The rectangle with white borders displays the area within which both tests have absolute value < 1.64 (dots inside are statistically not significant by  $0.1\ p$ -value threshold).

excess significance tests, it is not shown here

The observed patterns on the scatterplots differ, and some small study effect test statistics do align better than others. Regression based tests as Egger and Thompsons test which are methodically almost identical are closely aligned, which is reflected in large correlation coefficients. Continuous and IV outcome type tests align more closely than binary outcome tests. While correlation coefficients between binary outcome tests vary between each other, Harbord's test statistic has similar correlation coefficients with the other small study effect test statistics. Because scatterplots and correlation coefficients can be misleading, also a Tukey mean-difference or Bland-Altman of transformed p-values plot is shown for four scenarios in Figure 4.5:

- For Egger's and Thompson's tests, which is supposedly the most similar test and should show the least deviations and systematic errors
- For Egger's and excess significance tests
- For Harbord's and Rücker's tests
- For Harbord's and excess significance tests

This can be justified since all tests are supposed to measure the evidence for publication bias. For the plots, the *p*-values of the tests are transformed on the entire continuous scale by a logit transformation  $f(x) = \log(\frac{p}{1-p})$ . The mean *p*-value ((f(p-value no. 1) + f(p-value no. 2))/2) is

then displayed against the difference between the f(p-value). If no systematic errors and biases exist between the measurement methods, then

- the mean of the differences should be around zero (no systematic error)
- the points should scatter independently on the y-axis and no general increase or decrease with the mean of the transformed p-values should be visible (and the linear regression fit is flat)

There are likely systematic errors and bias between the tests, although the extent seems to vary. Most error seems to be between small study effect tests and excess significance tests, because the slope of the linear regression fit is likely positive. This means that the excess significance test finds less evidence in cases when both p-values are small and more evidence when both p-values are large, on average.

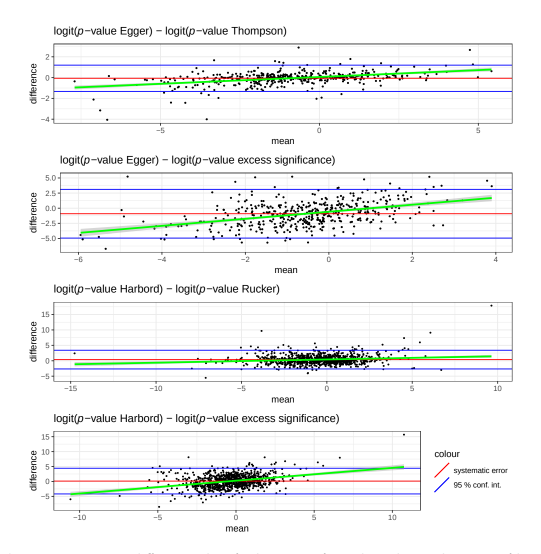
The confidence intervals from Figure 4.5 can be used as limits of agreement, which gives, after back-transformation, around 0.9 for Egger's and Thompson's test, and around 0.99 for Harbord's and Rücker's tests. This suggests that additionally to the bias correspondence between the tests is not very good in general.

The previous results suggest that the results will also differ substantially after applying the common dichotomization of p-values. Some proportion of the meta-analyses will only be significant using a single test, while being non-significant otherwise. This can be seen in Table 4.1. It displays the percentage of meta-analyses with a certain number of significant test results. For both outcome types, there is only marginal agreement, very few cases are significant in any test. Around 67% of the dataset is not significant, no matter which test is used. When only using small study effect tests, 29.5% of binary outcome tests and 31.9% of IV and continuous outcome tests had at least one significant result. After applying the Bonferroni correction for multiple testing, this shrinks to 12.3% for binary and 14.4% for continuous and IV outcomes. To compare significant findings for small study effect tests and excess significance tests, Harbord's or Egger's test results are compared with excess significance tests. 24.1% of binary outcome analyses had at least one of the two test p-values being significant, and equivalently, 28.4% for continuous and IV outcomes. The numbers change to 13.9% for binary outcomes and 18.1% for continuous and IV outcomes after applying the Bonferroni correction. 5.1% have significant Harbord's test result and significant excess significance test result (2.3% with Bonferroni). Of the continuous outcomes, we have 3.9% with Egger's test and 1.3% with Bonferroni correction.

Count	Binary Outcomes	Continuous and IV
0	66.1 %	65.3 %
1	12.3 %	11.9 %
2	7.7 %	6.9 %
3	6.8 %	14.2 %
4	5.7 %	1.7 %
5	1.3 %	=

**Table 4.1:** Counts Number of significant test results per meta-analysis, separated for outcome types. Last entry for continuous and IV outcomes is empty since one test less was applied

The precise proportions of agreement are provided in table 4.2. We see that only in few cases when excess significance test indicates statistical significance, small study effect tests do so. Correspondence between excess significance tests and small study effects thus is considered rather random. Linear regression based tests agree more often with other linear regression based tests, and agreement between small study effect tests is in general well above 60 % (at best, 95 % test agreement in significance for IV outcome.flag).



**Figure 4.5:** Mean - difference plots for logit transformed p-values. The mean of logit transformed p-values is displayed on the x-axis and the difference on the y-axis. Blue and red lines display the systematic error and the confidence intervals of the systematic error (limits of agreement). In green, a linear regression fit is shown with 95% CI bands.

## 4.2 Small Study Effects Adjustment

#### 4.2.1 Change in Effect Size after Adjustment

There are methods that can take into account the presence of publication bias in meta-analyses when estimating the overall treatment effect. The methods work in a semi-automatic manner; they will not only adjust for publication bias if smaller studies show larger effects, but also in the opposite case. The latter results in the adjusted overall treatment effect being *larger* than the unadjusted, overall treatment effect.

To compare the effects of adjustment between meta-analyses of different outcomes, the outcome measures are transformed to standardized mean differences and Fisher's z-scores (see section 2.6 for details). When comparing to unadjusted effects, fixed or random effects meta-analysis estimates are used as references

Figure 4.6 displays the difference  $\delta$  between the estimated meta-analysis treatment effect and the regression adjusted treatment effect 2.5.1,  $\hat{\theta}_M - \hat{\theta}_{Adj.}$ . The absolute value  $|\hat{\theta}_M|$  is taken and  $\hat{\theta}_{Adj.}$  is negative if the signs change in the original  $\theta$ . Thus, a positive difference indicates a reduction of the original effect size, and the magnitude of the difference indicates the extent of

	Agreement (overall)	Agreement (significance)
Excess significance, Schwarzer	0.85	0.27
Excess significance, Peter	0.77	0.32
Excess significance, Rucker	0.79	0.43
Excess significance, Harbord	0.81	0.46
Peter, Schwarzer	0.84	0.63
Schwarzer, Rucker	0.85	0.72
Schwarzer, Harbord	0.87	0.78
Rucker, Peter	0.88	0.67
Harbord, Peter	0.87	0.63
Excess significance, Egger	0.77	0.64
Excess significance, Thompson	0.80	0.59
Excess significance, Begg	0.78	0.36
Thompson, Egger	0.93	0.92
Thompson, Begg	0.87	0.70
Egger, Begg	0.84	0.70
Excess significance, Egger (IV)	0.71	0.12
Excess significance, Thompson (IV)	0.71	0.10
Excess significance, Begg (IV)	0.74	0.04
Thompson, Egger (IV)	0.95	0.94
Thompson, Begg (IV)	0.86	0.79
Egger, Begg (IV)	0.86	0.83

**Table 4.2:** Overall proportion of agreement if significant or unsignificant, and for significance only. Horizontal lines separate binary, continuous and IV outcomes (order as in table). The reference significance test is the one with more significant results.

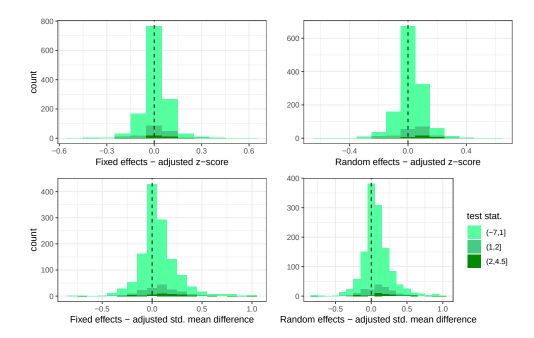
#### the adjustment.

Additionally, the test statistic of heterogeneity adjusted publication bias tests (Rücker's and Thompson's test) are displayed with green color. Test statistics smaller < 1 (light green colored) are equivalent to no evidence for publication test, test statistics between one and two to weak evidence, and above two they indicate evidence for publication bias (dark green). An adjusted effect with evidence for publication bias can be regarded as a more realistic estimate of treatment efficacy. Some very large and very small differences have been omitted in the z-score and std. mean difference histograms; they are shown in Table 4.4.

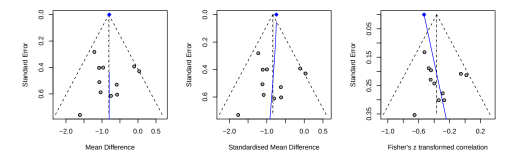
Most often, adjustment leads to a reduction of overall treatment effect estimates, because the bins on the positive side of the histograms are larger. Adjustment is stronger when random effects meta-analysis is used as reference, because it gives larger weights to small studies.

Contrary to naive expectation, we see cases with large adjustment, but no evidence for small study effects. This is because the linear regression parameter estimates are large, but estimated with high uncertainty, such that there will be few evidence for small study effects (publication bias), but nonetheless, the adjustment will be large. The authors that developped the methods also recommend to use the methods in cases where there is clear evidence for publication bias. The color legend in Figure 4.6 thus gives a sense of confidence that there is in the credibility of the adjustment.

Additionally, some meta-analyses with positively adjusted, larger overall treatment effects after adjustment (left side of the histogram) have at the same time evidence for publication bias. This is no wrong result, although the p-value of the test is one-sided,  $i.\,e.$  only allowing for publication bias for large effects (small studies with large effects). But the effects and their variances have been transformed, and it is possible that the shape of the funnel plot changes upon transformation; Figure 4.7 shows this for illustratory purposes. From the left to the right, the funnel plot is shown for mean differences (the original measure), standardized mean differences and Fisher's z scores. This is quite the most extreme case, with three different effects of adjustment, depending on outcome measure; there is no change with mean differences, reduction of effect sizes with std. mean differences and increase with Fisher's z transformed correlation. Note that while the rank of the effect sizes is usually preserved after transformation, the relative size and especially the variance may vary. One effect of the Fisher's z-transformation is that the effect sizes are



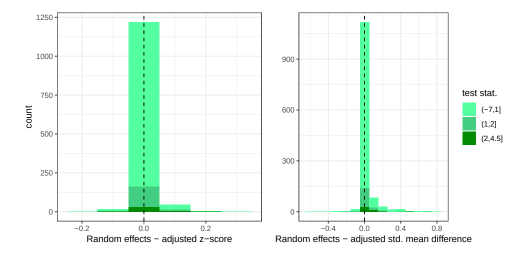
**Figure 4.6:** Histogram of the treatment effect differences between meta-analysis and regression adjusted meta-analysis. Negative differences indicate greater adjusted effect sizes than meta-analysis effect sizes. The bins are centered at zero and binwidth is equal to 0.1. Deeper green color indicates more evidence for small study effects.



**Figure 4.7:** Funnel plots for a meta-analysis based on three different effect size measures: Mean differences, std. mean differences and Fisher's z transformed correlations and corresponding standard errors. Vertical dashed lines indicate meta-analysis estimates, the rhombus with the curved blue line the adjusted treatment effect.

bounded on [-1,1], and thus, very large effect sizes will influence the fit of the linear regression less than for example in std. mean differences which are not bounded. In contrast, the variance of the correlation is directly tied to the sample size, which makes it a suitable proxy for study size (variance of the mean difference is in contrast strongly influenced by the standard deviations). Thus, cases where the direction of adjustment changes upon transformation are not very rare.

Figure 4.8 shows the same for effect sizes adjusted by Copas; Copas selection model substitutes its estimates with random effect estimates when it finds no evidence for small study effects. Therefore, the effect of adjustment by Copas can be seen when comparing adjusted with random effects meta analysis estimates. Again, we clearly see that more effect sizes are adjusted downwards. Additionally, there is more coincidence between the publication bias test statistics and adjustment, which is as expected.



**Figure 4.8:** Histogram of the treatment effect differences between meta-analysis and Copas adjusted meta-analysis. Negative differences indicate greater adjusted effect sizes than meta-analysis effect sizes. The bins are centered at zero and binwidth is equal to 0.1. Deeper green color indicates more evidence for small study effects.

Table 4.3 shows quantiles and means for the various differences and the overall proportion of downward adjusted effect sizes. When std. mean difference is used as an effect measure, there are (substantially) more reduced effect sizes. The means in Table 4.3 suggest that the average reduction is small. To recall some other findings out of Table 4.3: 5% or 69 meta-analyses have their z-score reduced by more than 0.13 by regression adjustment (and 5% or 69 increased by -0.11 or more, fixed effects reference). Also, std. mean difference is reduced by 0.39 compared to fixed effects estimates in 5% or 69 meta-analyses (or increased by 0.24).

	5%	25%	50%	75%	95%	mean	= 0 (%)	>= 0 (%)	> 0 (%)	No adj. est. (%)
z: Fixed - Copas	-0.04	-0.01	0.00	0.01	0.04	0.00	8.14	50.14	42.00	65.49
z: Random - Copas	-0.00	0.00	0.00	0.00	0.04	0.00	67.15	85.23	18.08	65.49
z: Fixed - Regression	-0.11	-0.03	0.01	0.05	0.13	0.01	0.00	52.02	52.02	0.00
z: Random - Regression	-0.13	-0.02	0.02	0.06	0.16	0.02	0.00	56.56	56.56	0.00
g: Fixed - Copas	-0.05	-0.01	0.00	0.01	0.11	0.01	19.38	56.41	37.03	56.92
g: Random - Copas	-0.00	0.00	0.00	0.00	0.16	0.02	60.09	85.01	24.93	56.92
g: Fixed - Regression	-0.21	-0.03	0.04	0.14	0.40	0.06	0.00	60.16	60.16	0.00
g: Random - Regression	-0.20	-0.02	0.05	0.17	0.44	0.08	0.00	62.68	62.68	0.00
IV: Fixed - Copas	-0.06	-0.01	0.00	0.01	0.07	0.00	23.97	42.15	66.12	59.50
IV: Random - Copas	-0.00	0.00	0.00	0.01	0.08	0.01	61.98	28.93	90.91	59.50
IV: Fixed - Regression	-0.24	-0.02	0.03	0.11	0.36	0.03	0.00	66.12	66.12	0.00
IV: Random - Regression	-0.25	-0.02	0.04	0.13	0.49	0.05	0.00	68.59	68.59	0.00

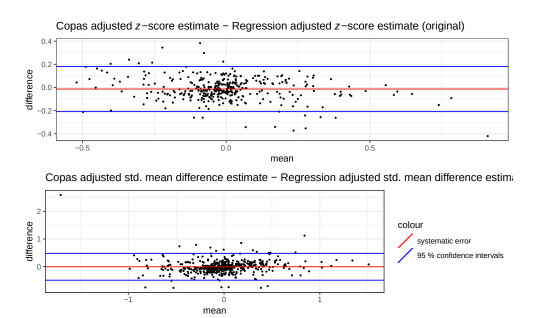
**Table 4.3:** Quantiles and Means of the differences between meta-analysis pooled treatment effects and small study adjusted treatment effects. The column with the names > 0" give the percentages of estimates larger than zero or larger or equal zero. The column "No adj. est." gives the percentage of missing estimates due to non-significant publication bias test (for Copas) and computational errors. The row names indicate which outcome measure, meta-analysis method and adjustment method is used. Abbreviations are used for z-score (= z) and std. mean difference (= d).

#### 4.2.2 Comparison of adjustment methods

Regression adjusted estimates are compared to the estimates of Copas selection model if these are not equal to random effects meta-analysis in Figure 4.9. A Tukey mean difference plot can serve to reveal systematic differences and biases between the two measurement methods. Note that most 121 IV outcome data is not included when using z-score and std. mean difference.

meta.id	id	comparison.nr	subgroup.nr	z fixed	z random	z Copas	z regression	g fixed	g random	g Copas	g regression
9957	CD000370	8	2	0.66	0.66	0.66	0.82	1.59	1.59	1.40	0.28
22324	CD001183	7	0	-0.48	-0.48	-0.48	-0.21	-1.10	-1.12	-0.50	-0.11
49559	CD002307	2	1	-0.10	-0.08	-0.10	0.03	-0.47	-0.42	-0.41	-3.00
197675	CD008625	2	2	-0.60	-0.60	-0.60	-0.39	-1.72	-2.02	-1.01	-0.69
222767	CD010060	1	0	0.31	0.32	0.31	0.23	0.50	0.52	0.20	-0.49

**Table 4.4:** Missing meta-analysis pooled treatment effect and adjusted treatment effects. Abbreviations are used for z-score (= z) and std. mean difference (= d).



**Figure 4.9:** Mean - difference plots for publication bias adjustment methods. The mean of the adjusted treatment effects is displayed on the x-axis and the difference on the y-axis. Blue and red lines display the systematic error and the confidence intervals of the systematic error (limits of agreement). Two values have been omitted in the middle plot for std. mean difference and one for z-score (see Table 4.4).

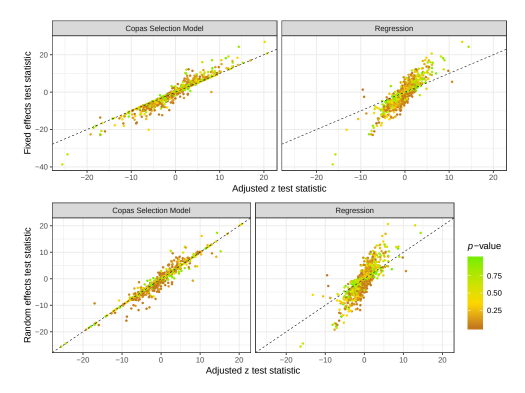
No formal tests are provided, but the at least there seems to be no clear bias or systematic error. The limits of agreement in Figure 4.9 are large. We conclude thus that the impact of regression adjustment on the effect sizes is in general not substantially larger than the impact of Copas selection model in the subset of data where the estimate of the Copas selection model is not equal to a random effects estimate. There is however a small difference, indicating that regression estimates have a little bit a larger absolute value. There might be some bias between adjusted z-scores, where regression estimates seem to be somewhat smaller when the mean is a little above zero, and somewhat larger when the mean is a little below zero.

#### 4.2.3 Change in Evidence for Treatment Effects

Adjustment for small study effects in meta-analysis will provide new effect sizes and standard errors. The evidence against the null hypothesis of no treatment effect can be computed newly. The test statistics  $\frac{\theta}{\sec(\theta)}$  of random effects and fixed effects meta-analysis and the corresponding adjusted test statistics are shown in Figure 4.10. The original scale of the effects sizes is used, since it is the measure on which policy-makers assess the treatment efficacy.

It can be seen that the alignment on the diagonal changes depending on the adjustment method. The effect of adjustment by copas can be seen when comparing it to random effects meta-analysis. There, less of an effect of adjustment on the evidence can be seen.

Thus, the decrease in evidence is larger when using adjustment with regression, which can be

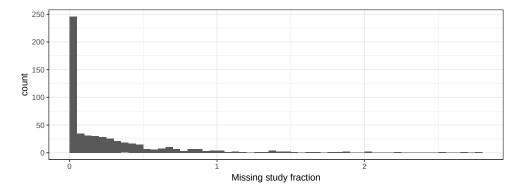


**Figure 4.10:** Test statistics of meta-analyses and adjustment methods. The color indicates the evidence for publication bias (p value of R "ucker's or Egger's test), and the dotted line depicts the diagonal through the origin.

expected as the uncertainity of the additional parameter from the linear regression fit to estimate publication bias is included in the uncertainity of the estimate. In contrast, as the Copas selection model algorithm does a sensitivity analysis, the uncertainity of additional model parameters does not affect the estimate as they are not estimated but treated as fixed (see section 3.2.3 and section 2.5.2).

Large adjustment in effect sizes and test statistics is not necessarily accompained by evidence for publication bias (color of the dots) in regression adjustment, as already discussed. The dark dots can be found more often far away from the diagonal. Some cases which have more evidence for treatment efficacy after adjustment have evidence for publication bias, which is not expected, since the publication bias tests applied are one-sided and only test for bias towards large effects. In some cases, the automatic detection of the side on which publication bias was expected (section 3.2.3) might failed, because few significant effects are given (used to define side of bias) and the weighted mean effect is close to zero, in which case it is difficult to detect the side of publication bias beforehand. There are some adjustments with large increase in evidence and clear dark orange color, as in the plots on the right hand side in Figure 4.10. Two of them are investigated in detail.

- z-value of regression adjustment 10.5 vs fixed effects z-value 5.7: When the funnnel plot of these meta-analyses are investigated, it becomes clear that there is no reason to assume publication bias since the smaller studies are near to risk ratios equal to one, and larger studies show larger ratios. Publication bias tests disagree in their findings (Rücker's and Harbord's tests: 0.058 and 0.05, and Peters and Schwarzer's test: 0.467 and 0.79).
- z-value of regression adjustment 9.24 vs fixed effects z-value 2.51: Asymmetry of the funnel plot could be confirmed when analysing it with mean differences. The change to a larger treatment effect is rather due to large between study heterogeneity and the specific routine of the regression adjustment. Because it takes into account between-study heterogeneity



**Figure 4.11:** Histogram of the fraction of missing studies from the total number of studies in a meta-analyses (only data shown where Copas estimate was obtained, thus n = 0)

(bias parameter times  $\tau^2$ ), while fixed effects meta-analysis does not, it returns a larger treatment effect. The adjusted effect (MD = - 10.3) lies between the fixed effect estimate (-2.56) and the random effects estimate (-18.7).

In the case of Copas selection model and when both test statistics are large (> 10), the explanation lies in a specialty of the model. The adjusted treatment effect estimates are not larger than the unadjusted, but their standard errors are different and sometimes smaller, because they are obtained from the Fisher information matrix of the log-likelihood (see section 2.5.2).

The Copas selection model also allows to compute the number of missing studies, given that the model's assumptions are correct. It finds that 2,618 are missing, which corresponds to 11.4% from all 22,916 analysed studies. Figure 4.11 shows a histogram of the overall fraction of missing studies. Note that random effects meta-analysis substitutes have been excluded (828 out of 1,388.

We can see that in some occasions, the method finds more than half of all studies are missing. In most occasions, the estimate of missing studies is zero, as can be seen in Table 4.5. The discrepancy between mean and median may indicate that the estimate of 11.4% missing studies depends somewhat on these extreme cases. As can be written of from Table 4.5, 5%, *i. e.* 28 meta-analyses have 17.6 or more studies missing: in fact, these 5% most extreme make up for 1,060, more than 30% of missing studies.

	= 0	5%	25%	50%	75%	95%	mean
Missing fraction	226	0	0	0.1	0.4	1.0	0.3
Missing study number	226	0	0	1.5	5.9	20.1	4.7

**Table 4.5:** Fraction of missing studies and estimates of missing studies with their zero counts ("= 0"), quantiles and means.

#### 4.3 Publication Bias over Time

To test if publication bias in meta-analyses has decreased over time and newer meta-analyses have less bias, a generalized linear model has been set up.

The dependent variable is the standardized mean differences (therefore, from outcome.flag = IV, only the std. mean differences are used). As before, the std. mean difference is mirrored to one side by multiplying with the sign of the expected side of bias, *i. e.*-1 or 1. The explanatory variable is the standard error, and the weights are the inverse of the variance of the std. mean difference. Additionally, random effects for the meta-analysis and for the review are added, the

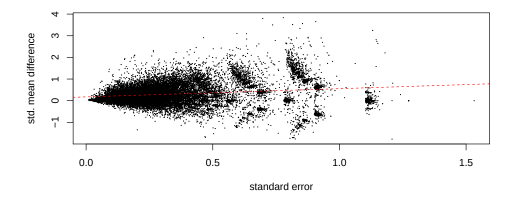


Figure 4.12: Std. mean differences of all meta-analyses plotted against their standard error, with the fit of a generalized linear model (dotted red line).

former being nested in the latter. Figure 4.12 shows the complete dataset with the fit of the previously described model (dotted red line). The details and evidence for the small study effects in the generalized linear model are given in table 4.6, the coefficients in table 4.7

	Model	df	AIC	BIC	logLik	Test	L.Ratio	p-value
null.fit	1	4.00	18768.23	18800.12	-9380.11			
small.study.fit	2	5.00	18230.52	18270.39	-9110.26	1  vs  2	539.71	0.00

**Table 4.6:** Anova table for a generalized linear model for small study effects compared to the null model. 'small.study.fit' denotes the model with random intercepts for meta-analyses and reviews.

	estimate	2.5%CL	97.5%CL
(Intercept)	0.18	0.16	0.19
se.smd.pool	0.38	0.35	0.41

**Table 4.7:** Coefficients and 95% confidence limits of the generalized linear model. 'se.smd.pool' denotes the standard error of the std. mean difference.

It is possible to include meta-analysis specific random slopes. Before continuing, it is tested if incorporation of random slopes improves model fit. Table 4.8 shows the anova table for a model with and without random slopes.

Model diagnostics indicate that there is no benefit in including random slopes for the single meta-analysis.

To test if publication bias varies over time, we include the year of the publication of the review as an additional explanatory variable. Ulitmately, it is of interest if there is an interaction between the small study effect (publication bias) and time, i.e. if the slope varies depending on the year. Because such a model is nested in a more simpler model where the study year is only an additive effect, all these models are fitted. Table 4.10 displays model fit diagnostics.

Because neither AIC, BIC of the F-test do show any improvement of model fit, it does not seem as if publication bias has substantially decreased or increased over the years. The review publication year is however not directly tied to the publication year of the studies that are affected by publication bias. Thus, using another measure of time, the results may be different. Thus, the models are re-fitted with the study publication years.

The analysis of model fit indicates that there is an improvement when using the study year as a covariate, but only as an additive effect. AIC and BIC are smaller and the p-value of the

	Model	df	AIC	BIC	logLik	Test	L.Ratio	p-value
small.study.fit	1	5.00	18230.52	18270.39	-9110.26			
small.study.rs.fit	2	7.00	18233.07	18288.88	-9109.53	1  vs  2	1.46	0.48

**Table 4.8:** Anova table for two generalized linear model fits. 'small.study.fit' denotes the model with random intercepts for meta-analyses and reviews, 'small.study.rs.fit' the model with additional random slopes per review.

	Model	df	AIC	BIC	logLik	Test	L.Ratio	p-value
small.study.fit	1	5.00	18230.52	18270.39	-9110.26			
year.fit	2	6.00	18231.61	18279.46	-9109.81	1  vs  2	0.91	0.34
year.int.fit	3	7.00	18233.59	18289.40	-9109.79	2  vs  3	0.03	0.87

**Table 4.9:** Anova table for three generalized linear model fits. 'small.study.fit' denotes the model with the standard error as explanatory variable, 'year.fit' the model with standard error and the centered (- 2013) review year as explanatory variable, and 'year.int.fit' the model with interaction between the two.

F-test is very small. However, not the size of the effect, but the slope of the linear regression fit would have to be decreased to ultimatively conclude a decrease of publication bias over the years.

	Model	df	AIC	BIC	logLik	Test	L.Ratio	p-value
small.study.fit	1	5.00	18115.23	18155.05	-9052.62			
year.fit	2	6.00	18095.04	18142.83	-9041.52	1  vs  2	22.19	0.00
year.int.fit	3	7.00	18096.90	18152.65	-9041.45	2  vs  3	0.14	0.71

**Table 4.10:** Anova table for three generalized linear model fits. 'Small study fit' denots the model with the standard error as explanatory variable, 'year.fit' the model with standard error and the centered (- 2000) study year as explanatory variable, and 'year.int.fit' the model with interaction between the two.

## Chapter 5

# Discussion

### 5.1 Results in the light of the Literature

Before discussing the results of the previous analyses, some results of similar studies are discussed briefly and compared to the results of this study, if possible. When clear methodological drawbacks are found in the analysis, the reader is referred to chapter 2 for comments on the disadvantages. The amount of studies analysing publication bias makes it not possible to discuss any of them, and thus we only some interesting and/or representative examples are shown.

#### 5.1.1 Egger et al. (1997)

The Cochrane database (1996 issue 2) and publications from the four leading medicine journals (the Lancet, BMJ, JAMA and Annals of Internal Medicine) between 1993 and 1996 were used to find systematic reviews with randomized controlled trials for the application of Egger's small study effect test. They included 38 meta-analyses with at least 5 studies and binary outcomes from the Cochrane Library and 37 from the journals. Five (13%) meta-analyses from Cochrane and 13 (38%) from the journals had significant two sided small study effect test results (p-value < 0.1). Also, they found that test-statistics were more often negative, which correspondend in their setup to larger effects in small studies. (63.2 among Cochrane meta-analyses and 70.3 for journal meta-analyses). The results from this report with Rücker's test are: 14.8%, and when using regression adjustment to decide about the direction of publication bias, we get 80.7%.

#### 5.1.2 (Leicester)

Out of 397 systematic reviews from the Cochrane database (complete number for 1998, issue 3), 49 had more than ten included studies and binary outcomes and 48 compared two treatments. They were analysed by trim-and-fill method (Duval and Tweedie, 2000) to detect and adjust for funnel plot asymmetry (similar to small study effect tests, but the method is known to overestimate bias). 23 were found to have missing studies, and eight had more than three missing studies which was considered to be significant publication bias. Additionally, they found that three estimates of random effects meta-analysis became non-significant after adjustment, and one became significant (by negative adjustment). The results are difficult to compare, but the methodological limitations make this findings unreliable.

#### 5.1.3 Ioannidis and Trikalinos (2007a)

Data processing steps were however different, and only binary outcomes as used in two-by-two tables were analysed. The approximately corresponding numbers are put in parentheses for comparison. The Cochrane Library from 2003 (issue 2) was used. After removal of duplicates and intractable meta-analyses, they had 6,873 meta-analyses with more than two studies left

(20,219). When only using one meta-analysis per review, this reduced to 846 (3,555. Then, the criteria that have also been applied in this masters thesis are applied:  $I^2 < 0.5$ , variance ratio of smallest and largest effects > 4, at least one significant study result and at least 10 studies to be used. Afterwards, they applied Harbord's, Egger's and Begg's Test to the dataset. The reader can compare some corresponding numbers in Table 5.1.

	Ioannidis	Thesis
Wider dataset (n)	6873	20219
study number $> 10$	13%	12%
variance ratio $> 4$	72%	75%
study sig. number $> 1$	55%	52%
all exclusion criteria	5%	5%
harbord test p-value $< 0.1$	12%	16%

**Table 5.1:** Comparison of results from Ioannidis et.al. (2007) to the results of the thesis. The percentage of meta-analysis which match all exclusion criteria denotes the ones that apply to all criteria in the table plus the small heterogeneity criterium. Harbord test is two-sided.

#### 5.1.4 Souza et al. (2007)

Reviews of the World Health Organization (WHO) Reproductive Health Library (RHL), issue 9, were analysed with the trim-and-fill method. The RHL reproduces and expands reviews from the Cochrane Library with implications for developping countries. 21 of 105 reviews contained more than ten studies and were used. Trim-and-fill found asymmetry in 18 of 21 studies, and 10 had more than 3 missing studies ("significance"). Two of those and one with one missing studies found no evidence for treatment effects after the 0.05 p-value threshold after adjustment by trim-and-fill.

#### 5.1.5 Kicinski *et al.* (2015)

The author uses a bayesian hierarchical selection model, but does not analyse treatment effects, but the parameters of the weight function of the selection model, which is estimated with a bayesian approach and MCMC sampling.

The author provides an estimate of the probability of including significant findings versus non-significant findings in Cochrane meta-analyses over time. The data is from the Cochrane Library from 2013 (issue number not provided). The author excluded treatment - treatment comparisons and analysed safety and efficacy meta-analyses separately (how this was achieved is not documented in the paper).

From 3845 reviews, the author separated 907 reviews with more than ten studies. From those, 539 compared placebo to treatment. After removing duplicates and sensitivity analyses, 358 analyses with 1297 meta-analyses remained. From these 191 were excluded because they comprised overall mortality and withdrawal, because these could not clearly be specified as safety or efficacy, respectively.

1106 meta-analyses from 329 meta-analyses, containing 802 efficacy and 304 safety meta-analyses. The median publication year per meta-analysis was 1997 for efficacy meta-analyses and 1999 for safety meta-analyses. Then, a bayesian two-step hierarchical selection model was applied (Kicinski (2013), Silliman (1997)). It assumes that the selection process is a two-step weight, which assigns different probabilities to non-significant and significant effects. Thus, a ratio of publication probability between significant and non-significant study estimates using a two-step weight function. The model was fitted with the Monte-Carlo Markov-Chain algorithm STAN and the

geometric mean was used as an estimate for the publication probability ratio. Simulations performed in (Kicinski, 2013) indicate that the method performes well if the true mean effect size was not small, and also robust to small study effects. It outperformed Egger's test and Begg's test in assessing publication bias, especially when small study effects were absent, and had lower false-positive rates.

The results showed a clear publication bias for significant results for efficacy meta-analyses (27% higher for significant studies, 95%CI credible intervals: 1.18 to 1.36). The probability was more than twice as high in 27% of the meta-analyses (95% CI: 23% to 31%). But the probability decreased: from 1.65 (95% CI: 1.31 to 2.15) in 1980 (average publication year) to 1.36 (95% CI: 1.17 to 1.62) in 1990 to 1.18 (95% CI: 1.04 to 1.33) in 2000. For sake of completeness, the probability of inclusion was 1.78 (95% CI: 1.51 to 2.13) larger for non-significant safety effect estimates, but again, decreased with time (1.77, 95% CI: 1.46 to 2.21 in 2000). The results are in line with the results from this study, but it was not possible to reafirm the finding of weaker publication bias in reviews published more frequently.

#### 5.1.6 van Aert et al. (2019)

The Authors of this recent study sample 366 meta-analyses randomly from the Cochrane Library (supposedly 2018 or 2019, not mentioned). They exclude any meta-analysis which include effect sizes identical to effect sizes in other meta-analyses (the larger meta-analysis is retained). Additional criteria were:  $I^2 < 0.5$  and at least 5 studies. The meta-analyses were analysed using standardized mean differences and four different tests for publication bias: Egger's test, p-uniform test (Van Assen *et al.*, 2015), Begg and Mazumdar's rank correlation test and the excess significance test.

The authors found, based on the significance threshold of p < 0.1, the following results (own results in brackets): 12.2 % significant results from Egger's test (16.3), 8.5 % significant results from Begg and Mazumdar's rank correlation test (10.5) and 4.4 % significant excess significance test results (9.7). It is known that the publication bias tests are lacking power in general as sample size is usually small. Decreasing sample size will result in lower power, especially if the maximal sample size is  $n \ge 5$ . Additionally, it was also not taken into consideration that it may be difficult to state that publication bias is present in a meta-analysis when no result within it is statistically significant (only 18.8 % of the effects of the sampled meta-analyses were). This, together that it has not been tried to exclude safety outcomes, may account for the large differences between the results.

The author's come to the conclusion in their study that in contrast to other studies, they find few evidence for publication

#### 5.1.7 Further Studies on Publication Bias

Zhang et al. (2013) find publication bias in critical care studies with similar methods. Nüesch et al. (2010) report publication bias in clinical osteoarthritis research because of funnel plot asymmetry and pledge for routine assessment of publication bias. Dechartres et al. (2013) analyse publication bias based on sample size in meta-analyses from top journals and Cochrane reviews in 93 Meta-analyses and find that, for example, effects in trials with less than 50 patients were 48% larger than in larger trials. Onishi and Furukawa (2014) find that in 36 Meta-Analyses without comprehensive literature research, there ar 19.4% significant publication bias tests (Egger's test).

### 5.2 Interpretation

In this study, a proportion of the meta-analyses that have been scrutinized have publication bias. Since regression and rank test have been applied one-sided, they have very likely larger effects

in smaller studies that in larger studies.

There have been proposed a number of reasons for this tendencies in smaller studies (see Egger et al. (1997) for a thorough discussion). In clinical settings, it might be that the requirement for more study participants in large studies might systematically change the study population. However, this systematic differences could influence the results in two ways: If recruitement of participants with a disease is difficult, and thus, also participants with less severe types of the disease are included, it might be that the effect of treatment is larger in the smaller study. But one could also argue that the opposite could eventually be the case.

The main cause for funnel plot asymmetry with smaller studies having larger effects is publication bias. Publication bias enforces questionable research practices from researchers such as selective outcome reporting, p-hacking and data fabrication. Larger studies are less affected, and have also in general higher methodological quality (Egger et al., 1997), which leads to smaller effect sizes.

Presence of publication bias in meta-analyses can ultimately lead to patient harm, because decision makers in clinical practice use the meta-analyses to take decisions if they use a treatment. On the swiss Cochrane website, it is stated that Cochrane is the "single most reliable source of evidence on the effects of health care". However, publication bias does in general lead to too large confidence in the results of the meta-analyses. In some cases, it might be that the evidence for a treatment in a meta-analysis is an artifact of publication bias. Besides harm to patient, this can cause a waste of ressources.

The comparison of the results with previous analyses of publication bias in the Cochrane Database of Systematic Reviews implies gives more confidence in the results of this study, because they are largely in line.

#### 5.3 Limitations

After application of the criteria of Ioannidis and Trikalinos (2007a), there are left 11.6 % of the reviews and 32.9% of the studies. It is not clear how the results would be in the remaining dataset if tests were applicable. The most restrictive criterium of exclusion was that a meta-analysis has to have at least ten studies. Thus, in the data that is scrutinized, there has been enforced and ongoing research. Ioannidis (2005) state that publication bias is also a consequence of modern discovery oriented research, and thus it may well be that publication bias is also an issue when the number of studies are lower.

It also has to be said that the removal of safety outcomes is only partially successfull. This is an issue, because it is not well known if and how publication bias affect safety outcomes. Kicinski et al. (2015) suggests that they are inversely affected, with editors publishing in favor of non-significant adverse effects.

Also, it has been previously discussed that the method to detect the side on which bias is to be expected can fail and provide meaningless results, in which case a two-sided test is more meaningful. However, one could also discuss in more general terms if one can speak of publication bias in meta-analyses that have only one significant effect out of ten (n = 417).

There is also criticism that one can address on the methods. It is possible that the results are caused by systematic differences in the population of study participants only. This is considered rather unlikely. Other cirticism is more severe. The small study effect tests fail to account for statistical significance, however, this is a major component of publication bias. Small study effect test work because they take into account that smaller studies need larger effects to be significant. The only test that is applied that takes into account statistical significance per se is the excess significance test. When looking at it's results, it seems as if it is rather a test for between study heterogeneity per se, because it does agree very poorly with small study effect tests, and finds almost no excess significance in tests with  $I^2 = 0$ . The choice of the effect size and proxy of

5.4. OUTLOOK 47

the study size is crucial when assessing publication bias. It is not clear which measure of effect size should be used when applying the test; the original measure ( $e.\ g.$  a mean difference) may have standard errors that do not only depend on study size, but journal editors use this measure as the basis for their decision of publication. Fisher's z transformed correlations have standard errors that are a function of study participant number only, but in contrast the effect size is bounded at [-1,1], making it less suitable for linear regression. Standardized mean differences are possibly a good compromise.

The ciriticism also applies for adjustment methods, as they also do not use statistical significance but the size of the effects. A weakness of the regression adjustment is that it also adjusts if the uncertainity in the small study effect is large. To compare it in this case with the meta-analysis estimate is difficult because the uncertainity in estimating the small study effect affects the uncertainity of the treatment effect. It is not clear how to proceed when one wants to adjust for publication bias. Copas selection model uses the p-value threshold of 0.1 to decide if adjustment is necessary or not. One can argue that this threshold is arbitrary. Additionally, the threshold is used in two manners: first to decide if adjustment is necessary at all, and secondly, if the adjusted model does suitably account for publication bias (p > 0.1). Hypothesis tests against the null hypothesis can however not be used to decide in favor of the null-hypothesis, and a separate test would be needed there.

With Copas selection model, it is in general the issue that it the model is not estimating, but rather applying some criteria of parsimony (the model with least publication bias with p-value for small study effect > 0.1 is chosen). Some authors find in their simulation studies that regression methods overperform the Copas selection model (Rücker  $et\ al.$ , 2011).

Thus, the results are not exact and could be improved if more suitable methods are applied. The bayesian approach taken by Kicinski *et al.* (2015) has many advantages over the applied methods, *e. g.* by accounting for statistical significance, but relies on assumptions over the weight function and is at least somewhat sensible on the choice of the prior distributions.

#### 5.4 Outlook

Given the size and abundancy of information in the dataset, the possibilities to investigate sources of bias in the Cochrane Database of Systematic Reviews are not exhausted (see for example in Fanelli *et al.* (2017)). In particular, one could easily come up with more hypotheses that could be tested in meta-regression, as introduced in the last section of the results.

There are numerous suggestions for different measures of publication bias and small study effects, which could be applied on the dataset (see Mueller et al. (2016)). This study mainly relies on suggestions from Sterne et al. (2001b), Ioannidis and Trikalinos (2007a) and Rücker et al. (2011). It uses most often methods that are well integrated in packages (Viechtbauer (2010), Schwarzer (2007)). Although there are well addressed, mathematical justifications for these methods, there might be better methods not yet tested on large datasets. An evaluation of all suggested methods was far beyond the scope of this study.

## 5.5 Implications

This study is so far the largest assessment of publication bias by small study effects and uses data that has been collected after major efforts have been made to curb publication bias. The results indicate, similarly to other studies, that the efforts did not yet resolve the issue.

The findings further imply that when using meta-analyses, it is key to investigate sources of between-study heterogeneity; an alternative is to rely solely on the largest trials (Stanley *et al.*, 2010). Since the Cochrane Organization uses results from grey literature to limit the effects of

publication bias, but still does not manage to abolish it, there is also the need for journal editors to change their publication policies.

# Appendix A

# Appendix

```
sessionInfo()
## R version 3.5.1 (2018-07-02)
## Platform: x86_64-apple-darwin15.6.0 (64-bit)
## Running under: macOS 10.14.4
##
## Matrix products: default
## BLAS: /Library/Frameworks/R.framework/Versions/3.5/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/3.5/Resources/lib/libRlapack.dylib
##
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
## attached base packages:
## [1] stats
                 graphics grDevices utils
                                               datasets methods
                                                                    hase
##
## other attached packages:
   [1] nlme_3.1-137
                        corrgram_1.13
                                        bindrcpp_0.2.2 rvest_0.3.2
   [5] xml2_1.2.0
                                        metafor_2.1-0
                                                        Matrix_1.2-15
##
                        xtable_1.8-3
  [9] gridExtra_2.3
                        metasens_0.3-2 meta_4.9-4
                                                        forcats_0.3.0
## [13] stringr_1.3.1
                        dplyr_0.7.8
                                        purrr_0.2.5
                                                        readr_1.2.1
                                        ggplot2_3.1.0
## [17] tidyr_0.8.2
                        tibble_1.4.2
                                                        tidyverse_1.2.1
## [21] testit_0.9
                        knitr_1.20
##
## loaded via a namespace (and not attached):
## [1] Rcpp_1.0.0
                           lubridate_1.7.4
                                              lattice_0.20-38
## [4] gtools_3.8.1
                           assertthat_0.2.0
                                              digest_0.6.18
  [7] foreach_1.4.4
                           R6_2.3.0
                                              cellranger_1.1.0
## [10] plyr_1.8.4
                           backports_1.1.2
                                              evaluate_0.12
## [13] httr_1.3.1
                           highr_0.7
                                              pillar_1.3.0
## [16] gplots_3.0.1.1
                           rlang_0.3.0.1
                                              lazyeval_0.2.1
## [19] readxl_1.1.0
                                              rstudioapi_0.8
                           gdata_2.18.0
## [22] labeling_0.3
                           munsell_0.5.0
                                              broom_0.5.0
## [25] compiler_3.5.1
                           modelr_0.1.2
                                              pkgconfig_2.0.2
## [28] tidyselect_0.2.5
                           seriation_1.2-7
                                              codetools_0.2-15
## [31] dendextend_1.12.0
                           viridisLite_0.3.0
                                              crayon_1.3.4
## [34] withr_2.1.2
                           bitops_1.0-6
                                              MASS_7.3-51.1
## [37] grid_3.5.1
                           jsonlite_1.5
                                              gtable_0.2.0
```

```
## [40] registry_0.5-1 magrittr_1.5 scales_1.0.0

## [43] KernSmooth_2.23-15 cli_1.0.1 stringi_1.2.4

## [46] viridis_0.5.1 iterators_1.0.10 tools_3.5.1

## [49] glue_1.3.0 gclus_1.3.2 hms_0.4.2

## [52] colorspace_1.3-2 cluster_2.0.7-1 caTools_1.17.1.1

## [55] TSP_1.1-7 bindr_0.1.1 haven_2.0.0
```

### A.1 Data Pre-processing

```
#Dataset processing function to get transformed effect sizes, p-values, event
# counts with increments, etc. :
pb.process3 <- function(data){</pre>
  data <- data %>% mutate(meta.id =
            group_indices(., id, comparison.id, outcome.id, subgroup.id)) %>%
    group_by(meta.id) %>% mutate(study.id2 = row_number()) %>% ungroup()
  #Mark duplicate effects between meta-analyses:
  data <- data %>% group_by(meta.id) %>%
    mutate(n = n()) %>% ungroup() %>% group_by(id) %>%
    mutate(dupl.id =
         dupl.finder(effects = effect, names = study.name, metas = meta.id),
           dupl.remove =
  dupl.max.finder(duplicate.index = dupl.id, study.number = n, metas = meta.id)) %%
    ungroup()
  data <- data %>% mutate(lrr = NA,
    var.lrr = NA,
    hedgesg = NA,
    var.hedgesg = NA,
    cohensd = NA,
    var.cohensd = NA,
    smd.ordl = NA,
    var.smd.ordl = NA,
    pval.single = NA,
    cor.pearson = NA,
    var.cor.pearson = NA,
    z = NA,
    var.z = NA,
    events1c = NA,
    events2c = NA,
    sig.single = NA,
    smd.pool = NA,
    se.smd.pool = NA)
  cont.ind <- which(data$outcome.flag == "CONT")</pre>
  bin.ind <- which(data$outcome.flag == "DICH")</pre>
  IV.ind <- which(data$outcome.flag == "IV")</pre>
```

```
#Outcome.flag == "DICH"
data[bin.ind,] <- escalc(data = data[bin.ind,], ai = events1, n2i = total2,</pre>
                         ci = events2, n1i = total1,
                         to = "only0",
                         add = 1/2,
                         measure = "RR", append = T,
                    var.names = c("lrr", "var.lrr")) #Calculate Risk Ratios
data[bin.ind,] <- escalc(data = data[bin.ind,], ai = events1, n2i = total2,
                         ci = events2, n1i = total1,
                         to = "only0",
                         add = 1/2,
                         measure = "OR2DL", append = T,
var.names = c("smd.ordl", "var.smd.ordl")) #Calculate SMD (w. logistic transf.)
#What to do if there are O/total cells:
data[bin.ind,] <- data[bin.ind,] %>%
  mutate(events1c = case_when(events1 == 0 ~ events1 + 0.5,
                               events2 == 0 ~ events1 + 0.5,
                               events1 - total1 == 0 ~ events1 - 0.5,
                               events2 - total2 == 0^{\circ} events1 - 0.5,
                               TRUE ~ events1),
         events2c = case_when(events2 == 0 ~ events2 + 0.5,
                              events1 == 0 ~ events2 + 0.5,
                              events2 - total2 == 0 ~ events2 - 0.5,
                              events1 - total1 == 0 ~ events2 - 0.5,
                              TRUE ~ events2))
#What to do if there is one zero and one total:
data[bin.ind,] <- data[bin.ind,] %>%
 mutate(events1c = case_when(events2 == 0 & events1 - total1 == 0 ~ events1,
                               events1 == 0 & events2 - total2 == 0 ~ events1,
                               TRUE ~ events1c),
         events2c = case_when(events2 == 0 & events1 - total1 == 0 ~ events2,
                              events1 == 0 & events2 - total2 == 0 ~ events2,
                              TRUE ~ events2c))
data[bin.ind, "pval.single"] <- data[bin.ind, ] %>%
 mutate(pval.single = 2*(1-pnorm(abs(lrr/sqrt(var.lrr))))) %>% select(pval.single)
#Outcome.flag == "CONT"
data[cont.ind,] <- escalc(data = data[cont.ind,], m1i = mean1, m2i = mean2,</pre>
                          sd1i = sd1, sd2i = sd2, n1i = total1, n2i = total2,
                          measure = "SMD" , append = T,
              var.names = c("hedgesg", "var.hedgesg")) #Calculate Hedge's g
data[cont.ind, ] <- data[cont.ind, ] %>%
  mutate(cohensd = (mean1 - mean2)/sqrt((((total1 - 1)*sd1^2) +
                  (total2 - 1)*sd2^2/(total1 + total2 - 2)),
         var.cohensd = ((total1 + total2)/(total1 * total2)) +
```

```
(cohensd^2)/(2*(total1 + total2)))
data[cont.ind, ] <- data[cont.ind, ] %>%
    mutate(cohensd = case_when(sd1 == 0 ~ NA_real_,
                                                               sd2 == 0 ~ NA_real_,
                                                               total2 == 0 ~ NA_real_,
                                                               total1 == 0 ~ NA_real_,
                                                                TRUE ~ cohensd))
#p-value for continuous outcomes (Student):
data[cont.ind, "pval.single"] <- data[cont.ind, ] %>%
    mutate(t = (mean1 - mean2)/(sqrt((((total1-1)*sd1^2) + (total2-1)*sd2^2)/(sqrt(((total1-1)*sd1^2) + (total2-1)*sd2^2)/(sqrt((total1-1)*sd1^2) + (total2-1)*sd2^2)/(sqrt((total1-1)*sd1^2)/(sqrt((total1-1)*sd1^2))/(sqrt((total1-1)*sd1^2)/(sqrt((total1-1)*sd1^2)/(sqrt((total1-1)*sd1^2)/(sqrt((total1-1)*sd1^2)/(sqrt((total1-1)*sd1^2)/(sqrt((total1-1)*sd1^2)/(sqrt((total1-1)*sd1^2)/(sqrt((total1-1)*sd1^2)/(sqrt((total1-1)*sd1^2)/(sqrt((total1-1)*sd1
                                                     (total1 + total2 -2))*sqrt((1/total1)+(1/total2))),
                    pval.single = 2*(1-pt(abs(t), df = total1 + total2 - 2))) %>%
    select(pval.single)
sd1.0 <- which(data$outcome.flag == "CONT" & data$sd1 == 0)
sd2.0 <- which(data$outcome.flag == "CONT" & data$sd2 == 0)
sd.0 \leftarrow union(sd1.0, sd2.0)
data[sd.0, "pval.single"] <- NA #Such that p-value here is not equal to zero
#Outcome.flag == "IV"
data[IV.ind, "pval.single"] <- data[IV.ind, ] %>%
    mutate(pval.single = 2*(1-pnorm(abs((effect)/se)))) %>%
    select(pval.single)
#Declare smd.ordl, cohensd and SMD of IV flag all as "smd.pool":
data <- data %>%
    mutate(smd.pool = case_when(outcome.flag == "DICH" ~ smd.ordl,
                                                                  outcome.flag == "CONT" ~ cohensd,
                                                                  outcome.flag == "IV" &
                                                                      outcome.measure.merged == "SMD" ~ effect,
                                                                  TRUE ~ NA_real_),
                    se.smd.pool = case_when(outcome.flag == "DICH" ~ sqrt(var.smd.ordl),
                                                                         outcome.flag == "CONT" ~ sqrt(var.cohensd),
                                                                         outcome.flag == "IV" &
                                                                             outcome.measure.merged == "SMD" ~ se,
                                                                        TRUE ~ NA_real_))
#Transform effect sizes:
smd.pool.ind <- which(!is.na(data$smd.pool))</pre>
#Pearson correlation:
data[smd.pool.ind, "cor.pearson"] <- data[smd.pool.ind, ] %>% mutate(
    a = ((total1 + total2)^2)/(total1*total2),
    cor.pearson = smd.pool/sqrt((smd.pool^2) + a),
    var.cor.pearson = ((a^2)*se.smd.pool^2)/(((smd.pool^2)+a)^3)) %>%
   select(cor.pearson)
```

```
data[smd.pool.ind, "var.cor.pearson"] <- data[smd.pool.ind, ] %>%
   mutate(
   a = ((total1 + total2)^2)/(total1*total2),
   cor.pearson = smd.pool/sqrt((smd.pool^2) + a),
   var.cor.pearson = ((a^2)*se.smd.pool^2)/(((smd.pool^2)+a)^3)) %>%
   select(var.cor.pearson)
  #Fisher's z-score:
 data[smd.pool.ind, "z"] <- data[smd.pool.ind, ] %>%
   mutate(z = atan(x = cor.pearson),
          var.z = 1/(total1 + total2 - 3)) \% % select(z)
 data[smd.pool.ind, "var.z"] <- data[smd.pool.ind, ] %>%
   mutate(z = atan(x = cor.pearson),
          var.z= 1/(total1 + total2 - 3)) %>% select(var.z)
 data$sig.single <- ifelse(data$pval.single < 0.05, 1, 0)
 return(data)
#Function to find sensitivity analyses. The function gives all meta-analyses
#that are subdivided
#into sensitivity analyses a common number.
#*to use with "data %>% mutate(.. = dupl.finder(..))"*
dupl.finder <- function(effects, names, metas){</pre>
 results <- rep(NA, times = length(effects))
 meta.double.marker <- 1</pre>
 for(u in seq_along(effects)){
   if(is.na(results[u])){
     double.indices <- which(effects[u] == effects & names[u] == names)</pre>
     if(length(double.indices) > 1){
    #If any meta-analysis has already duplicates, give all the same double marker
       if(any(!is.na(results[double.indices]))){
         already.marked <- which(!is.na(results[double.indices]))</pre>
         meta.double.marker.2 <- unique(results[double.indices[already.marked]])</pre>
         #Collect meta.ids with same double markers
         meta.ids.2 <- unique(metas[which(results %in% meta.double.marker.2)])</pre>
         meta.ids <- metas[double.indices] #Collect metas with the same results
```

```
meta.ids.pooled <- union(meta.ids, meta.ids.2)</pre>
           meta.indices <- which(metas %in% meta.ids.pooled)</pre>
           results[meta.indices] <- meta.double.marker
        } else{
           meta.ids <- metas[double.indices]</pre>
           meta.indices <- which(metas %in% meta.ids)</pre>
           results[meta.indices] <- meta.double.marker</pre>
      } else results[u] <- 0</pre>
      meta.double.marker <- meta.double.marker + 1</pre>
    }}
  return(results)
}
#Function picks the largest meta-analysis within a set of sensitivity analyses,
#and assigns it a "0" *to use with "data %>% mutate(.. = dupl.finder(..))":*
dupl.max.finder <- function(duplicate.index, study.number, metas){</pre>
  results <- rep(NA, length(duplicate.index))</pre>
  for(u in seq_along((duplicate.index))){
    duplicate.indices <- which(duplicate.index %in% duplicate.index[u])</pre>
    if(duplicate.index[u] != 0){
      meta.ids <- metas[duplicate.indices]</pre>
      max.meta.id <- meta.ids[which.max(study.number[duplicate.indices])]</pre>
      max.meta.indices <- which(metas %in% max.meta.id)</pre>
      if(length(unique(max.meta.id)) < 2){</pre>
        results[max.meta.indices] <- 0</pre>
      } else print("error")
    } else results[duplicate.indices] <- 0</pre>
  }
  results[is.na(results)] <- 1
  return(results)
}
```

### A.2 Analysis

Data is subdivided into meta-analyses, and after all meta-analyses are excluded that are not suited for analysis, meta-analyses are done and saved. "data.ext2" is the dataset processed with

the previously introduced "pb.process3" function. Functions defined in the global environment are marked with a "\*USER.DEF\*". They can be find in the next section.

```
#Load data:
\# rm(list = ls())
# PATH_HOME = path.expand("~") # user home
# PATH = file.path(PATH_HOME, 'Data/PubBias')
# PATH2 = file.path(PATH_HOME, 'PubBias')
# FILE = 'cochrane_2019-07-04.csv'
# PATH_DATA = file.path(PATH, 'data')
# PATH_CODE = file.path(PATH2, 'code')
# PATH_RESULTS = file.path(PATH2, 'results_new')
# PATH_FIGURES = file.path(PATH_RESULTS, 'figures')
source(file.path(PATH_CODE, 'PubBias_functions.R')) #Load user defined functions
load(file.path(PATH_DATA, "PubBias_2019-07-19.RData")) #Load data as contributed
#by Simon Schwab.
# load(file.path(PATH_RESULTS, "data_used_for_analysis.RData"))
data.ext2 <- pb.process3(data) #*USER.DEF*</pre>
# PRE-ANALYSIS PART - EXCLUDE UNSUITABLE META-ANALYSES:
#To skip pre-analysis:
# load(file.path(PATH_RESULTS, "meta_id_vector.RData"))
# # ----- Uncomment to run pre-analysis -----
\# # Get exclusion critetia (I-squared, n. sig. findings, s.e. ratio, dupl.index):
meta.info.bin <- data.ext2 %>% filter(outcome.flag == "DICH") %>%
  group_by(meta.id) %>%
 mutate(n.pre = n()) %>% filter(n.pre >= 10) %>%
  filter(!all(events1 == 0) & !all(events2 == 0)) %>% #No events
 mutate(se.lrr = sqrt(var.lrr)) %>%
  summarize(dupl.remove = unique(dupl.remove),
            id = unique(id),
            outcome.desired = unique(outcome.desired),
            n.sig.single = sum(sig.single, na.rm = T),
            se.min = min(se.lrr, na.rm = T),
            se.max = max(se.lrr, na.rm = T))
meta.info.cont <- data.ext2 %% filter(outcome.flag == "CONT") %>%
  group_by(meta.id) %>%
 mutate(n.pre = n()) %>% filter(n.pre >= 10) %>%
  filter(!all(mean1 == 0) & !all(mean2 == 0)) %>% #No means
  filter(!all(sd1 == 0) | !all(sd2 == 0)) %>% #No sd's
  summarize(dupl.remove = unique(dupl.remove),
            id = unique(id),
            outcome.desired = unique(outcome.desired),
```

```
n.sig.single = sum(sig.single, na.rm = T),
            se.min = min(se, na.rm = T),
            se.max = max(se, na.rm = T))
meta.info.iv <- data.ext2 %>%
  group_by(meta.id) %>%
  mutate(n.pre = n()) \%\% filter(n.pre >= 10) \%\%
  filter(outcome.flag == "IV") %>%
  filter(outcome.measure.merged == "Rate Ratio" |
           outcome.measure.merged == "SMD" |
           outcome.measure.merged == "MD" |
           outcome.measure.merged == "Hazard Ratio" |
           outcome.measure.merged == "OR" |
           outcome.measure.merged == "RR") %>% #Remove outcomes w. unclear def.
  summarize(dupl.remove = unique(dupl.remove),
            id = unique(id),
            outcome.desired = unique(outcome.desired),
            n.sig.single = sum(sig.single, na.rm = T),
            se.min = min(se, na.rm = T),
            se.max = max(se, na.rm = T))
meta.info.pre <- rbind(meta.info.bin, meta.info.cont, meta.info.iv)</pre>
#To save to control how many are excluded
meta.info.pre <- meta.info.pre %>%
  filter(id %in% data.review$id[which(data.review$withdrawn == FALSE)])
#No withdrawn meta-analyses
meta.info <- meta.info.pre %>%
  filter(n.sig.single > 0) %>% #At least one significant result
  filter((se.max^2)/(se.min^2) > 4) %>% #variance ratio > 4
  filter(dupl.remove == 0) %>% #no duplicates
  filter(outcome.desired == "efficacy") #Only efficacy outcomes
#Meta-analysis to get I2 (random effects meta-analysis):
meta.id.vector.org.ranef <- meta.info$meta.id #Do meta-analysis for each of those
meta.org.ranef.list <- list()</pre>
counter <- 0
for(u in meta.id.vector.org.ranef){
  counter <- counter + 1
  temp <- data.ext2 %>% filter(meta.id == u)
 print(c(u, counter))
 meta.org.ranef.list[[counter]] <- meta.fct.ranef.mom(temp)</pre>
exclusion.estimates <- cbind(
  meta.id = meta.id.vector.org.ranef,
 n = unlist(lapply(meta.org.ranef.list,
```

```
FUN = function(meta.analysis){meta.analysis$k})),
  Q = unlist(lapply(meta.org.ranef.list,
                  FUN = function(meta.analysis){meta.analysis$QE})))
meta.info.pre2 <- merge(meta.info, exclusion.estimates, by = "meta.id")</pre>
meta.info.I2 <- meta.info.pre2 %>%
 rowwise() \%% mutate(I2 = max(0, (Q - n + 1)/Q)) #calculate I2
meta.info <- meta.info.I2 %>% filter(n > 9) %>%
 filter(I2 < 0.5) #exclude I2 >= 0.5
meta.id.vector <- meta.info$meta.id #Vector of meta-ids that match the criteria.
save(meta.id.vector, file = file.path(PATH_RESULTS, "meta_id_vector.RData"))
save(meta.info.I2, file = file.path(PATH_RESULTS, "meta_id_I2.RData"))
#Get more extensive information from the original dataset about the meta-analyses:
meta.info.extended <- data.ext2 %>%
 filter(meta.id %in% meta.id.vector) %>%
 group_by(meta.id) %>%
 summarize(id = unique(id),
           outcome.flag = unique(outcome.flag),
           comparison.nr = unique(comparison.nr),
           comparison.name = unique(comparison.name),
           outcome.name = unique(outcome.name),
           outcome.nr = unique(outcome.nr),
           subgroup.name = unique(subgroup.name),
           subgroup.nr = unique(subgroup.nr),
           outcome.measure.merged = unique(outcome.measure.merged),
           outcome.measure = unique(outcome.measure),
           outcome.desired = unique(outcome.desired),
           mean.samplesize = mean(total1 + total2, na.rm = T),
           total.samplesize = sum(total1 + total2),
           var.samplesize = var(total1 + total2, na.rm = T),
           min.samplesize = min(min(total1, na.rm = T), min(total2, na.rm = T)),
           total.events = sum(events1 + events2),
           mean.events = mean(events1 + events2),
           mean.publication.year = mean(study.year, na.rm = TRUE),
           first.publication.year = min(study.year, na.rm = T),
           #*USER.DEF* function to detect the expected side of bias
           side = bias.side.fct2(outcome = outcome.flag, outcome.measure.merged,
                               lrr = lrr, var.lrr = var.lrr, smd = cohensd,
                               var.smd = var.cohensd, effect = effect, se = se),
           n.sig.single = sum(sig.single, na.rm = T),
           NA.sig.single = sum(is.na(sig.single)),
           se.min = min(se, na.rm = T),
           se.max = max(se, na.rm = T))
```

```
#Do all analyses with Fixed effects meta-analysis method and paule mandel tau^2
#estimator
settings.meta(method.tau = "PM", method = "Inverse")
#Load previously constructed lists (to skip analysis):
# load(file.path(PATH_RESULTS, "meta_complete_list_bin.RData"))
# load(file.path(PATH_RESULTS, "meta_complete_list_cont.RData"))
# load(file.path(PATH_RESULTS, "meta_complete_list_iv.RData"))
# load(file.path(PATH_RESULTS, "meta_complete_list_zscore.RData"))
\#\ load(file.path(PATH\_RESULTS,\ "meta\_complete\_list\_cohensd.RData"))
# BINARY DATA META-ANALYSES: -----
# # ----- Uncomment to run analysis -----
meta.id.vector.bin <-</pre>
 meta.info.extended$meta.id[which(meta.info.extended$outcome.flag == "DICH")]
meta.analyses <- list()</pre>
meta.analyses.asd <- list()</pre>
meta.analyses.reg <- list()</pre>
meta.analyses.copas <- list()</pre>
meta.tests.harbord <- list()</pre>
meta.tests.peter <- list()</pre>
meta.tests.rucker.mm <- list()</pre>
meta.tests.rucker.linreg <- list()</pre>
meta.tests.schwarzer <- list()</pre>
meta.tests.excess <- list()</pre>
counter <- 0
for(u in meta.id.vector.bin){
counter <- counter + 1</pre>
print(c(u, counter))
meta.analyses[[counter]] <-</pre>
 metabin(event.e = events1c, n.e = total1, event.c = events2c, n.c = total2,
        studlab = study.name, sm = "RR",
        method = "Inverse", data = data.ext2[data.ext2$meta.id == u,])
 #Arcsine transformed proportions to use in combination with R\widetilde{A}_{\overline{4}}cker's test:
meta.analyses.asd[[counter]] <-</pre>
 metabin(event.e = events1c, n.e = total1, event.c = events2c, n.c = total2,
        studlab = study.name, sm = "ASD",
        method = "Inverse", data = data.ext2[data.ext2$meta.id == u,])
```

```
#Adjustment for publication bias
meta.analyses.reg[[counter]] <- limitmeta(meta.analyses[[counter]])</pre>
meta.analyses.copas[[counter]] <- auto.copas(meta.analyses[[counter]],</pre>
                          sig.level = 0.1) #*USER.DEF* Copas selection model.
 #Publication bias tests.
meta.tests.excess[[counter]] <- tes.fct2(data = data.ext2[data.ext2$meta.id == u,])</pre>
meta.tests.harbord[[counter]] <-</pre>
  metabias(meta.analyses[[counter]], method.bias = "score", k.min = 2)
meta.tests.peter[[counter]] <-</pre>
  metabias(meta.analyses[[counter]], method.bias = "peters", k.min = 2)
meta.tests.schwarzer[[counter]] <-</pre>
  metabias(meta.analyses[[counter]], method.bias = "count", k.min = 2)
meta.tests.rucker.mm[[counter]] <-</pre>
  metabias(meta.analyses.asd[[counter]], method.bias = "mm", k.min = 2)
meta.tests.rucker.linreg[[counter]] <-</pre>
  metabias(meta.analyses.asd[[counter]], method.bias = "linreg", k.min = 2)
}
# counter.unknown.bin.copas.error <- 0
# for(u in meta.id.vector.bin){
# if(length(meta.analyses.copas[[counter]]) < 3){
# meta.analyses.copas[[counter]] <- c(NA, NA, NA)</pre>
# counter.unknown.bin.copas.error <- counter.unknown.bin.copas.error + 1
# }
# }
# print(c("unknown.bin.copas.errors = ", counter.unknown.bin.copas.error))
# bin.meta.list <- list(data.ext2, meta.id.vector.bin, meta.analyses,
# meta.analyses.asd,
# meta.analyses.reg,
# meta.analyses.copas,
# meta.tests.excess,
# meta.tests.harbord,
# meta.tests.peter,
# meta.tests.rucker.mm,
# meta.tests.schwarzer,
# meta.tests.excess,
# meta.tests.rucker.linreq)
\# save(bin.meta.list, file = file.path(PATH_RESULTS, "meta_complete_list_bin.RData"))
#
# #Load analysis data:
# meta.id.vector.bin <- bin.meta.list[[2]]</pre>
# meta.analyses <- bin.meta.list[[3]]</pre>
# meta.analyses.reg <- bin.meta.list[[5]]</pre>
# meta.analyses.copas <- bin.meta.list[[6]]</pre>
# meta.analyses.trimfill <- bin.meta.list[[7]]</pre>
# meta.tests.harbord <- bin.meta.list[[8]]</pre>
# meta.tests.peter <- bin.meta.list[[9]]</pre>
```

```
# meta.tests.rucker.mm <- bin.meta.list[[10]]</pre>
# meta.tests.schwarzer <- bin.meta.list[[11]]</pre>
# meta.tests.excess <- bin.meta.list[[12]]</pre>
# meta.tests.rucker.linreg <- bin.meta.list[[13]]</pre>
#Extract from lists:
bin.results <- data.frame(</pre>
  meta.id = meta.id.vector.bin,
  meta.es.measure = rep(times = length(meta.id.vector.bin), "log risk ratio"),
  n.sig.single2 = unlist(lapply(meta.analyses,
    FUN = function(meta.analysis){length(which(meta.analysis$pval < 0.05))})),
  #Meta-analysis:
  est.fixef = unlist(lapply(meta.analyses,
    FUN = function(meta.analysis){meta.analysis$TE.fixed})),
  est.ranef = unlist(lapply(meta.analyses,
                  FUN = function(meta.analysis) {meta.analysis$TE.random})),
  zval.fixef = unlist(lapply(meta.analyses,
                  FUN = function(meta.analysis){meta.analysis$zval.fixed})),
  zval.ranef = unlist(lapply(meta.analyses,
                  FUN = function(meta.analysis) {meta.analysis$zval.random})),
  pval.fixef = unlist(lapply(meta.analyses,
                   FUN = function(meta.analysis){meta.analysis$pval.fixed})),
  pval.ranef = unlist(lapply(meta.analyses,
                  FUN = function(meta.analysis) {meta.analysis$pval.random})),
  se.est.fixef = unlist(lapply(meta.analyses,
                  FUN = function(meta.analysis) {meta.analysis$seTE.fixed})),
  se.est.ranef = unlist(lapply(meta.analyses,
                  FUN = function(meta.analysis) {meta.analysis$seTE.random})),
  Q = unlist(lapply(meta.analyses, FUN = function(meta.analysis){meta.analysis$Q})),
  pval.Q = unlist(lapply(meta.analyses,
                         FUN = function(meta.analysis){meta.analysis$pval.Q})),
  #I2 = unlist(lapply(meta.analyses, FUN = function(meta.analysis){meta.analysis$I2.w})),
  tau = unlist(lapply(meta.analyses,
                      FUN = function(meta.analysis){meta.analysis$tau})),
  \#se.tau = unlist(lapply(meta.analyses, FUN = function(meta.analysis)\{meta.analysis\}se.tau\}
  sparse = unlist(lapply(meta.analyses,
                         FUN = function(meta.analysis){meta.analysis$sparse})),
  k = unlist(lapply(meta.analyses,
                    FUN = function(meta.analysis){meta.analysis$k})),
  #Adjustment:
  method.reg = unlist(lapply(meta.analyses.reg,
                  FUN = function(meta.adjust) {meta.adjust$method.adjust})),
  est.reg = unlist(lapply(meta.analyses.reg,
                  FUN = function(meta.adjust) {meta.adjust$TE.adjust})),
```

```
se.est.reg = unlist(lapply(meta.analyses.reg,
                FUN = function(meta.adjust) {meta.adjust$seTE.adjust})),
zval.reg = unlist(lapply(meta.analyses.reg,
                FUN = function(meta.adjust) {meta.adjust$zval.adjust})),
pval.reg = unlist(lapply(meta.analyses.reg,
                FUN = function(meta.adjust) {meta.adjust$pval.adjust})),
alpha.r = unlist(lapply(meta.analyses.reg,
                FUN = function(meta.adjust) {meta.adjust$pval.adjust})),
beta.r = unlist(lapply(meta.analyses.reg,
                FUN = function(meta.adjust) {meta.adjust$pval.adjust})),
Q.small = unlist(lapply(meta.analyses.reg,
                FUN = function(meta.adjust){meta.adjust$Q.small})),
Q.resid = unlist(lapply(meta.analyses.reg,
                FUN = function(meta.adjust){meta.adjust$Q.resid})),
G.squared = unlist(lapply(meta.analyses.reg,
                FUN = function(meta.adjust){meta.adjust$G.squared})),
est.copas = unlist(lapply(meta.analyses.copas,
                FUN = function(meta.adjust){meta.adjust[1]})),
se.est.copas = unlist(lapply(meta.analyses.copas,
               FUN = function(meta.adjust){meta.adjust[2]})),
missing.copas = unlist(lapply(meta.analyses.copas,
                FUN = function(meta.adjust){meta.adjust[3]})),
#Tests;
pval.d.tes.org = unlist(lapply(meta.tests.excess,
                               FUN = function(object){object[3]})),
pval.d.tes = unlist(lapply(meta.tests.excess,
                           FUN = function(object){object[4]})),
stat.d.tes = unlist(lapply(meta.tests.excess,
                           FUN = function(object){object[2]})),
excess.d = unlist(lapply(meta.tests.excess,
                         FUN = function(object){object[5]})) -
  unlist(lapply(meta.tests.excess,
                FUN = function(object){object[6]})),
pval.harbord = unlist(lapply(meta.tests.harbord,
                             FUN = function(meta.test){meta.test$p.value})),
stat.harbord = unlist(lapply(meta.tests.harbord,
                             FUN = function(meta.test){meta.test$statistic})),
pval.peter = unlist(lapply(meta.tests.peter,
                           FUN = function(meta.test){meta.test$p.value})),
stat.peter = unlist(lapply(meta.tests.peter,
                           FUN = function(meta.test){meta.test$statistic})),
pval.rucker = unlist(lapply(meta.tests.rucker.mm,
                            FUN = function(meta.test){meta.test$p.value})),
stat.rucker = unlist(lapply(meta.tests.rucker.mm,
                            FUN = function(meta.test){meta.test$statistic})),
pval.rucker.linreg = unlist(lapply(meta.tests.rucker.linreg,
                            FUN = function(meta.test){meta.test$p.value})),
stat.rucker.linreg = unlist(lapply(meta.tests.rucker.linreg,
                            FUN = function(meta.test){meta.test$statistic})),
```

```
pval.schwarzer = unlist(lapply(meta.tests.schwarzer,
                                FUN = function(meta.test){meta.test$p.value})),
  stat.schwarzer = unlist(lapply(meta.tests.schwarzer,
                                FUN = function(meta.test) {meta.test$statistic}))
)
# CONTINUOUS DATA META-ANALYSES: -----
# # ----- Uncomment to run analysis -----
meta.id.vector.cont <-</pre>
  meta.info.extended$meta.id[which(meta.info.extended$outcome.flag == "CONT")]
meta.analyses <- list()</pre>
meta.analyses.reg <- list()</pre>
meta.analyses.copas <- list()</pre>
meta.analyses.trimfill <- list()</pre>
meta.tests.excess <- list()</pre>
meta.tests.linreg <- list()</pre>
meta.tests.begg <- list()</pre>
meta.tests.mm <- list()</pre>
counter <- 0
for(u in meta.id.vector.cont){
counter <- counter + 1</pre>
print(c(u, counter))
outcome.measure.merged <-</pre>
  as.character(unique(data.ext2[data.ext2$meta.id == u,
                  "outcome.measure.merged"]) $outcome.measure.merged)
meta.analyses[[counter]] <-</pre>
  metacont(n.e = total1, mean.e = mean1, sd.e = sd1, n.c = total2,
mean.c = mean2, sd.c = sd2, sm = outcome.measure.merged,
studlab = study.name, data = data.ext2[data.ext2$meta.id == u,])
meta.analyses.reg[[counter]] <- limitmeta(meta.analyses[[counter]])</pre>
meta.analyses.trimfill[[counter]] <- trimfill(meta.analyses[[counter]])</pre>
meta.analyses.copas[[counter]] <- auto.copas(meta.analyses[[counter]],</pre>
                                                sig.level = 0.1)
meta.tests.excess[[counter]] <-</pre>
  tes.fct2(data = data.ext2[data.ext2$meta.id == u,])
meta.tests.linreg[[counter]] <- metabias(meta.analyses[[counter]],</pre>
                                            method.bias = "linreg", k.min = 2)
meta.tests.begg[[counter]] <- metabias(meta.analyses[[counter]],</pre>
                                         method.bias = "rank", k.min = 2)
meta.tests.mm[[counter]] <- metabias(meta.analyses[[counter]],</pre>
                                       method.bias = "mm", k.min = 2)
# cont.meta.list <- list(data.ext2, meta.id.vector.cont, meta.analyses,
# meta.analyses.req,
```

```
# meta.analyses.copas,
# meta.analyses.trimfill,
# meta.tests.linreg,
# meta.tests.begg,
# meta.tests.mm,
# meta.tests.excess)
# save(cont.meta.list, file = file.path(PATH_RESULTS, "meta_complete_list_cont.RData"))
#Load analysis data:
# meta.id.vector.cont <- cont.meta.list[[2]]</pre>
# meta.analyses <- cont.meta.list[[3]]</pre>
# meta.analyses.reg <- cont.meta.list[[4]]</pre>
# meta.analyses.copas <- cont.meta.list[[5]]</pre>
# meta.analyses.trimfill <- cont.meta.list[[6]]</pre>
# meta.tests.linreg <- cont.meta.list[[7]]</pre>
# meta.tests.begg <- cont.meta.list[[8]]</pre>
# meta.tests.mm <- cont.meta.list[[9]]</pre>
# meta.tests.excess <- cont.meta.list[[10]]</pre>
#Extract from lists:
cont.results <- data.frame(</pre>
 meta.id = meta.id.vector.cont,
 meta.es.measure = rep("std. mean difference", times = length(meta.id.vector.cont)),
 n.sig.single2 = unlist(lapply(meta.analyses, FUN = function(meta.analysis){length(which(meta.
  #Meta-Analysis
  est.fixef = unlist(lapply(meta.analyses,
                    FUN = function(meta.analysis){meta.analysis$TE.fixed})),
  est.ranef = unlist(lapply(meta.analyses,
                    FUN = function(meta.analysis){meta.analysis$TE.random})),
 zval.fixef = unlist(lapply(meta.analyses,
                    FUN = function(meta.analysis){meta.analysis$zval.fixed})),
 zval.ranef = unlist(lapply(meta.analyses,
                    FUN = function(meta.analysis){meta.analysis$zval.random})),
 pval.fixef = unlist(lapply(meta.analyses,
                    FUN = function(meta.analysis){meta.analysis$pval.fixed})),
 pval.ranef = unlist(lapply(meta.analyses,
                    FUN = function(meta.analysis) {meta.analysis$pval.random})),
 se.est.fixef = unlist(lapply(meta.analyses,
                    FUN = function(meta.analysis){meta.analysis$seTE.fixed})),
 se.est.ranef = unlist(lapply(meta.analyses,
                    FUN = function(meta.analysis) {meta.analysis$seTE.random})),
 Q = unlist(lapply(meta.analyses,
                    FUN = function(meta.analysis){meta.analysis$Q})),
 pval.Q = unlist(lapply(meta.analyses,
```

```
FUN = function(meta.analysis){meta.analysis$pval.Q})),
tau = unlist(lapply(meta.analyses,
                  FUN = function(meta.analysis){meta.analysis$tau})),
k = unlist(lapply(meta.analyses, FUN = function(meta.analysis){meta.analysis$k})),
#Adjustment:
method.reg = unlist(lapply(meta.analyses.reg,
                    FUN = function(meta.adjust){meta.adjust$method.adjust})),
est.reg = unlist(lapply(meta.analyses.reg,
                    FUN = function(meta.adjust) {meta.adjust$TE.adjust})),
se.est.reg = unlist(lapply(meta.analyses.reg,
                    FUN = function(meta.adjust) {meta.adjust$seTE.adjust})),
zval.reg = unlist(lapply(meta.analyses.reg,
                    FUN = function(meta.adjust) {meta.adjust$zval.adjust})),
pval.reg = unlist(lapply(meta.analyses.reg,
                    FUN = function(meta.adjust) {meta.adjust$pval.adjust})),
alpha.r = unlist(lapply(meta.analyses.reg,
                    FUN = function(meta.adjust) {meta.adjust$pval.adjust})),
beta.r = unlist(lapply(meta.analyses.reg,
                    FUN = function(meta.adjust) {meta.adjust$pval.adjust})),
Q.small = unlist(lapply(meta.analyses.reg,
                    FUN = function(meta.adjust){meta.adjust$Q.small})),
Q.resid = unlist(lapply(meta.analyses.reg,
                    FUN = function(meta.adjust){meta.adjust$Q.resid})),
G.squared = unlist(lapply(meta.analyses.reg,
                    FUN = function(meta.adjust){meta.adjust$G.squared})),
est.copas = unlist(lapply(meta.analyses.copas,
                    FUN = function(meta.adjust){meta.adjust[1]})),
se.est.copas = unlist(lapply(meta.analyses.copas,
                    FUN = function(meta.adjust){meta.adjust[2]})),
missing.copas = unlist(lapply(meta.analyses.copas,
                    FUN = function(meta.adjust){meta.adjust[3]})),
#Tests:
pval.d.tes.org = unlist(lapply(meta.tests.excess,
                            FUN = function(object){object[3]})),
pval.d.tes = unlist(lapply(meta.tests.excess,
                            FUN = function(object){object[4]})),
stat.d.tes = unlist(lapply(meta.tests.excess,
                            FUN = function(object){object[2]})),
excess.d = unlist(lapply(meta.tests.excess,
                            FUN = function(object){object[5]})) -
  unlist(lapply(meta.tests.excess,
                            FUN = function(object){object[6]})),
pval.egger = unlist(lapply(meta.tests.linreg,
                           FUN = function(meta.test){meta.test$p.value})),
stat.egger = unlist(lapply(meta.tests.linreg,
                           FUN = function(meta.test){meta.test$statistic})),
pval.begg = unlist(lapply(meta.tests.begg,
                          FUN = function(meta.test){meta.test$p.value})),
```

```
stat.begg = unlist(lapply(meta.tests.begg,
                              FUN = function(meta.test){meta.test$statistic})),
  pval.thompson = unlist(lapply(meta.tests.mm,
                              FUN = function(meta.test){meta.test$p.value})),
  stat.thompson = unlist(lapply(meta.tests.mm,
                              FUN = function(meta.test){meta.test$statistic}))
)
# IV OUTCOME META-ANALYSIS: -----
# # ----- Uncomment to run analysis -----
meta.id.vector.iv <-</pre>
  meta.info.extended$meta.id[which(meta.info.extended$outcome.flag == "IV")]
meta.analyses <- list()</pre>
meta.analyses.reg <- list()</pre>
meta.analyses.copas <- list()</pre>
meta.analyses.trimfill <- list()</pre>
meta.tests.linreg <- list()</pre>
meta.tests.begg <- list()</pre>
meta.tests.mm <- list()</pre>
meta.tests.excess <- list()</pre>
counter <- 0
for(u in meta.id.vector.iv){
counter <- counter + 1</pre>
meta.analyses[[counter]] <- metagen.iv(data = data.ext2[data.ext2$meta.id == u,])</pre>
print(c(u, counter))
meta.analyses.reg[[counter]] <- limitmeta(meta.analyses[[counter]])</pre>
meta.analyses.trimfill[[counter]] <- trimfill(meta.analyses[[counter]])</pre>
meta.analyses.copas[[counter]] <- auto.copas(meta.analyses[[counter]],</pre>
                                                sig.level = 0.1)
meta.tests.excess[[counter]] <-</pre>
  tes.fct2(data = data.ext2[data.ext2$meta.id == u,])
meta.tests.linreg[[counter]] <- metabias(meta.analyses[[counter]],</pre>
                                 method.bias = "linreg", k.min = 2)
meta.tests.begg[[counter]] <- metabias(meta.analyses[[counter]],</pre>
                                 method.bias = "rank", k.min = 2)
meta.tests.mm[[counter]] <- metabias(meta.analyses[[counter]],</pre>
                                 method.bias = "mm", k.min = 2)
\# iv.meta.list \leftarrow list(data.ext2, meta.id.vector.iv, meta.analyses,
   meta.analyses.reg,
   meta.analyses.copas,
   meta.analyses.trimfill,
   meta.tests.linreq,
# meta.tests.begg,
# meta.tests.mm,
```

```
# meta.tests.excess)
# save(iv.meta.list, file = file.path(PATH_RESULTS, "meta_complete_list_iv.RData"))
# #Load analysis data:
# meta.id.vector.iv <- iv.meta.list[[2]]</pre>
# meta.analyses <- iv.meta.list[[3]]</pre>
# meta.analyses.reg <- iv.meta.list[[4]]</pre>
# meta.analyses.copas <- iv.meta.list[[5]]</pre>
# meta.analyses.trimfill <- iv.meta.list[[6]]</pre>
# meta.tests.linreg <- iv.meta.list[[7]]</pre>
# meta.tests.begg <- iv.meta.list[[8]]</pre>
# meta.tests.mm <- iv.meta.list[[9]]</pre>
# meta.tests.excess <- iv.meta.list[[10]]</pre>
#Extract from lists:
iv.results <- data.frame(</pre>
  meta.id = meta.id.vector.iv,
 meta.es.measure = rep(times = length(meta.id.vector.iv), "unknown.effect(IV)"),
  n.sig.single2 = unlist(lapply(meta.analyses,
   FUN = function(meta.analysis){length(which(meta.analysis$pval < 0.05))})),
  #Meta-Analysis
  est.fixef = unlist(lapply(meta.analyses,
    FUN = function(meta.analysis){meta.analysis$TE.fixed})),
  est.ranef = unlist(lapply(meta.analyses,
    FUN = function(meta.analysis){meta.analysis$TE.random})),
  zval.fixef = unlist(lapply(meta.analyses,
    FUN = function(meta.analysis){meta.analysis$zval.fixed})),
  zval.ranef = unlist(lapply(meta.analyses,
    FUN = function(meta.analysis){meta.analysis$zval.random})),
  pval.fixef = unlist(lapply(meta.analyses,
    FUN = function(meta.analysis){meta.analysis$pval.fixed})),
  pval.ranef = unlist(lapply(meta.analyses,
    FUN = function(meta.analysis){meta.analysis$pval.random})),
  se.est.fixef = unlist(lapply(meta.analyses,
    FUN = function(meta.analysis){meta.analysis$seTE.fixed})),
  se.est.ranef = unlist(lapply(meta.analyses,
    FUN = function(meta.analysis) {meta.analysis$seTE.random})),
  Q = unlist(lapply(meta.analyses,
    FUN = function(meta.analysis){meta.analysis$Q})),
  pval.Q = unlist(lapply(meta.analyses,
    FUN = function(meta.analysis){meta.analysis$pval.Q})),
  tau = unlist(lapply(meta.analyses,
    FUN = function(meta.analysis){meta.analysis$tau})),
  k = unlist(lapply(meta.analyses,
   FUN = function(meta.analysis){meta.analysis$k})),
```

```
#Adjustment:
method.reg = unlist(lapply(meta.analyses.reg,
  FUN = function(meta.adjust){meta.adjust$method.adjust})),
est.reg = unlist(lapply(meta.analyses.reg,
 FUN = function(meta.adjust) {meta.adjust$TE.adjust})),
se.est.reg = unlist(lapply(meta.analyses.reg,
  FUN = function(meta.adjust) {meta.adjust$seTE.adjust})),
zval.reg = unlist(lapply(meta.analyses.reg,
 FUN = function(meta.adjust) {meta.adjust$zval.adjust})),
pval.reg = unlist(lapply(meta.analyses.reg,
  FUN = function(meta.adjust) {meta.adjust$pval.adjust})),
alpha.r = unlist(lapply(meta.analyses.reg,
  FUN = function(meta.adjust) {meta.adjust$pval.adjust})),
beta.r = unlist(lapply(meta.analyses.reg,
  FUN = function(meta.adjust) {meta.adjust$pval.adjust})),
Q.small = unlist(lapply(meta.analyses.reg,
  FUN = function(meta.adjust){meta.adjust$Q.small})),
Q.resid = unlist(lapply(meta.analyses.reg,
 FUN = function(meta.adjust){meta.adjust$Q.resid})),
G.squared = unlist(lapply(meta.analyses.reg,
  FUN = function(meta.adjust){meta.adjust$G.squared})),
est.copas = unlist(lapply(meta.analyses.copas,
                         FUN = function(meta.adjust){meta.adjust[1]})),
se.est.copas = unlist(lapply(meta.analyses.copas,
                         FUN = function(meta.adjust){meta.adjust[2]})),
missing.copas = unlist(lapply(meta.analyses.copas,
                         FUN = function(meta.adjust){meta.adjust[3]})),
#Tests:
pval.d.tes.org = unlist(lapply(meta.tests.excess,
                           FUN = function(object){object[3]})),
pval.d.tes = unlist(lapply(meta.tests.excess,
                           FUN = function(object){object[4]})),
stat.d.tes = unlist(lapply(meta.tests.excess,
                           FUN = function(object){object[2]})),
excess.d = unlist(lapply(meta.tests.excess,
                           FUN = function(object){object[5]})) -
  unlist(lapply(meta.tests.excess,
                FUN = function(object){object[6]})),
pval.egger = unlist(lapply(meta.tests.linreg,
                           FUN = function(meta.test){meta.test$p.value})),
stat.egger = unlist(lapply(meta.tests.linreg,
                           FUN = function(meta.test){meta.test$statistic})),
pval.begg = unlist(lapply(meta.tests.begg,
                           FUN = function(meta.test){meta.test$p.value})),
stat.begg = unlist(lapply(meta.tests.begg,
                           FUN = function(meta.test){meta.test$statistic})),
pval.thompson = unlist(lapply(meta.tests.mm,
                           FUN = function(meta.test){meta.test$p.value})),
```

```
stat.thompson = unlist(lapply(meta.tests.mm,
                               FUN = function(meta.test) {meta.test$statistic}))
)
# Z-SCORE BASED META-ANALYSES: -----
# # ----- Uncomment to run analysis -----
meta.id.vector.noiv <-</pre>
  meta.info.extended$meta.id[-which(meta.info.extended$outcome.flag == "IV")]
meta.id.vector.smd <-</pre>
  meta.info.extended$meta.id[which(meta.info.extended$outcome.measure.merged == "SMD")]
meta.id.vector.z <- union(meta.id.vector.smd, meta.id.vector.noiv)</pre>
meta.id.vector.z.c <- meta.id.vector.z[-c(which(meta.id.vector.z == 104613),</pre>
                                             which(meta.id.vector.z == 104616),
                                             which (meta.id.vector.z == 153452),
                                             which(meta.id.vector.z == 163246),
                       which(meta.id.vector.z == 182958))] #Have zero total1..
tmp.z \leftarrow data.ext2 \%\% group_by(meta.id) \%\% mutate(n = n()) \%\% filter(n > 9) \%\%
  filter(total1 + total2 > 3) %>% filter(total1 != 0 & total2 != 0)
meta.analyses <- list()</pre>
meta.analyses.reg <- list()</pre>
meta.analyses.copas <- list()</pre>
counter <- 0
for(u in meta.id.vector.z.c){
counter <- counter + 1</pre>
print(c(u,counter))
meta.analyses[[counter]] <- metacor(cor = z, n = total1 + total2,</pre>
                    studlab = study.name, tmp.z[tmp.z$meta.id == u,])
meta.analyses.reg[[counter]] <- limitmeta(meta.analyses[[counter]])</pre>
meta.analyses.copas[[counter]] <- auto.copas(meta.analyses[[counter]],</pre>
                                                sig.level = 0.1)
}
# zscore.meta.list <- list(tmp.z, meta.id.vector.z.c, meta.analyses,
#meta.analyses.reg, meta.analyses.copas)
# save(zscore.meta.list, file = file.path(PATH_RESULTS,
#"meta_complete_list_zscore.RData"))
#Load analysis data:
meta.id.vector.z.c <- zscore.meta.list[[2]]</pre>
meta.analyses <- zscore.meta.list[[3]]</pre>
meta.analyses.reg <- zscore.meta.list[[4]]</pre>
meta.analyses.copas <- zscore.meta.list[[5]]</pre>
#List extraction:
zscore.meta.analysis.estimates <- cbind(
  meta.id = meta.id.vector.z.c,
 est.z.fixef = unlist(lapply(meta.analyses,
```

```
FUN = function(meta.analysis) {meta.analysis$TE.fixed})),
  est.z.ranef = unlist(lapply(meta.analyses,
                   FUN = function(meta.analysis) {meta.analysis$TE.random})),
  se.est.z.fixef = unlist(lapply(meta.analyses,
                   FUN = function(meta.analysis){meta.analysis$seTE.fixed})),
  se.est.z.ranef = unlist(lapply(meta.analyses,
                    FUN = function(meta.analysis) {meta.analysis$seTE.random})),
  est.z.reg = unlist(lapply(meta.analyses.reg,
                    FUN = function(meta.adjust) {meta.adjust$TE.adjust})),
  se.est.z.reg = unlist(lapply(meta.analyses.reg,
                    FUN = function(meta.adjust) {meta.adjust$seTE.adjust})),
  est.z.copas = unlist(lapply(meta.analyses.copas,
                    FUN = function(meta.adjust){meta.adjust[1]})),
  se.est.z.copas = unlist(lapply(meta.analyses.copas,
                    FUN = function(meta.adjust){meta.adjust[2]})))
# HEDGES G BASED META-ANALYSES: -----
# # ----- Uncomment to run analysis -----
meta.id.vector.d <- meta.id.vector.z[-c(which(meta.id.vector.z == 25407))]</pre>
meta.analyses <- list()</pre>
meta.analyses.reg <- list()</pre>
meta.analyses.copas <- list()</pre>
counter <- 0
for(u in meta.id.vector.d){
  counter <- counter + 1
  print(c(u,counter))
 meta.analyses[[counter]] <- metagen.bincont2(data.ext2[data.ext2$meta.id == u,])</pre>
 meta.analyses.reg[[counter]] <- limitmeta(meta.analyses[[counter]])</pre>
  meta.analyses.copas[[counter]] <- auto.copas(meta.analyses[[counter]],</pre>
                                                 sig.level = 0.1)
}
# cohensd.meta.list <- list(data.ext2, meta.id.vector.d, meta.analyses,
#meta.analyses.reg, meta.analyses.copas)
# save(cohensd.meta.list, file = file.path(PATH_RESULTS,
#"meta_complete_list_cohensd.RData"))
#Load analysis data:
meta.id.vector.d <- cohensd.meta.list[[2]]</pre>
meta.analyses <- cohensd.meta.list[[3]]</pre>
meta.analyses.reg <- cohensd.meta.list[[4]]</pre>
meta.analyses.copas <- cohensd.meta.list[[5]]</pre>
#List extraction:
cohensd.meta.analysis.estimates <- cbind(</pre>
```

```
meta.id = meta.id.vector.d,
  est.d.fixef = unlist(lapply(meta.analyses,
                  FUN = function(meta.analysis) {meta.analysis$TE.fixed})),
  est.d.ranef = unlist(lapply(meta.analyses,
                 FUN = function(meta.analysis){meta.analysis$TE.random})),
  se.est.d.fixef = unlist(lapply(meta.analyses,
                  FUN = function(meta.analysis) {meta.analysis$seTE.fixed})),
  se.est.d.ranef = unlist(lapply(meta.analyses,
                  FUN = function(meta.analysis){meta.analysis$seTE.random})),
  est.d.reg = unlist(lapply(meta.analyses.reg,
                  FUN = function(meta.adjust){meta.adjust$TE.adjust})),
  se.est.d.reg = unlist(lapply(meta.analyses.reg,
                  FUN = function(meta.adjust){meta.adjust$seTE.adjust})),
  est.d.copas = unlist(lapply(meta.analyses.copas,
                  FUN = function(meta.adjust){meta.adjust[1]})),
  se.est.d.copas = unlist(lapply(meta.analyses.copas,
                  FUN = function(meta.adjust){meta.adjust[2]})))
# BUILD FINAL DATASET OF RESULTS
meta.tp5 <- bind_rows(bin.results, cont.results, iv.results)</pre>
meta.tp1 <- merge(meta.info.extended, meta.tp5,</pre>
                  by = c("meta.id")) #Temporary versions
meta.tp4 <- merge(by = "meta.id", x = meta.tp1,</pre>
                  y = zscore.meta.analysis.estimates, all.x = T)
meta.f <- merge(by = "meta.id", x = meta.tp4,</pre>
                y = cohensd.meta.analysis.estimates, all.x = T)
# ROUND-UP: DETECT AND REPLACE MISSING VALUES
#se.est.copas.na = 1 -> is missing
meta.f$se.est.copas.na <- ifelse(is.na(meta.f$se.est.copas), 1, 0)</pre>
meta.f$se.est.z.copas.na <- ifelse(is.na(meta.f$se.est.z.copas), 1, 0)</pre>
meta.f$se.est.d.copas.na <- ifelse(is.na(meta.f$se.est.d.copas), 1, 0)</pre>
meta.f$se.est.reg.na <- ifelse(is.na(meta.f$se.est.reg), 1, 0)</pre>
meta.f$se.est.z.reg.na <- ifelse(is.na(meta.f$se.est.z.reg), 1, 0)</pre>
meta.f$se.est.d.reg.na <- ifelse(is.na(meta.f$se.est.d.reg), 1, 0)</pre>
meta.f$est.copas.na <- ifelse(is.na(meta.f$est.copas), 1, 0)</pre>
meta.f$est.z.copas.na <- ifelse(is.na(meta.f$est.z.copas), 1, 0)</pre>
meta.f$est.d.copas.na <- ifelse(is.na(meta.f$est.d.copas), 1, 0)</pre>
meta.f$est.reg.na <- ifelse(is.na(meta.f$est.reg), 1, 0)</pre>
```

```
meta.f$est.z.reg.na <- ifelse(is.na(meta.f$est.z.reg), 1, 0)</pre>
meta.f$est.d.reg.na <- ifelse(is.na(meta.f$est.d.reg), 1, 0)</pre>
#Inpute random effect estimate for missing copas:
copas.names <- c("est.copas", "se.est.copas", "est.z.copas", "se.est.z.copas",
                 "est.d.copas", "se.est.d.copas")
ranef.names <- c("est.ranef", "se.est.ranef", "est.z.ranef", "se.est.z.ranef",</pre>
                 "est.d.ranef", "se.est.d.ranef")
missing.names <- paste(copas.names, ".missing", sep = "")</pre>
missing <- c()
for(u in 1:length(copas.names)){
 missing.count <- 0
 for(k in 1:dim(meta.f)[1]){
    if(is.na(meta.f[k, copas.names[u]])){
      missing.count <- missing.count + 1</pre>
      meta.f[k, copas.names[u]] <- meta.f[k, ranef.names[u]]</pre>
    }
 }
 missing[u] <- missing.count</pre>
 meta.f[, missing.names[u]] <- missing.count</pre>
#Some useful variables:
meta.f <- meta.f %>% rowwise() %>%
 mutate(pval1.egger = onesided.p(stat = stat.egger, side = side, n = k,
                                   test.type = "reg"),
         pval1.thompson = onesided.p(stat = stat.thompson, side = side, n = k,
                                     test.type = "reg"),
         pval1.begg = onesided.p(stat = stat.begg, side = side, n = k,
                                  test.type = "rank"),
         pval1.harbord = onesided.p(stat = stat.harbord, side = side, n = k,
                                    test.type = "reg"),
         pval1.rucker = onesided.p(stat = stat.rucker, side = side, n = k,
                                    test.type = "reg"),
         pval1.rucker.linreg = onesided.p(stat = stat.rucker.linreg, side = side, n = k,
                                           test.type = "reg"),
         pval1.peter = onesided.p(stat = stat.peter, side = side, n = k,
                                   test.type = "reg"),
         pval1.schwarzer = onesided.p(stat = stat.schwarzer, side = side, n = k,
                                       test.type = "rank"),
         pval1.d.tes = pval.d.tes)
#Summarise different tests for binary and continuous variables to one:
meta.f <- meta.f %>% rowwise() %>%
  mutate(I2 = max(0, (Q - k + 1)/Q),
         var.ratio = (se.max^2)/(se.min^2),
         pval.se = case_when(outcome.flag == "DICH" ~ pval1.rucker.linreg,
```

```
TRUE ~ pval1.egger),
         pval.se.het = case_when(outcome.flag == "DICH" ~ pval1.rucker,
                                  TRUE ~ pval1.thompson))
#significance of pb tests:
sig.level <- 0.1
meta.f <- meta.f %>% rowwise() %>%
  mutate(egger.test = ifelse(pval1.egger < sig.level, 1, 0),</pre>
         thompson.test = ifelse(pval1.thompson < sig.level, 1, 0),
         begg.test = ifelse(pval1.begg < sig.level, 1, 0),</pre>
         tes.d.test = ifelse(pval1.d.tes < sig.level, 1, 0),</pre>
         schwarzer.test = ifelse(pval1.schwarzer < sig.level, 1, 0),</pre>
         rucker.test = ifelse(pval1.rucker < sig.level, 1, 0),</pre>
         rucker.test.linreg = ifelse(pval1.rucker.linreg < sig.level, 1, 0),</pre>
         harbord.test = ifelse(pval1.harbord < sig.level, 1, 0),</pre>
         peter.test = ifelse(pval1.peter < sig.level, 1, 0))</pre>
#calculate test statistics:
meta.f <- meta.f %>% mutate(
  z.fixef = (est.fixef/se.est.fixef)
  z.ranef = (est.ranef/se.est.ranef),
  z.reg = (est.reg/se.est.reg),
  z.copas = ((est.copas)/se.est.copas))
#Separate outcome.types:
meta.bin <- meta.f %>% filter(outcome.flag == "DICH")
meta.cont <- meta.f %>% filter(outcome.flag == "CONT")
meta.iv <- meta.f %>% filter(outcome.flag == "IV")
#Save the result datasets:
save(meta.f, file = file.path(PATH_RESULTS, "meta_analyses_summary_complete.RData"))
save(meta.bin, file = file.path(PATH_RESULTS, "meta.bin.RData"))
save(meta.cont, file = file.path(PATH_RESULTS, "meta.cont.RData"))
save(meta.iv, file = file.path(PATH_RESULTS, "meta.iv.RData"))
save(meta.id.vector, file = file.path(PATH_RESULTS, "meta_id_vector.RData"))
save(data.ext2, file = file.path(PATH_RESULTS, "data_used_for_analysis.RData")) #Save data
# x <- foreach(i = 1:length(meta.id.vector.bin)) %dopar%
# # u <- meta.id.vector.bin[i]</pre>
# # counter <- counter + 1</pre>
  metabin(event.e = events1c, n.e = total1, event.c = events2c, n.c = total2,
           studlab = study.name, sm = "RR",
            method = "Inverse", data = data.ext2[data.ext2$meta.id == meta.id.vector.bin[i]
# cl<-makeCluster(7)</pre>
```

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