orchaRd: An R package for plotting prediction intervals and orchard plots

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Introduction

orchaRd allows users to create pretty orchard plots that contain both confidence and prediction intervals around mean effect size estimates, plots the effect size distribution over top such estimates and weights effect sizes by their precision (1/sampling error) or sample size. orchaRd takes a metafor object of class rma.mv or rma (Viechtbauer, 2010) and plots the results for the meta-analytic or meta-regression model. Currently, only meta-regression models with a single moderator variable are allowed or intercept only meta-analytic models. orchaRd uses ggplot (Wickham, 2009) for plotting, and as such, layers can be added directly to make plots customizable to the users needs.

Citing orchaRd

To cite orchaRd in publications one can use the following reference:

Nakagawa, S. et al. 2020. The Orchard Plot: Cutlivating the Forest Plot for Use in Ecology, Evolution and Beyond. Research Synthesis Methods, in review

Installation

To install orchaRd use the following code in R:

```
install.packages("devtools")
install.packages("tidyverse")
install.packages("metafor")
install.packages("patchwork")
devtools::install_github("itchyshin/orchard_plot", subdir = "orchaRd", force = TRUE)
library(orchaRd)
library(patchwork)
library(tidyverse)
library(metafor)
```

Installation will make the primary functions accessible to users along with their help files. You will also need the tidyverse and metafor packages.

Examples of how it works

In this vignette we'll walk the reader through a number of case studies and show you how you can create beautiful looking orchard plots. We overview three different case studies that make use of different effect sizes and moderators. The datasets associated with each case study come as part of the orchard package.

Example 1: Dietary Restriction and Lifespan

#> ---

English and Uller (2016) performed a systematic review and meta-analysis on the effects of early life dietary restriction (a reduction in a major component of the diet without malnutrition; e.g. caloric restriction) on average at death, using the standardised mean difference (often called d). They found that across the whole dataset, there was little evidence for an effect of dietary restriction on mean age at death. Here we'll use the dataset to first calculate the effect size measures and then fit an intercept only, meta-analytic model.

```
data(english)
# We need to calculate the effect sizes, in this case d
english <- metafor::escalc(measure = "SMD", n1i = NStartControl, sd1i = SD_C, m1i = MeanC,
   n2i = NStartExpt, sd2i = SD_E, m2i = MeanE, data = english)
english_MR_int <- metafor::rma.mv(yi = yi, V = vi, random = list(~1 | EffectID),
   data = english)
summary(english_MR_int)
#>
\#> Multivariate Meta-Analysis Model (k = 77; method: REML)
#>
#>
     logLik Deviance
                           AIC
                                      BIC
                                              AICc
                       97.3748 102.0362
#> -46.6874
            93.3748
                                            97.5391
#>
#> Variance Components:
#>
#>
               estim
                        sqrt nlvls fixed
                                             factor
#> siqma^2
             0.1271
                     0.3566
                                77
                                       no EffectID
#>
#> Test for Heterogeneity:
\#>Q(df=76)=297.4722, p-val<.0001
#>
#> Model Results:
#>
                              pval
#> estimate
            se
                      zval
                                       ci.lb
    0.0239 0.0510 0.4676 0.6401 -0.0761 0.1238
#>
```

We can see from the above that we have fit a meta-analytic model without an intercept and thus the mean estimates is the overall effect size on the effects of caloric restriction on mean death across all studies examined. Now that we have fit our meta-analytic model we can get the confidence intervals and prediction intervals with a few functions in the orchaRd package. If one is interested in getting the table of results we can use the mod_results function. This will allow users to make nice tables of the results if needed. We can do that as follows:

#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

```
model_results <- orchaRd::mod_results(english_MR_int, mod = "Int")</pre>
print(model_results)
        name
               estimate
                             lowerCL
                                       upperCL
                                                   lowerPR
                                                              upperPR
#> 1 Intrcpt 0.02385105 -0.07612047 0.1238226 -0.6821196 0.7298217
If we instead want to create an orchard plot and visualise the results we can do so quite simply as:
orchard_plot(english_MR_int, mod = "Int", xlab = "Standardised mean difference",
    transfm = "none") +
scale_fill_manual(values = "grey") +
scale_colour_manual(values = "grey")
#> Scale for 'fill' is already present. Adding another scale for 'fill', which will replace the
#> existing scale.
#> Scale for 'colour' is already present. Adding another scale for 'colour', which will replace the
```

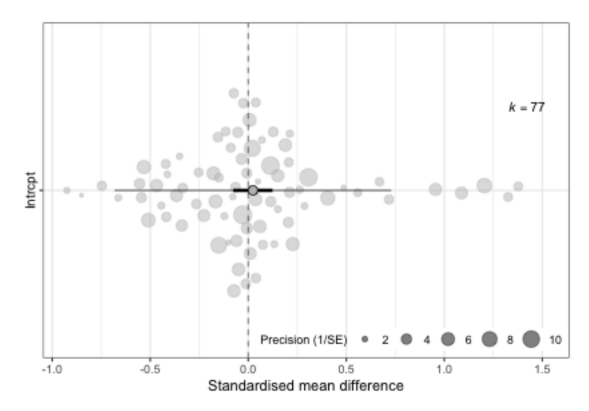


Figure 1: Orchard plot of the impact caloric restriction using standardised mean difference

In 1 we simply add in the metafor model and it will create a default orchard plot. Alternatively, we could also add in the table of results.

```
orchard_plot(model_results, mod = "Int", xlab = "Standardised mean difference", transfm = "none") +
    scale_fill_manual(values = "grey") +
scale_colour_manual(values = "grey")

#> Scale for 'fill' is already present. Adding another scale for 'fill', which will replace the
#> existing scale.

#> Scale for 'colour' is already present. Adding another scale for 'colour', which will replace the
#> existing scale.
```

Fugure 2 and Figure 1 above show that overall estimate from a random-effects meta-analysis of 77 effect sizes is centered on zero, with a 95% CI that spans the line of no-effect. The prediction intervals clearly demonstrate the high level of heterogeneity, with effects size less than -0.5 and greater than 0.5 predicted to be observed.

In a subsequent publication, Senior et al. (2017) analysed this dataset for effects of dietary-restriction on among-individual variation in the age at death using the log coefficient of variation ratio. A major prediction in both English & Uller (2016) and Senior et al. (2017) was that the type pf manipulation, whether the study manipulated quality of food versus the quantity of food would be important. As such, we can fit a meta-regression model to test whether the moderator "Manipulation Type" impacts our results on the mean and variance

```
# First we need to calculate the lnCVR
english$lnCVR <- metafor::escalc(measure = "CVR", n1i = NStartControl, sd1i = SD_C,
    m1i = MeanC, n2i = NStartExpt, sd2i = SD_E, m2i = MeanE, data = english)

# Now we can fit the meta-regression models
english_MR <- metafor::rma.mv(yi = yi, V = vi, mods = ~ManipType - 1, random = list(~1 |
    EffectID), data = english)
summary(english_MR)
#>
#> Multivariate Meta-Analysis Model (k = 77; method: REML)
```

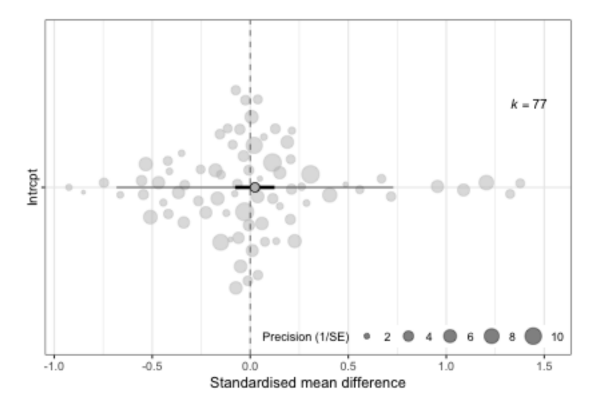


Figure 2: Orchard plot of the impact caloric restriction using standardised mean difference but instead of using the metafor model, using the model results table

```
#>
#>
     logLik Deviance
                            AIC
                                      BIC
                                               AICc
   -46.5719
             93.1437
                        99.1437 106.0962
                                            99.4818
#>
#>
#>
   Variance Components:
#>
                                              factor
#>
                        sqrt nlvls fixed
               estim
              0.1296
                      0.3600
                                       no EffectID
#> sigma^2
                                 77
#>
#> Test for Residual Heterogeneity:
\#> QE(df = 75) = 295.5324, p-val < .0001
#>
#> Test of Moderators (coefficients 1:2):
   QM(df = 2) = 0.2832, p-val = 0.8680
#>
#> Model Results:
#>
#>
                      estimate
                                          zval
                                                  pval
                                                          ci.lb
                                                                  ci.ub
                                    se
#> ManipTypeQuality
                      0.0125 0.0666 0.1874
                                               0.8513
                                                        -0.1181 0.1431
#> ManipTypeQuantity
                     0.0401 0.0805 0.4981
                                               0.6184
                                                        -0.1178 0.1980
#>
#>
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# Again, we can create a table of results
model_results <- orchaRd::mod_results(english_MR, mod = "ManipType")</pre>
print(model_results)
#>
        name estimate
                            lowerCL
                                      upperCL
                                                 lowerPR
#> 1 Quality 0.01248610 -0.1180989 0.1430711 -0.7050164 0.7299886
#> 2 Quantity 0.04011772 -0.1177539 0.1979893 -0.6828489 0.7630844
```

```
# Lets fit a meta-regression - I am modelling non-independence here (article).
senior MR <- metafor::rma.mv(yi = lnCVR, V = vi, mods = ~ManipType - 1, random = list(~1 |
    EffectID), data = english)
\#> Error in metafor::rma.mv(yi = lnCVR, V = vi, mods = ~ManipType - 1, random = list(~1 | : Number of rows of
summary(senior_MR)
#>
#> Multivariate Meta-Analysis Model (k = 77; method: REML)
#>
#>
                                       BIC
     logLik Deviance
                            AIC
                                                AICc
#>
   -33.6583
             67.3166
                        73.3166
                                   80.2690
                                             73.6546
#>
#> Variance Components:
#>
#>
                        sqrt nlvls fixed
               estim
                                               factor
#>
  sigma^2
              0.0726
                      0.2695
                                 77
                                         no
                                            EffectID
#>
#> Test for Residual Heterogeneity:
\#> QE(df = 75) = 215.7242, p-val < .0001
#> Test of Moderators (coefficients 1:2):
  QM(df = 2) = 5.6956, p-val = 0.0580
#>
#> Model Results:
#>
                      estimate
                                     se
                                            zval
                                                    pval
                                                            ci.lb
                                                                     ci.ub
#> ManipTypeQuality
                       -0.1258 0.0529
                                         -2.3805
                                                  0.0173
                                                          -0.2294
                                                                   -0.0222
#> ManipTypeQuantity
                        0.0106 0.0625
                                          0.1695
                                                  0.8654
                                                          -0.1118
                                                                    0.1330
#>
#>
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# creating a table of results
senior_results <- mod_results(senior_MR, "ManipType")</pre>
print(senior_results)
#>
         n.a.me.
                 estimate
                             lowerCL
                                                     lowerPR
                                          upperCL
#> 1 Quality -0.12582843 -0.2294275 -0.02222936 -0.6641038 0.4124469
#> 2 Quantity 0.01058903 -0.1118324 0.13301043 -0.5316237 0.5528017
# We can now plot SMD and lnCVR beside each other and compar ethe results
p1 <- orchard_plot(english_MR, mod = "ManipType", xlab = "Standardised mean difference",
    transfm = "none")
p2 <- orchard_plot(senior_MR, mod = "ManipType", xlab = "log(CV ratio) (lnCVR)",
    transfm = "none")
p1/p2
```

Our orchard plot for the log coefficient of variation demonstrates that, while restrictions on dietary quality and quantity do not affect the average age at death (top of Figure 3), among-individual variation may be altered by quality restrictions (bottom of Figure 3). The effect is negative suggesting that the coefficient of variation in the control group is lower than that in the treatment group, and the 95% CI does not span zero. Again though, the effect is heterogeneous; a substantial number of positive effects are still predicted.

Example 2: Predation and Invertebrate Community

Eklof et al. (2012) evaluated the effects of predation on benthic invertebrate communities. Using the log response ratio they quantified differences in abundance and/or biomass of gastropods and Amphipods in groups with and without predation in

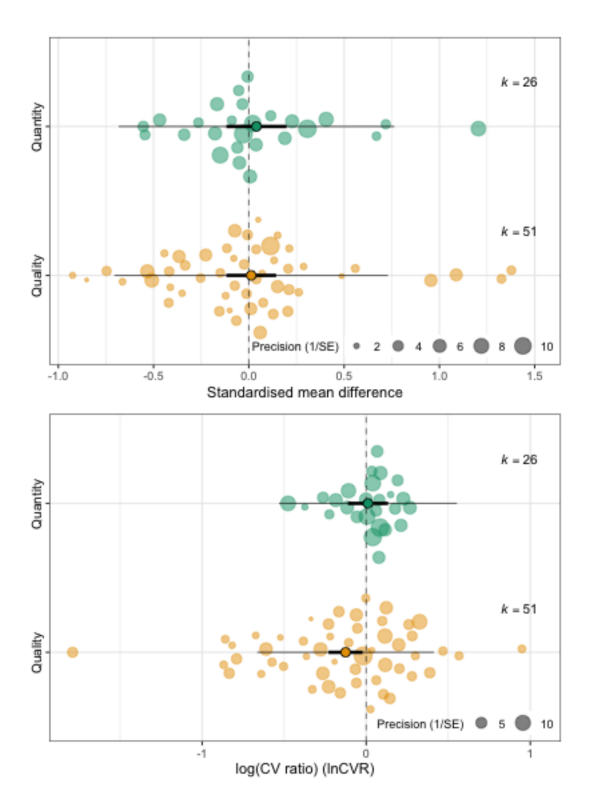


Figure 3: Orchard plot of diet qualities impact on SMD (top) and log coefficient of variation (bottom)

an experimental setting.

Here again, we can create orchard plots of the model results, but we'll show how a few simple things can be modified. Again, we can fit the meta-analytic model first:

```
data(eklof)
```

Above we have fit a meta-regression model using "Grazer Type" as a moderator which is predicted to explain variation in log response ratios. We can demonstrate a few simple changes users can make, but we note here that users can make far more complex changes down the line if needed, but we'll save those for the last example. The first is the angle at which the y-axis labels are positioned (bottom of Figure 4):

The other thing we can change is the type of scaling we wish to use. Lets say we are interested in scaling the effect size by the total sample size of the study we use a vector of N, sample size (bottom of Figure 5):

Overall, our orchard plot shows the results of a re-analysis of their data. The effects are negative for both Gastropods and Amphipods suggesting that mean abundance/biomass in the control group is lower than in the treatment groups, although the effect is largest, and is statistically significant, for Amphipods. In both cases the prediction intervals reveal the extent of heterogeneity, with positive effects predicted to be observed.

Example 3: Maternal-Offspring Morphological Correlations

Finally, we also look at the case discussed by Lim et al. (2014), who meta-analysed the strength of correlation between maternal and offspring size within-species, across a very wide range of taxa. The found, that typically, there is a moderate positive correlation between maternal size and offspring size within species (i.e. larger mothers have larger offspring). However, they also found evidence for relatively strong phylogenetic effects suggesting the strength of the association is dependent on evolutionary lineage.

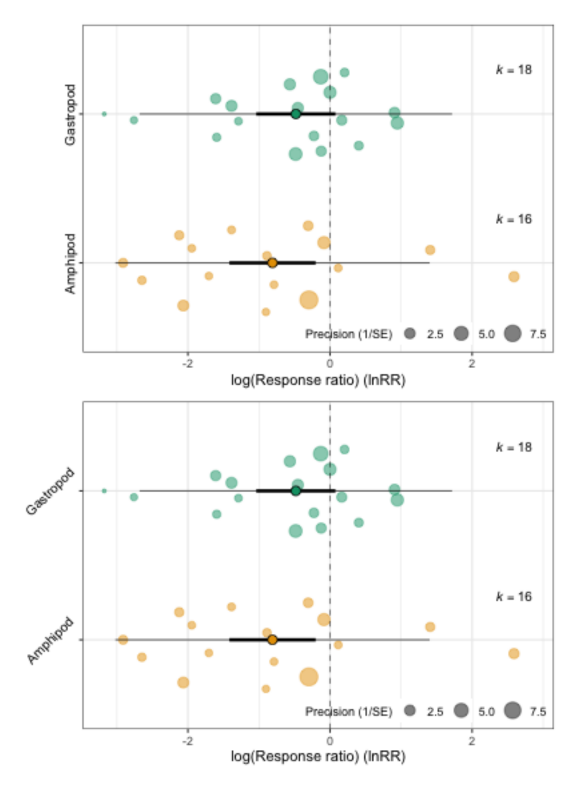


Figure 4: Orchard plots of the effects of predation on benthic invertebrate communities compared using the log response ratio. Top panel is the default plot and bottom panel a plot containing changes in label axes

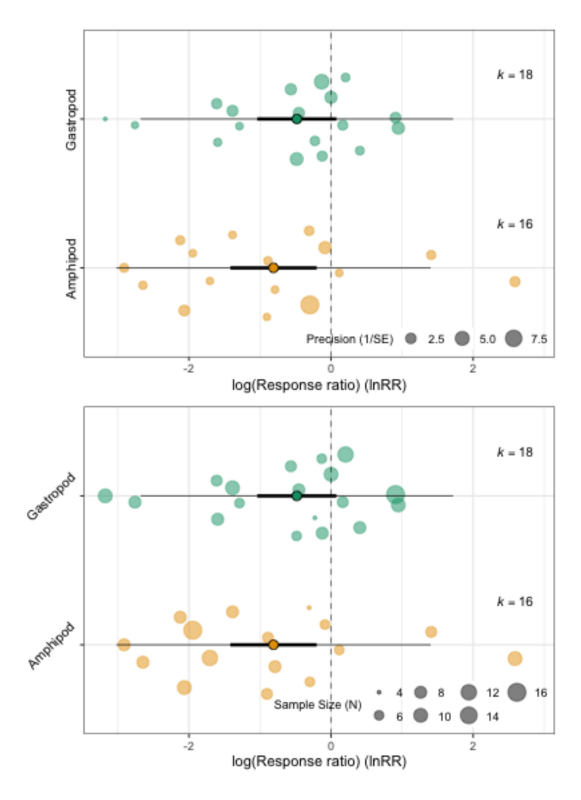


Figure 5: Orchard plots of the effects of predation on benthic invertebrate communities compared using the log response ratio. Top panel is the default plot and bottom panel a plot containing changes in label axes and scaling with sample size instead of precision

Here we have used an orchard plot to represent the results obtained when meta-analysing the data from Lim et al. (2014) by taxonomic Phylum.

```
data(lim)
```

```
# Add in the sampling variance
lim$vi <- (1/sqrt(lim$N - 3))^2
# Lets fit a meta-regression - I will do Article non-independence. The
# phylogenetic model found phylogenetic effects, however, instead we could fit
# Phylum as a fixed effect and explore them with an Orchard Plot
lim_MR <- metafor::rma.mv(yi = yi, V = vi, mods = ~Phylum - 1, random = list(~1 |</pre>
   Article, ~1 | Datapoint), data = lim)
summary(lim_MR)
\#> Multivariate Meta-Analysis Model (k=357; method: REML)
#>
#>
   logLik Deviance
                         AIC
                                    BIC
                                            AICc
#> -97.6524 195.3049 213.3049 248.0263 213.8343
#>
#> Variance Components:
#>
#>
              estim
                      sqrt nlvls fixed
                                            factor
#> sigma^2.1 0.0411 0.2029
                              220
                                      no
                                           Article
#> sigma^2.2 0.0309 0.1757
                              357
                                     no Datapoint
#>
#> Test for Residual Heterogeneity:
\#> QE(df = 350) = 1912.9637, p-val < .0001
#>
#> Test of Moderators (coefficients 1:7):
\#> QM(df = 7) = 356.6775, p-val < .0001
#> Model Results:
#>
#>
                       estimate
                                                         ci.lb
                                                                  ci.ub
                                      se
                                           zval
                                                  pval
                                         5.2829 <.0001 0.1692 0.3687
#> PhylumArthropoda
                        0.2690 0.0509
                         0.3908 0.0224 17.4190 <.0001
#> PhylumChordata
                                                         0.3468 0.4347
                                                                          ***
                        0.8582 0.3902
#> PhylumEchinodermata
                                         2.1992 0.0279
                                                          0.0934
                                                                  1.6230
#> PhylumMollusca
                         0.4867 0.1275
                                         3.8175 0.0001
                                                         0.2368 0.7366
                                                                          ***
#> PhylumNematoda
                        0.4477 0.3054 1.4658 0.1427 -0.1509 1.0463
#> PhylumPlatyhelminthes 0.4935 0.2745
                                          1.7980 0.0722 -0.0444 1.0314
#> PhylumRotifera
                          0.4722 0.3021
                                          1.5634 0.1180 -0.1198 1.0642
#>
#> ---
#> Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Noe we can plot a default orchard plot, scaling each effect size by N. Also, because we are using Zr, we can use transfm = "tanh" and it will do the conversions for us:

Now that we have Figure 6 we can do some small changes to make it pretty. Currently, the cb argument is "FALSE", we can change this to "TRUE" to use colour blind friendly colours. Additionally, because we are using ggplot we can add element to the figure to make it look pretty.

```
orchard_plot(lim_MR, mod = "Phylum", xlab = "Correlation coefficient (r)", alpha = 0.5, transfm = "tanh", theme(legend.position= c(0.05, 0.99),
```

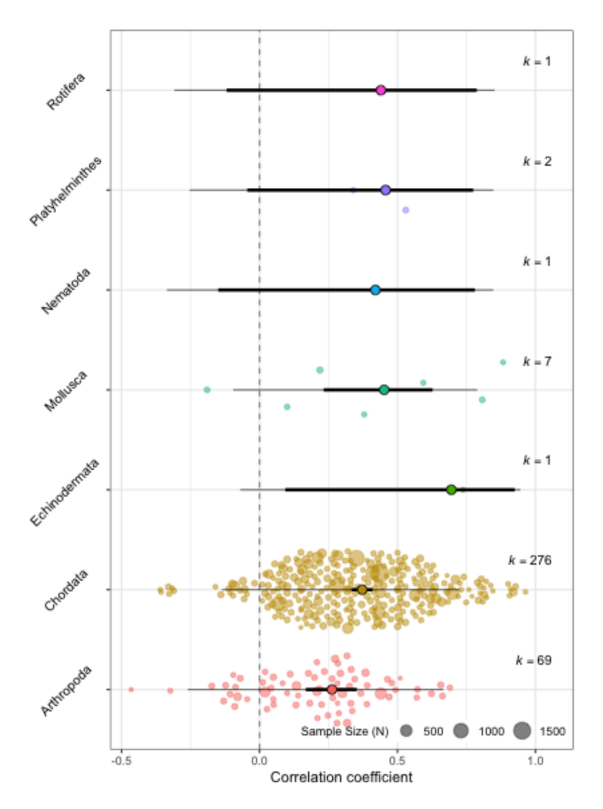


Figure 6: Orchard plot of the the strength of correlation between maternal and offspring size within-species

```
legend.justification = c(0,1),
legend.key.size = unit(1, "mm")) +

theme(legend.direction="horizontal",

legend.title = element_text(size =8),

legend.text = element_text(size = 10)) +

scale_x_continuous(expand = c(0.1,0.1))

#> Warning in if (N != "none") {: the condition has length > 1 and only the first element will be used
```

As in Figure 7, New elements can be added to the orchard_plot to modify it as one sees fit. It will overwrite existing elements. From our orchard plots above, it is clear that the analysis is dominated by data from Chordates and Arthropods, with the other Phyla being much more poorly represented. Second, there is a difference between the strength of a typical correlation within these two well represented groups (the correlation is stronger in Chordates), which arguably would explain the phylogenetic signals detected by Lim et al. (2014). Lastly, although there are differences within the typical correlation between Chordates and Arthropods, there remains a large overlap in predicted range of individual effect sizes; individual species within Phyla are still highly variable.

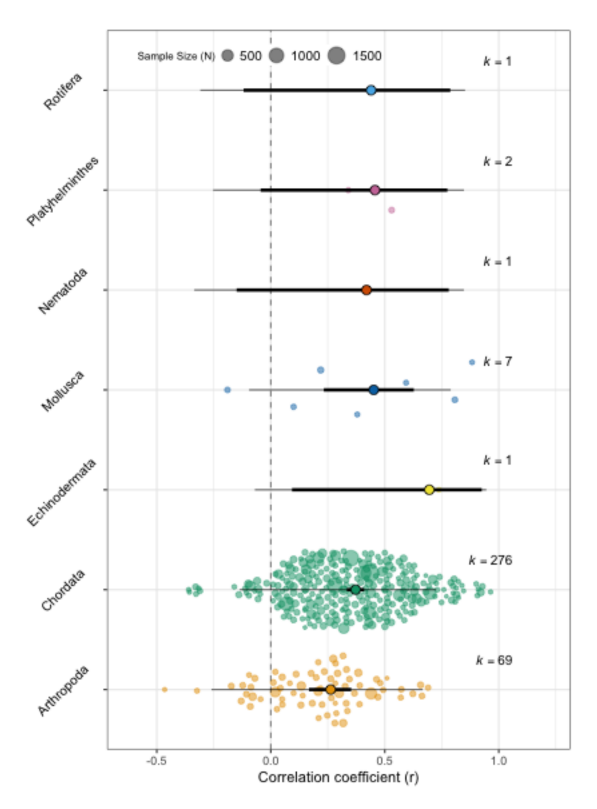


Figure 7: Orchard plot of the the strength of correlation between maternal and offspring size within-species

References

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Wickham H. 2009. ggplot2: elegant graphics for data analysis. New York; London: Springer.