Multilevel Meta-Analysis Tutorial

Hu, M., Koh, P.S., Soh, X.C., Hartanto, A., Majeed, N.M.

Introduction

The following script is designed to perform a traditional meta-analysis using the metafor package in R.

The analysis is based on data from Hartanto et al. (2024) and focuses on the relationship between smart-phone presence and cognitive functions. The original paper can be found here: https://doi.org/10.1037/tmb0000123.

The script includes steps for data preparation, effect size calculation, overall effect size computation, forest plot generation, tests for publication bias, and moderation analysis.

The script is structured to be run in RStudio, and it includes comments to guide users through each step of the process.

Setting Up

This section sets up the working environment, installs necessary packages, loads the required libraries, and reads in the data.

If the metafor and lmerTest package is not already installed, use the install.packages() function to install it.

- The setwd() function sets the working directory to the location of the script, ensuring that all file paths are relative to the script's location.
- The library() function loads the metafor package.
- The options() function is used to adjust the display settings, specifically to disable scientific notation and set the number of digits displayed.
- The read.csv() function reads in the data from a CSV file named "SPC.csv", which contains the data drawn from Hartanto et al. (2024).
- A new column named "ID" is created in the data frame to assign unique IDs to each row of data.

```
### Set Up ------
# Set working directory to that of script's current location
setwd(dirname(rstudioapi::getActiveDocumentContext()$path))
# R version 4.5.0
```

```
# Load packages
library(metafor) # version 4.8-0
## Loading required package: Matrix
## Loading required package: metadat
## Loading required package: numDeriv
## Loading the 'metafor' package (version 4.8-0). For an
## introduction to the package please type: help(metafor)
library(lmerTest) # version 3.1-3
## Loading required package: lme4
##
## Attaching package: 'lmerTest'
## The following object is masked from 'package:lme4':
##
##
       lmer
## The following object is masked from 'package:stats':
##
##
       step
# Display settings (to disable scientific notation)
options(scipen = 9999, digits = 4)
# Read in data drawn from Hartanto et al. 2024
multilevelmeta_raw = read.csv("SPC.csv")
# Create new column with unique IDs
multilevelmeta_raw$ID = 1:nrow(multilevelmeta_raw)
```

Prepare Data

This section prepares the data for analysis by computing effect sizes for each study and organizing the data frame.

The multimeta_raw data frame does not include pre-computed effect sizes for each study. Therefore, the escalc() function from metafor package is used to calculate them.

For more information on the escalc() function, refer to ?escalc in R.

Explanation of the Code

- The measure function specifies the type of effect size to be calculated, in this case, "SMD" (Standardized Mean Difference).
- The n1i and n2i functions specify the columns for the sample sizes of each group
- The m1i and m2i functions specify the columns for the means of each group.
- The sdli and sdli functions specify the columns for the standard deviations of each group.
- Afterwards, the escalc() function computes the effect sizes (yi) and their corresponding sampling variances (vi) for each study.
- The multilevelmeta\$publication variable is converted to a factor with specified levels to ensure that the creativity measure types are ordered correctly in the forest plot.
- The multilevelmeta data frame is then sorted by the type of creativity measure and corresponding effect sizes to facilitate clearer visualization in the forest plot.

```
### Prepare Data -----
multilevelmeta = escalc(
       # Type of effect size measure
       measure = "SMD",
       # Columns for sample size of each group
      n1i = n_p,
      n2i = n_a,
       # Columns for means of each group
       m1i = cog_M_p,
       m2i = cog_M_a,
       # Columns for standard deviation of each group
       sd1i = cog_SD_p,
       sd2i = cog_SD_a,
       # Specify data.frame that the information will be extracted from
       data = multilevelmeta_raw
)
# Convert Publication to a factor with specified levels
multilevelmeta$publication = factor(
      multilevelmeta$publication,
      levels = c("Journal article", "Thesis/dissertation", "Conference")
# Order the data frame based on publication
multilevelmeta = multilevelmeta[order(multilevelmeta$publication), ]
```

Computing the Overall Effect Size

This section estimates the overall effect size using the rma.mv() function from the metafor package.

Specifically, we demonstrate the difference in calculating the overall effect size using a traditional meta-analysis approach versus a multilevel meta-analysis approach.

The traditional meta-analysis approach is not recommended for multilevel data, as it does not account for the nested structure of the data (i.e., studies nested within labs).

The multilevel meta-analysis approach is more appropriate for this type of data, as it accounts for the hierarchical structure and allows for the inclusion of random effects.

Explanation of the Multilevel Meta-Analysis Code

- The rma.mv() function is used to compute the overall effect size, accounting for the nested structure of the data.
- The random function specifies the random effects structure, where ~ 1 | lab_id / ID indicates that random intercepts are included for both the lab and individual studies.
- The yi and vi functions specify the effect size estimates and their corresponding sampling variances, respectively.
- The data function specifies the data frame containing the effect size estimates and variances.
- The summary() function is used to display the results of the meta-analysis, including the overall effect size estimate and its confidence interval.

```
##
## Random-Effects Model (k = 136; tau^2 estimator: REML)
##
##
                            AIC
                                      BIC
                                                AICc
    logLik deviance
##
   -5.3141
              10.6282
                        14.6282
                                  20.4387
                                             14.7191
##
## tau^2 (estimated amount of total heterogeneity): 0.0106 (SE = 0.0049)
## tau (square root of estimated tau^2 value):
                                                     0.1027
## I^2 (total heterogeneity / total variability):
                                                     25.36%
## H^2 (total variability / sampling variability):
## Test for Heterogeneity:
## Q(df = 135) = 213.5023, p-val < .0001
##
## Model Results:
##
## estimate
                        zval
                                pval
                                         ci.lb
                                                  ci.ub
                 se
  -0.0481 0.0181 -2.6651 0.0077
                                      -0.0835 -0.0127
```

```
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# Using multilevel code
# The code below is the recommended approach for multilevel meta-analysis
mlmmetaresults = rma.mv(
       # Effect size estimates
      yi = yi,
       # Sampling variances
      V = vi,
       # Include random effects for grouping variable (i.e., lab)
      random = ~ 1 | lab_id / ID,
       # Specify where to get the data from
       data = multilevelmeta
)
# summary function used to provide detailed results of the meta-analysis
summary(mlmmetaresults)
##
```

```
## Multivariate Meta-Analysis Model (k = 136; method: REML)
##
                            AIC
                                      BIC
                                               AICc
##
     logLik Deviance
     3.5443
             -7.0885
                        -1.0885
##
                                   7.6273
                                            -0.9053
##
## Variance Components:
##
##
               estim
                        sqrt nlvls fixed
                                               factor
## sigma^2.1 0.0305 0.1747
                                 22
                                               lab id
                                        no
## sigma^2.2 0.0000
                     0.0000
                                136
                                        no
                                           lab_id/ID
##
## Test for Heterogeneity:
## Q(df = 135) = 213.5023, p-val < .0001
##
## Model Results:
##
## estimate
                                        ci.lb
                        zval
                                pval
                                                ci.ub
                 se
   -0.0315 0.0498 -0.6335 0.5264
                                     -0.1291
                                               0.0660
##
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
```

Forest Plot

This section generates a forest plot to visually represent the effect sizes and confidence intervals for each study included in the meta-analysis.

The forest plot is created using the forest() function from the metafor package.

The plot includes the following features:

• Arrangement of studies by effect sizes

- Sample size information for both dyslexia and control groups
- Custom headers for the plot
- Custom labels for the studies

Saving the forest plot as a PDF file allows for easy sharing and presentation, and allows adjustment to the plot's dimensions.

- The pdf() function starts the graphics device driver to create PDF files, and the file function specifies the name of the file.
- The width and height function adjust the dimensions of the PDF file.
- The forest() function is used to create the forest plot, and the tradmetaresults object contains the results of the meta-analysis.
- The order function specifies the arrangement of studies, with "obs" indicating that the studies should be arranged by effect sizes. To organise by column, replace "obs" with the specific column in the data frame.
- The ylim function sets the y-axis limits for the plot.
- The ilab function is used to add sample size information for both dyslexia and control groups into the forest plot.
- The cbind() function combines the columns indicating the sample size of the groups (dys and control).
- The ilab.xpos function specifies the horizontal arrangement of the columns.
- The slab function is used to label each effect size with its respective study, using the paste() function to combine the "Paper" and "Study" columns. The paste() function creates the label, and the sep function specifies the separator between the columns.
- The xlim function sets the x-axis limits for the plot.
- The alim function sets the confidence interval limits, and the steps function determines the number of intervals in the x-axis.
- The efac function changes the size of effect size polygons.
- The header function is set to FALSE to allow for manual specification of headers.
- The text() function is used to manually include text within the plot, such as the "Author(s) Year" header and specific sample size column headers.
- The x and y function in the text() function adjust the position of the headers, with the x function specifying the horizontal arrangement of the columns and the y function specifying the vertical arrangement of the columns.
- The x function specifies the horizontal arrangement of the columns and the y function specifies the vertical arrangement of the columns.
- The font function adjusts the font size.
- The pos function in the text() function specifies the position of the text relative to the specified coordinates.

- The font function adjusts the font size.
- The dev. off() function is used to close the graphics device and finalize the plot as a saved file.

```
### Forest Plot -----
# Save the forest plot as a PDF file
# Name the pdf file of the forest plot
pdf(file = "mlmforestplot.pdf", width = 15, height = 40)
# Start creating the forest plot itself
# Specify dataset
forest(
       mlmmetaresults,
       # Arrangement of studies
       order = "obs",
       # Add y-axis limits
      ylim = c(-3, 140),
       # Add sample size information for presence and absence of smartphones group
       # -3 for presence of smartphones (n_p)
       # -2.55 for absence of smartphones (n_a)
      ilab = cbind(n_p, n_a),
      ilab.xpos = c(-3, -2.55),
       # Label studies on the forest plot
       slab = paste(author, year_published, sep = ", "),
       \# Add x-axis limits
      xlim = c(-5, 3),
       # Add confidence interval limits
       # Adjust intervals based on the number of steps
       alim = c(-2, 2),
       steps = 9,
       # Change size of effect size polygons
      efac = 0.3,
       # Show (TRUE) or hide (FALSE) default headers
       # Hide when we want to manually specify our own headers
      header = FALSE,
       # Add label for confidence interval, in this case, "Hedge's g"
       xlab = "Hedge's g"
)
# For the following lines of code,
# Use text function to manually include text within the plot
# Add "Author(s) Year" header
text(x = -4.6, y = 139, "Author(s) Year", font = 2)
```

```
# Add "Sample Size" header
text(x = -2.8, y = 139.6, "Sample Size", font = 2)

# Add specific sample size column headers, "Presence" and "Absence"
# x = -3 for for presence of smartphones
# x = -2.55 for absence of smartphones
# y = 139 for both
text(c(x = -3, x = -2.55), y = 139, c("Presence", "Absence"), font = 2)

# Add "g [95% CI]" header
text(x = 2.7, y = 139, "g [95% CI]", font = 2)

# Close the forest plot and finalise it as a saved file
dev.off()
```

pdf ## 2

Tests for Publication Bias

This section performs tests for publication bias, including a funnel plot, rank correlation test and Egger's test.

Funnel Plot

The funnel plot visually represents the distribution of effect sizes and their standard errors, allowing for the identification of potential publication bias. Saving the funnel plot as a PDF file allows for easy sharing and presentation, and allows adjustment to the plot's dimensions.

- The pdf() function starts the graphics device driver to create PDF files, and the file function specifies the name of the file.
- The width and height function adjust the dimensions of the PDF file.
- The funnel() function is used to create the funnel plot, and the tradmetaresults object contains the results of the meta-analysis.
- The legend function specifies whether to include a legend in the plot. TRUE indicates that a legend should be included, FALSE indicates that it should not.
- The xlab function specifies the confidence interval label for the funnel plot, in this case, "Hedge's g"
- The dev.off() function is used to close the graphics device and finalize the plot as a saved file.

```
### Tests for Publication Bias -----
## Funnel Plot
# Save the funnel plot as a PDF file
```

```
# Name the pdf file of the funnel plot
# Adjust the width and height of the pdf file
pdf(file = "mlmfunnelplot.pdf", width = 8, height = 5)
# funnel function to create the funnel plot, specify the data to create the plot
funnel(mlmmetaresults, legend = TRUE, xlab = "Hedge's g")
# Close the funnel plot and finalise it as a saved file
dev.off()
```

```
## pdf
## 2
```

Rank Correlation Test

The rank correlation test serves as a complementary method to the funnel plot in assessing publication bias by examining the correlation between effect sizes and their standard errors.

Explanation of the Code

- The ranktest() function computes the Kendall tau value, which indicates the strength and direction of the association between the ranks of effect sizes and their standard errors.
- The mlmmetaresults object contains the results of the meta-analysis, and the ranktest() function uses this object to perform the rank correlation test.

```
### Checking for Publication Bias -----
## Rank Correlation Test
ranktest(mlmmetaresults)

##
## Rank Correlation Test for Funnel Plot Asymmetry
##
## Kendall's tau = 0.0684, p = 0.2387
```

Egger's Test

The Egger's test is a statistical test that quantifies the degree of asymmetry in the funnel plot, providing a more formal assessment of publication bias.

- The multilevelmeta\$sei_corrected variable is created to store the corrected standard error for each effect size, calculated using the formula $\sqrt{\frac{(n_p+n_a)}{(n_p\cdot n_a)}}$. Using Egger' test (unadjusted) on SMDs results in inflated type 1 error as SMD and SE are not independent. Hence, use corrected formula for SE.
- The lmer() function is used to fit a linear mixed-effects model, where the effect size weighted by the corrected standard error is predicted by the intercept and the inverse of the corrected standard error. metafor::rma.mv does not have a weights argument, and metafor::regtest does not support rma.mv objects. For three (or more) level meta-analysis, use lmerTest::lmer instead.

- The I(yi / sei_corrected) expression indicates that the effect size (yi) is divided by the corrected standard error (sei_corrected), which is used to weight the effect sizes in the model.
- The I(1 / sei_corrected) expression indicates that the inverse of the corrected standard error is included in the model as a predictor.
- The 1 | lab_id expression indicates that random intercepts are included for each lab, accounting for the nested structure of the data.
- The data function specifies the dataset to be used for the analysis.
- The summary() function provides the results of the Egger's test, including the slope estimate and its significance.

```
## Linear mixed model fit by REML. t-tests use Satterthwaite's method [
## lmerModLmerTest]
## Formula: I(yi/sei_corrected) ~ 1 + I(1/sei_corrected) + (1 | lab_id)
      Data: multilevelmeta
##
##
## REML criterion at convergence: 444.7
##
## Scaled residuals:
##
      Min
               1Q Median
                                3Q
                                       Max
## -2.6417 -0.7225 0.0334 0.7713 2.6991
##
## Random effects:
##
  Groups
                         Variance Std.Dev.
            Name
  lab_id
             (Intercept) 0.445
                                  0.667
                         1.335
## Residual
                                  1.155
## Number of obs: 136, groups: lab_id, 22
##
## Fixed effects:
                      Estimate Std. Error
##
                                               df t value Pr(>|t|)
## (Intercept)
                        0.6753
                                   0.4536 61.7763
                                                     1.49
                                                             0.142
## I(1/sei_corrected) -0.1803
                                   0.0874 79.4062
                                                    -2.06
                                                             0.042 *
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
```

Moderation Analysis

This section performs moderation analysis to examine the influence of categorical and continuous moderators on the effect sizes.

Explanation of the Code

- The rma.mv() function is used to compute the multilevel meta-analysis with moderators.
- The yi and vi functions specify the effect size estimates and their corresponding sampling variances, respectively.
- The random function specifies the random effects structure, where ~ 1 | lab_id / ID indicates that random intercepts are included for both the lab and individual studies, accounting for the nested structure of the data.
- The mods function specifies the moderator variable, which can be either continuous or categorical.
- The method function specifies the method used to estimate heterogeneity, in this case, "REML" (Restricted Maximum Likelihood).
- The subset function is used to specify the subset of data for each moderator analysis.
- The data function specifies the data frame containing the effect size estimates and variances.
- The summary() function is used to display the results of the meta-analysis, including the overall effect size estimate and its confidence interval.

Note that if convergence issues arise, the control function can be used to address it. Researchers may also refer more to the metafor package documentation for more information on how to address convergence issues.

```
## Test of Moderators ------

# Categorical variable (i.e., publication)
rma.mv(

yi = yi,
    V = vi,
    random = ~ 1 | lab_id / ID,
    # Specify categorical moderator (i.e., Journal Article)
    subset = (publication == "Journal article"),
    data = multilevelmeta,
    # To address convergence issues (if it exists)
    control=list(rel.tol=1e-8)
)
```

```
##
## Multivariate Meta-Analysis Model (k = 66; method: REML)
##
## Variance Components:
##
## estim sqrt nlvls fixed factor
## sigma^2.1 0.0270 0.1644 14 no lab_id
## sigma^2.2 0.0000 0.0000 66 no lab_id/ID
```

```
##
## Test for Heterogeneity:
## Q(df = 65) = 100.7856, p-val = 0.0029
## Model Results:
##
## estimate
              se
                       zval
                            pval ci.lb ci.ub
## -0.0476 0.0571 -0.8328 0.4050 -0.1596 0.0644
##
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
rma.mv(
      yi = yi,
      V = vi,
      random = ~ 1 | lab_id / ID,
      # Specify categorical moderator (i.e., Thesis/dissertation)
      subset = (publication == "Thesis/dissertation"),
      data = multilevelmeta
)
## Multivariate Meta-Analysis Model (k = 68; method: REML)
## Variance Components:
##
                                             factor
              estim
                       sqrt nlvls fixed
## sigma^2.1 0.0321 0.1792
                             8
                                   no
                                             lab_id
## sigma^2.2 0.0029 0.0543
                                68
                                     no lab_id/ID
## Test for Heterogeneity:
## Q(df = 67) = 100.9032, p-val = 0.0047
##
## Model Results:
##
## estimate
                                     ci.lb
                se
                      zval
                             pval
## 0.0131 0.0872 0.1506 0.8803 -0.1578 0.1841
##
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
rma.mv(
      yi = yi,
      V = vi,
      random = ~ 1 | lab_id / ID,
      # Specify categorical moderator (i.e., Conference)
      subset = (publication == "Conference"),
      data = multilevelmeta
)
## Warning: Single-level factor(s) found in 'random' argument. Corresponding
## 'sigma2' value(s) fixed to 0.
```

```
## Multivariate Meta-Analysis Model (k = 2; method: REML)
## Variance Components:
##
##
                       sqrt nlvls fixed
              estim
                                              factor
## sigma^2.1 0.0000 0.0000
                             1
                                      yes
                                              lab id
## sigma^2.2 0.2598 0.5097
                                 2
                                       no lab_id/ID
##
## Test for Heterogeneity:
## Q(df = 1) = 4.3422, p-val = 0.0372
## Model Results:
##
## estimate
              se
                       zval
                               pval
                                       ci.lb ci.ub
## -0.2118  0.4108  -0.5155  0.6062  -1.0170  0.5934
##
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
# Continuous variable (i.e., female proportion)
rma.mv(
      yi = yi,
      V = vi,
      random = ~ 1 | lab_id / ID,
       # Specify continuous moderator (i.e., female proportion)
      mods = ~female_proportion,
      method = "REML",
      data = multilevelmeta
) |>
      summary()
## Multivariate Meta-Analysis Model (k = 136; method: REML)
##
##
     logLik Deviance
                           AIC
                                     BIC
                                              AICc
##
    5.4145 -10.8291
                      -2.8291
                                  8.7623
                                           -2.5190
## Variance Components:
##
##
              estim
                       sqrt nlvls fixed
                                              factor
## sigma^2.1 0.0246 0.1567
                                22
                                              lab_id
                                       no
## sigma^2.2 0.0000 0.0000
                               136
                                       no lab_id/ID
##
## Test for Residual Heterogeneity:
## QE(df = 134) = 200.0850, p-val = 0.0002
## Test of Moderators (coefficient 2):
## QM(df = 1) = 5.1951, p-val = 0.0227
##
## Model Results:
##
##
                     estimate
                                   se
                                          zval
                                                  pval
                                                          ci.lb
                                                                   ci.ub
                      -0.3041 0.1286 -2.3650 0.0180 -0.5562 -0.0521 *
## intrcpt
```

```
## female_proportion 0.4104 0.1801 2.2793 0.0227 0.0575 0.7634 *
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Forest Plot with Moderators

This section generates a forest plot that includes moderators to visually represent the effect sizes and confidence intervals for each study included in the meta-analysis.

The plot is saved as a PDF file for easy sharing and presentation.

The forest plot includes the following features:

- Arrangement of studies by effect sizes and types of moderators
- Sample size information for both dyslexia and control groups
- Custom headers for the plot
- Custom labels for the studies
- Summary effect sizes for each moderator

- The pdf() function starts the graphics device driver to create PDF files, and the file function specifies the name of the file.
- The width and height function adjust the dimensions of the PDF file.
- The forest() function is used to create the forest plot, and the mlmmetaresults object contains the results of the meta-analysis.
- The rows function specifies the arrangement of studies, with the order of rows adjusted to group studies by publication type.
- The ylim function sets the y-axis limits for the plot.
- The ilab function is used to add sample size information for both dyslexia and control groups into the forest plot.
- The cbind() function combines the columns indicating the sample size of the groups (dys and control).
- The ilab.xpos function specifies the horizontal arrangement of the columns, while the xlim function sets the x-axis limits for the plot.
- The slab function is used to label each effect size with its respective study, using the paste() function to combine the "Paper" and "Study" columns.
- The paste() function creates the label, and the sep function specifies the separator between the columns.
- The alim function sets the confidence interval limits, and the steps function determines the number of intervals in the x-axis.
- The header function is set to FALSE to allow for manual specification of headers.

- The xlab function specifies the label for the confidence interval, in this case, "Hedge's g".
- The text() function is used to manually include text within the plot, such as the "Author(s) Year" header and specific sample size column headers.
- The x and y functions in the text() function adjust the position of the headers.
- The x function specifies the horizontal arrangement of the columns and the y function specifies the vertical arrangement of the columns.
- The font function adjusts the font size.
- The pos function specifies the position of the text relative to the specified coordinates
- The rma.mv() function is used to compute the multilevel meta-analysis with moderators.
- The rest.j, res.t, and res.c variables store the results of the moderation analysis for journal articles, thesis/dissertations, and conference papers, respectively.
- The subset function is used to specify the subset of data for categorical moderators, allowing for separate analyses for each category.
- The addpoly() function is used to add summary effect sizes for each of the moderators, with the row function specifying the position of the summary in the plot.
- The dev.off() function is used to close the graphics device and finalize the plot as a saved file.

```
### Forest Plot of Moderators -----
# Save the forest plot as a PDF file
# Name the pdf file of the forest plot
# Adjust the width and height of the pdf file
pdf(file = "mlmforestplotwithmoderators.pdf", width = 15, height = 45)
forest(
      mlmmetaresults,
       # Manually arrange effect sizes by creativity measure type
       # - Journal article: Rows 143 to 79
       # - Thesis/Dissertations: Rows 75 to 7
       # - Conference: Rows 3 to 2
       # The arrangement must consider spacing and must end at row 2
       rows = c(143:79, 75:7, 3:2),
       # Add y-axis limits
       ylim = c(-3, 147),
       # Add sample size information for presence and absence of smartphones group
       # -4.2 for presence of smartphones (n_p)
       # -3.6 for absence of smartphones (n_a)
       ilab = cbind(n_p, n_a),
       ilab.xpos = c(-4.2, -3.6),
       # Label studies on the forest plot
       slab = paste(author, year_published, sep = ", "),
       # Add x-axis limits
       xlim = c(-7, 4),
```

```
# Add confidence interval limits
       # Adjust intervals based on the number of steps
       alim = c(-1.5, 1.5),
       steps = 11,
       # Change size of effect size polygons
       efac = 0.3,
       # Remove headers (if any), for manual input
       header = FALSE,
       # Add label for confidence interval, in this case, "Hedge's q"
       xlab = "Hedge's g"
)
# For the following lines of code,
# Use text function to manually include text within the plot
# Add text labels for moderator (type of publication)
# Labels for different creativity task types (Moderator Analysis)
\# - "Journal article" at y = 7
\# - "Thesis/Dissertations" at y = 15
\# - "Non-Conference" at y = 21
text(
       x = -7,
       y = c(4, 76, 144),
       pos = 4,
       c("Conference", "Thesis/dissertation", "Journal article"),
       font = 2
)
# Moderation analysis
res.j = rma.mv(
       уi,
       vi,
       random = ~ 1 | lab_id / ID,
       subset = (publication == "Journal article"),
       data = multilevelmeta,
       # To address convergence issues (if it exists)
       control=list(rel.tol=1e-8)
)
res.t = rma.mv(
       yi,
       νi,
       random = ~ 1 | lab_id / ID,
       subset = (publication == "Thesis/dissertation"),
       data = multilevelmeta
)
res.c = rma.mv(
       yi,
       νi,
       random = ~ 1 | lab_id / ID,
       subset = (publication == "Conference"),
```

```
data = multilevelmeta
## Warning: Single-level factor(s) found in 'random' argument. Corresponding
## 'sigma2' value(s) fixed to 0.
# Add summary effect sizes for each of the moderators
addpoly(res.c, row = 1) # summary effect for "Conference" group
addpoly(res.t, row = 6) # summary effect for "Thesis/Dissertation" group
addpoly(res.j, row = 78) # summary effect for "Journal article" group
# Add"Author(s) Year" header
text(x = -6.5, y = 146, "Author(s) Year", font = 2)
# Add "Sample Size" header
text(x = -3.9, y = 146.7, "Sample Size", font = 2)
# Add specific sample size column headers, "Presence" and "Absence"
# x = -4.2 for for presence of smartphones
# x = -3.6 for absence of smartphones
# y = 146  for both
text(c(x = -4.2, x = -3.6), y = 146, c("Presence", "Absence"), font = 2)
# Add "q [95% CI]" header
text(x = 3.6, y = 146, "g [95\% CI]", font = 2)
# Close the forest plot and finalise it as a saved file
dev.off()
## pdf
```

End of Code

##