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- ¹⁵ Running Headline: Phylogenetic multilevel meta-analysis

16 Abstract

1. Meta-analyses in ecology and evolution require special attention due to certain study 17 characteristics in these fields. First, the primary articles in these fields usually report results 18 that are observed from studies conducted on different species, and the phylogeny among 19 the species violates the independence assumption. Second, the primary articles frequently 20 report multiple results which cannot be accounted for by conventional meta-analytic models. 21 Although there is a model that accounts for these two problems in theory, its performance 22 has not been examined extensively. In this article, we investigate the performance of this 23 model in comparison with simpler models. 24

25 **2.** We conducted an extensive simulation study where data with different levels of com-26 plexities were generated and then various models were fitted to examine their performance. 27 The models we used include the conventional random-effects and multilevel random-effects 28 models along with more complex multilevel models that account for species-level variance 29 with different variance components. Furthermore, we present an illustrative application of 30 these models based on the data from a meta-analysis on size-assortative mating and comment 31 on the results in light of the findings from the simulation study.

3. Our simulation results show that, when the phylogenetic relationships among the 32 species are at least moderately strong, only the most complex model that decomposes the 33 species-level variance into non-phylogenetic and phylogenetic components provides approxi-34 mately unbiased estimates of the overall mean and variance components and yields confidence 35 intervals with an approximately nominal coverage rate. On the other hand, removing the 36 phylogenetic or non-phylogenetic component leads to biased variance component estimates 37 and an increased risk for incorrect inferences about the overall mean. These findings are 38 supported by the results derived from the illustrative application. 39

4. Based on our results, we suggest that meta-analyses in ecology and evolution should
4. use the model that accounts for both the non-phylogenetic and phylogenetic species-level
4. variance in addition to the multilevel structure of the data. Any attempts to simplify this

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⁴³ model, such as using only the phylogenetic variance component, may lead to erroneous
⁴⁴ inferences from the data.

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Keywords: comparative analysis, mixed-effects models, model efficiency, multilevel
 models, phylogenetic meta-analysis, random-effects variance estimation.

48 1 Introduction

Meta-analysis encompasses an array of methods for synthesizing information from stud-49 ies examining some phenomenon of interest and evaluating the consistency of their results 50 (Glass, 1976; Hedges and Olkin, 1985; Cooper et al., 2009; Senior et al., 2016). Although 51 these methods have been mostly developed in the medical and social sciences (Egger et al., 52 2001; Sutton and Higgins, 2008; Cooper et al., 2009), ecologists and evolutionary biologists 53 have successfully adopted these techniques for conducting research syntheses in their respec-54 tive fields (Gurevitch et al., 2001; Koricheva et al., 2013; Gurevitch et al., 2018). However, 55 meta-analyses in ecology and evolution typically have several features that require special 56 attention so that trustworthy evidence can be obtained. 57

To start, meta-analyses in these fields often incorporate data from multiple species which 58 share an evolutionary history, known as phylogeny (Arnqvist and Wooster, 1995; Gurevitch 59 and Hedges, 1999; Chamberlain et al., 2012). As a result, the samples (and the effect 60 sizes obtained from these samples) are not independent which violates the independence 61 assumption underlying conventional meta-analytic models. For example, the standard fixed-62 and random-effects models (Hedges and Olkin, 1985; Hedges and Vevea, 1998), often used 63 for ecological meta-analyses (Nakagawa and Santos, 2012), assume independence among the 64 effect sizes and therefore do not account for phylogeny (Chamberlain et al., 2012; Noble 65 et al., 2017). This issue was first addressed by Adams (2008) and Lajeunesse (2009) who 66 incorporated phylogenies into the fixed- and random-effects models, respectively. 67

⁶⁸ Chamberlain et al. (2012) empirically investigated how the inclusion of phylogeny af-⁶⁹ fects the estimate of the overall mean based on data from 30 meta-analyses in ecology and ⁷⁰ evolution. While the estimate of the overall mean did not change considerably in most ⁷¹ cases (especially when using a random-effects model), a substantial portion of the meta-⁷² analyses, which reported significant results before, produced non-significant results when ⁷³ the phylogeny was incorporated into the model. Therefore, including phylogeny might be ⁷⁴ an important factor to reduce Type I error rates and to obtain an accurate reflection of the

⁷⁵ uncertainty of meta-analytic estimates.

Although Chamberlain et al. (2012) is the most extensive study to date examining the 76 effects of phylogeny in meta-analysis, their work was based on available meta-analyses. To 77 investigate the issue of phylogeny more broadly, we require a simulation study to explore a 78 wider parameter space and under controlled conditions. Moreover, Chamberlain et al. (2012) 79 did not address the fact that ecological and evolutionary studies usually report multiple effect 80 sizes per study, which leads to another source of non-independence (Nakagawa and Santos, 81 2012; Noble et al., 2017). Although past and current meta-analyses have sometimes avoided 82 this issue by selecting a single effect size from each study or by collapsing multiple effect sizes 83 into one, these procedures can lead to a severe loss of information (Nakagawa and Santos, 84 2012). 85

As an alternative, Hadfield and Nakagawa (2010) proposed a mixed-effects model that 86 accounts for the multilevel structure via a study-level random effect (i.e., multiple effect 87 sizes per study are nested within this random effect). In the same model, they include two 88 additional random effects to estimate the non-phylogenetic and the phylogenetic variance. 89 This way, among-species variance is decomposed into two components, the one resulting 90 from species similarities due to evolutionary history and the other from species similarities 91 due to shared ecology and other factors (Lynch, 1991). Although the model by Hadfield 92 and Nakagawa (2010) addresses two major statistical issues in ecological and evolutionary 93 meta-analyses, the complexity of the model poses certain challenges. 94

Partitioning the species variance into its two components is a challenging endeavor, because both components are modeled using random effects at the species level, with the only difference that the phylogenetic component assumes that the random effects are correlated according to a phylogenetic correlation matrix – which is derived from a phylogenetic tree constructed based on the similarities and differences of species in terms of their (usually) genetic (but sometimes also physical) characteristics (Felsenstein, 2004). This raises concerns about the identifiability of the variance components and potential bias in their estimates, issues that have also been raised outside the meta-analytic context when analyzing the data
of primary studies including multiple species (Paradis, 2012).

Moreover, the complexity of the model poses a threat to the convergence of optimization 104 algorithms (Bates et al., 2015). Accordingly, Nakagawa and Santos (2012) suggested that 105 model fitting may only be feasible with larger datasets, which would limit the applicability 106 of the model in practice. To avoid these problems, some ecological and evolutionary meta-107 analyses have been carried out using a simplified model without the non-phylogenetic random 108 effect and that therefore accounts for species variance only via the phylogenetic component 109 (e.g., Garamszegi et al., 2012; Moore et al., 2016). However, the consequences of doing so, 110 and the performance of the more complex model, has yet to be evaluated in a simulation 111 study. 112

We therefore investigated the performance of models for conducting a phylogenetic mul-113 tilevel meta-analysis in a comprehensive simulation study. We simulate studies that report 114 multiple effect sizes and use several models that vary in their complexity, starting from a 115 simple model (including only a random effect at the effect sizes level) to the most complex 116 model which incorporates a study-level and two among-species random effects. Further, we 117 generate specific conditions to examine the performance of the most complex model when 118 phylogenetic relationships are weak and the consequences of removing the non-phylogenetic 119 component. Finally, we present an illustrative application of these models based on the data 120 from a meta-analysis on size-assortative mating and comment on the results in light of the 121 findings from the simulation study. 122

¹²³ 2 Materials and Methods

¹²⁴ 2.1 Meta-Analytic Models

To conduct a meta-analysis, the phenomenon of interest (e.g., the size of a treatment effect or the strength of the association between two variables) needs to be quantified in terms of an effect size estimate for each study to be included in the analysis. We use the term 'study'
broadly here, as a single study may contribute multiple estimates (e.g., for multiple species,
subgroups, treatments), but for the moment we assume that each study contributes a single
estimate to the meta-analysis.

The specific effect size measure to be used in a meta-analysis depends on the phenomenon 131 of interest and the information reported in the studies (Nakagawa and Santos, 2012). For 132 example, raw or standardized mean differences and response ratios (Hedges et al., 1999) 133 are typically used to quantify group differences or treatment effects based on quantitative 134 variables, correlation coefficients (or Fisher r-to-z transformed values thereof) reflect the 135 (linear) relationship between two variables, while (log-transformed) odds/risk ratios and 136 risk differences (calculated from 2×2 contingency tables) indicate group differences (e.g., 137 treated vs. untreated, exposed vs. non-exposed) with respect to dichotomous dependent 138 variables (e.g., cured vs. not cured, diseased vs. not diseased). For all of these measures, 139 we can also compute the sampling variances of the estimates, that is, the variability in each 140 estimate that would be expected under repeated sampling of new study units under identical 141 circumstances (Nakagawa and Cuthill, 2007; Cooper et al., 2009; Borenstein et al., 2011). 142 Regardless of the specific measure used in a meta-analysis, let y_i denote the effect size 143

estimate for the *i*th study (with $i = 1, ..., N_{studies}$) and v_i the corresponding sampling variance. The most basic model that can be considered for synthesizing the estimates is the fixed-effects model, which is given by

$$y_i = \mu + e_i,\tag{1}$$

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$$\mathbf{e} \sim N(\mathbf{0}, \mathbf{V}),\tag{2}$$

where μ is the overall mean, e_i is the sampling error for the *i*th study, **e** is a 1 × $N_{studies}$ column vector with the e_i values (which are assumed to be normally distributed with mean 0 and variance v_i), **0** is a column vector of zeros, and **V** is an $N_{studies} \times N_{studies}$ matrix with ¹⁵¹ the v_i values along the diagonal.

The fixed-effects model assumes that the included studies share a single common true effect. This assumption, however, is rarely met in multi-population and multi-species metaanalyses of ecology and evolution studies (Gurevitch and Hedges, 1999; Higgins et al., 2009). The random-effects model addresses this potential 'heterogeneity' among the true effects by adding a random effect corresponding to each estimate and is given by

$$y_i = \mu + u_i + e_i \tag{3}$$

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$$\mathbf{u} \sim N(\mathbf{0}, \sigma_u^2 \mathbf{I}_u),\tag{4}$$

where u_i is the random effect corresponding to the *i*th estimate, **u** is a 1 × $N_{studies}$ column vector with the u_i values (which are assumed to be normally distributed with mean 0 and variance σ_u^2), and \mathbf{I}_u is an $N_{studies} \times N_{studies}$ identity matrix.

Although the models above are suitable for conducting a meta-analysis in many circumstances, they do not account for the multilevel structure that arises when at least some studies provide multiple effect size estimates (e.g., when the same experiment was conducted under varying circumstances within the same study) and they do not account for phylogenetic dependence (when studies are conducted with multiple species that differ in similarity due to differences in their shared evolutionary history).

To address the first issue, we can use a multilevel meta-analytic model (Konstantopoulos, 2011; Nakagawa and Santos, 2012) which includes a random effect at the effect size level (as in model 3), but which now captures variability in the true effects within studies, and a random effect at the study level, which captures between-study variability. Let y_{ij} denote the *j*th effect in the *i*th study (with $j = 1, ..., N_i$, where N_i is the number of effect sizes reported in the *i*th study), v_{ij} the corresponding sampling variance, and let $N_{total} = \sum_{i=1}^{N_{studies}} N_i$ denote ¹⁷³ the total number of effects. The model is then given by

$$y_{ij} = \mu + u_{ij} + s_i + e_{ij} \tag{5}$$

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$$\mathbf{s} \sim N(\mathbf{0}, \sigma_s^2 \mathbf{I}_s),\tag{6}$$

where u_{ij} is a random effect corresponding to the *j*th effect size in the *i*th study, s_i is a 175 random effect at the study level, **u** is now a $1 \times N_{total}$ column vector with the u_{ij} values, **s** is 176 a $1 \times N_{studies}$ column vector with the s_i values (which are assumed to be normally distributed 177 with mean 0 and variance σ_s^2), and \mathbf{I}_u and \mathbf{I}_s are $N_{total} \times N_{total}$ and $N_{studies} \times N_{studies}$ identity 178 matrices, respectively. Finally, **e** is now a $1 \times N_{total}$ column vector with the e_{ij} values and **V** 179 is the corresponding (diagonal) variance-covariance matrix with dimensions $N_{total} \times N_{total}$. 180 When the effect size estimates were computed based on a set of $N_{species}$ different species, 181 we will need an additional index. Let y_{ijk} denote the *j*th effect in the *i*th study as before, but 182 now let $k = 1, \ldots, N_{species}$ be the index that indicates for which species a particular effect 183

size estimate was computed. Model 5 can then be extended to account for species-level
variability as follows:

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$$y_{ijk} = \mu + u_{ij} + s_i + n_k + e_{ij}, \tag{7}$$

$$\mathbf{n} \sim N(\mathbf{0}, \sigma_n^2 \mathbf{I}_n),\tag{8}$$

where n_k is a species-specific random effect, **n** is a $1 \times N_{species}$ column vector with the n_k values (which are assumed to be normally distributed with mean 0 and between-species variance σ_n^2), and \mathbf{I}_n has dimensions $N_{species} \times N_{species}$. Note that n_k is a crossed random effect (e.g., Fernández-Castilla et al., 2019) and not nested within studies and we therefore do not put subscript k on u_{ij} , s_i , or e_{ij} .

¹⁹² Model 7, however, does not account for phylogeny. For this, we further extend the ¹⁹³ model by including an additional species-level random effect (Hadfield and Nakagawa, 2010), ¹⁹⁴ but instead of assuming independence for different species (as for the n_k values), we allow the values of the random effect to be correlated according to a phylogenetic correlation matrix, which in turn is derived from a phylogenetic tree based on some model of evolution (e.g., Brownian motion) prior to the analysis (e.g., Lajeunesse, 2009; Felsenstein, 1985, 2004; Freckleton et al., 2002). The model is given by

$$y_{ijk} = \mu + u_{ij} + s_i + n_k + p_k + e_{ij},\tag{9}$$

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$$\mathbf{p} \sim N(\mathbf{0}, \sigma_p^2 \mathbf{A}),\tag{10}$$

where p_k denotes the phylogenetic random effect for the kth species, **p** is a $1 \times N_{species}$ column 200 vector with the p_k values (which are assumed to follow a multivariate normal distribution with 201 mean 0 and variance-covariance matrix $\sigma_p^2 \mathbf{A}$, where \mathbf{A} is the $N_{species} \times N_{species}$ phylogenetic 202 correlation matrix). Hence, the model includes a non-phylogenetic species-level random effect 203 (i.e., the n_k values) to account for heterogeneity in the effects sizes due to differences between 204 species unrelated to phylogeny (e.g., the influence of differences in the environments they 205 live in) and a phylogenetic random effect (i.e., the p_k values) that captures dependencies in 206 the effect sizes according to the similarities between species due to phylogenetic relatedness. 207 A concern with model 9 arises when phylogenetic relationships are weak. In that case, 208 A starts to resemble I_n and hence σ_p^2 and σ_n^2 are confounded and may not be uniquely 209 identifiable. This may lead to bias in the estimates of the variance components. This concern, 210 or the complexity of model 9 in general, has led some researchers to adopt a simplified model 211 in their meta-analyses where the non-phylogenetic variance component is removed. This 212 leads to the model 213

$$y_{ijk} = \mu + u_{ij} + s_i + p_k + e_{ij}, \tag{11}$$

with all terms as explained before. Whether this simplified version is an adequate substitute
for model 9 is currently unknown.

The models described above can be fitted within a Bayesian or likelihood framework (Hadfield and Nakagawa, 2010). For the latter, the metafor package (Viechtbauer, 2010)

for R (R Core Team, 2021) is particularly attractive as it is freely available and was written 218 specifically for the purposes of conducting meta-analyses. Maximum likelihood (ML) or 219 restricted maximum likelihood (REML) estimation can be used for model fitting (the latter 220 usually being the preferred choice; see Patterson and Thompson, 1971), providing estimates 221 of the variance components included in a particular model, the estimate of μ (i.e., $\hat{\mu}$), and its 222 standard error (i.e., $SE[\hat{\mu}]$). Likelihood ratio tests and profile likelihood confidence intervals 223 provide inferences for the variance components. An approximate 95% Wald-type confidence 224 interval for μ can be obtained with $\hat{\mu} \pm 1.96 \text{SE}[\hat{\mu}]$. Analogously, $H_0: \mu = 0$ can be tested 225 by comparing $z = \hat{\mu}/\text{SE}[\hat{\mu}]$ against the critical values (i.e., ±1.96) of a standard normal 226 distribution. 227

Although fitting the models and deriving inference from them is feasible, the consequences of using the various models have not been examined systematically. We therefore conducted an extensive simulation study to investigate the performance of the various model under varying circumstances.

232 2.2 Simulation Setup

In our setup, the primary studies could provide one or multiple effect size estimates for one 233 or multiple species. We set $(N_{studies}, N_{species})$ either to (20, 40) or (50, 100) to examine the 234 difference between a smaller versus larger meta-analysis. Furthermore, we set σ_u^2 , σ_s^2 , σ_n^2 , and 235 σ_p^2 to either 0, 0.05, or 0.3 (plus an additional parameter α to be described below to either 236 0.5, 1, or 2) to define a particular condition within the simulation study. Table 1 provides 237 an overview of the 158 conditions that were studied in this manner. Note that we used 238 a 'conditional factorization' of the variance components to keep the number of conditions 239 manageable and to generate scenarios where one of the models described in equations 3, 5, 7, 240 and 9 corresponds to the true data generating mechanism (see Table 1). Within a particular 241 condition, the following steps were repeated 1000 times. 242

First, the number of effect sizes provided by the studies (i.e., the N_i values) were simulated

$N_{studies}$	$N_{species}$	σ_u^2	σ_s^2	σ_n^2	σ_p^2	α	Conditions	True model
20	40	0, 0.05, 0.30	0	0	0	1	3	Model 3
20	40	0.05, 0.30	0.05, 0.30	0	0	1	4	Model 5
20	40	0.05, 0.30	0.05, 0.30	0.05, 0.30	0	0.5, 1, 2	24	Model 7
20	40	0.05, 0.30	0.05, 0.30	0.05, 0.30	0.05, 0.30	0.5,1,2	48	Model 9
50	100	0, 0.05, 0.30	0	0	0	1	3	Model 3
50	100	0.05, 0.30	0.05, 0.30	0	0	1	4	Model 5
50	100	0.05, 0.30	0.05,0.30	0.05,0.30	0	0.5, 1, 2	24	Model 7
50	100	0.05, 0.30	0.05, 0.30	0.05, 0.30	0.05, 0.30	0.5,1,2	48	Model 9

Table 1: Overview of the conditions examined in the simulation study. The first two columns show the number of studies and species, respectively. The next four columns indicate the true values of the variance components. The α column represent the power parameter. All values were crossed within a particular row of the table. The last two columns respectively represent the number of conditions generated in each row and the model that corresponds to the true data generating mechanism for the conditions in a particular row.

from a right-skewed distribution, as typically observed in practice. For this, we generated $N_{studies}$ random values from a Beta(1.5, 3) distribution, which were then multiplied by 39, rounded to the closest integer, and increased by 1. Therefore, the number of estimates per study could vary between 1 and 40 (with a mean, median, and mode of approximately 14, 13, and 9, respectively).

In the next step, we simulated the species indices (i.e., the k values) by generating N_{total} random values from a Beta(2, 2) distribution, which were multiplied by $N_{species} - 1$, rounded to the closest integer, and then increased by 1. Accordingly, the number of times that the various species were studied followed a symmetric unimodal distribution (with mean equal to ($N_{species} + 1$)/2). In order to guarantee that all species appear at least once in each metaanalysis, a randomly chosen $N_{species}$ random numbers generated this way were replaced with the integers from 1 to $N_{species}$.

Next, we generated a phylogenetic tree for the species using the rtree() function from the R package ape (Paradis and Schliep, 2019), which uses a recursive random splitting algorithm to simulate a phylogeny (Paradis, 2012). The branch lengths were then computed



Figure 1: An example of a simulated phylogenetic tree for 40 species modified with different values of the power parameter α (i.e., 0.5, 1, and 2).

using the compute.brlen() function based on the method by Grafen (1989), using the power 250 parameter α to adjust the 'height' of branch lengths at the tips of the phylogenetic tree, 260 leading to phylogenetic relationships that are generally stronger when branches are shorter 261 at the tips or weaker when branches are longer at the tips. Fig. 1 shows an example of such 262 a simulated tree for 40 species modified by different α values. Finally, the correlation matrix 263 that represents the phylogenetic relationships (matrix \mathbf{A} in equation 10) was calculated 264 from the tree by using the vcv() function based on a Brownian model of evolution (i.e., 265 $\mathbf{A}_{k,k'} = 1 - b_{k,k'}$, where $b_{k,k'}$ is the branch length for a pair of species to their most recent 266 common ancestor). 267

We then generated the values for the four random effects, corresponding to the variance components σ_u^2 , σ_s^2 , σ_n^2 , and σ_p^2 , either as independent draws from normal distributions for the first three components or from a multivariate normal distribution for the last one. In conditions where a variance component is equal to 0, the corresponding random effect values are then just a series of 0s of the appropriate length. To complete the data generating step, the sampling variances (i.e., the v_{ij} values) were simulated from a right-skewed Beta(2, 20) distribution (and hence had a value of .091 on average) which were then used to generate the N_{total} sampling errors from a normal distribution with mean 0 and variance v_{ij} . We then summed up the random effects and sampling errors as shown in equation 9, setting $\mu = 0$ without loss of generality.

After generating the data, we fitted the four models shown in equations 3, 5, 7, and 9, 278 using REML estimation as implemented in the rma.mv() function from the metafor package. 279 For model 3, we simply treated each estimate as a separate study (one can also think of this 280 as model 5 without the addition of the study-level random effect). For each model, we then 281 saved the estimate of μ , the variance component estimates, the bounds of the 95% Wald-type 282 confidence interval for μ , and the model fitting time. In case any one of the four models 283 did not converge within a particular iteration (with the default settings of the rma.mv() 284 function), the iteration was discarded and a new iteration was run to guarantee that a 1000 285 successful model fits were available for all four models. 286

After the 1000 iterations, we computed the mean of the $\hat{\mu}$ values for each model, the mean of the variance component estimates, the proportion of iterations where 0 was included in the confidence interval (i.e., the empirical coverage rate for μ), the mean confidence interval width, the convergence rate, and the mean model fitting time. The simulation was run on a workstation with an Intel Xeon E5-2630v4 processor utilizing 15 cores in parallel. Completion time for the simulation was approximately 7 days (roughly 2520 core hours).

We generated two other sets of conditions to investigate specific questions. First, we 293 examined conditions where the phylogenetic relationships could also be weaker than in the 294 main scenarios to test the performance of model 9 under such conditions. These conditions 295 were generated by setting α to (0.1, 0.2, 0.3, 0.4, 0.5, 1, 2) when $(N_{studies}, N_{species}) = (50, 100)$, 296 the estimate- and study-level variance components were both large (0.3), and the levels of 297 the remaining variance components were factorized with values of 0.05 and 0.3 (for a total of 298 28 different conditions). Second, we compared the performance of model 9 and the simplified 299 model 11 (that leaves out the non-phylogenetic species-level random effect). For this, we set 300

 $(N_{studies}, N_{species}) = (50, 100), \sigma_u^2 = 0.05, \sigma_s^2 = 0.05, \text{ and } \alpha = 1, \text{ and then generated different}$ conditions by factorizing different values of only σ_n^2 and σ_p^2 , where the former was set to values from 0 to 0.3 with increments of 0.05, whereas the latter was set to either 0, 0.05, or 0.3 (for a total of 21 different conditions).

305 **3** Results

306 3.1 Simulation Results

Fig. 2a displays boxplots of the mean $\hat{\mu}$ values for each of the four models across the 158 conditions, separated by which model was the true data generating mechanism. Generally, the means were clustered tightly around 0, indicating little to no bias in $\hat{\mu}$, although in a small set of conditions there was some slight positive bias in the estimates of the overall mean. These conditions were characterized by non-zero values for all four variance components (i.e., when model 9 was the true model), $(N_{studies}, N_{species}) = (20, 40)$, a weak phylogenetic relationship ($\alpha = 0.5$), and a large phylogenetic variance ($\sigma_p^2 = 0.3$).

In contrast to the results for the overall mean, the coverage rates of the 95% confidence 314 interval for μ differed markedly across models (Fig. 2b). For conditions where model 3 was 315 the true data generating mechanism, all models achieved coverage rates close to or slightly 316 above the nominal 95% confidence level regardless of the condition. As the other variance 317 components were introduced into the data, however, the coverage rates of models that did 318 not account for these additional sources of variability started to decrease, at times severely 319 so. Only model 9 was able to achieve rates close to the nominal level across the majority 320 of conditions, although the rates also fell somewhat below the nominal level for certain 321 conditions when all variance components were larger than zero. 322

Given that estimates of μ were relatively unbiased for all models, the closer to nominal coverage rates of model 9 would be expected to be mainly a consequence of wider confidence intervals (that consequently have a better chance of capturing the true value of μ). Fig. 2c



Figure 2: Boxplots based on the (a) mean $\hat{\mu}$ values, (b) coverage rates of the 95% confidence interval for μ , and (c) mean confidence interval widths for each of the four models across the 158 conditions, separated by which model was the true data generating mechanism.

confirms this, showing the mean confidence interval widths for the various models across the various conditions. However, what is particularly noteworthy is that the use of model 9 under conditions where actually a simpler model is the true data generating mechanism only leads to a relatively minor increase in the mean interval width.

Fig. 3 displays the bias in the variance component estimates of model 9 under the 28 different conditions generated by varying α , σ_n^2 , and σ_p^2 (while holding σ_u^2 and σ_s^2 constant at 0.3). The results show no bias in the estimates of σ_u^2 and σ_s^2 . Furthermore, the model



Figure 3: Mean bias of the variance component estimates of model 9 under different combinations of the power parameter (α) and the non-phylogenetic and phylogenetic variance components (σ_n^2 and σ_p^2 , respectively).

is able to estimate σ_n^2 and σ_p^2 with little to no bias, except when the strength of the phylogenetic relationships decreased. As expected, under such conditions, the model struggles to provide unbiased estimates of the non-phylogenetic and phylogenetic species-level variance components (especially when $\sigma_n^2 \neq \sigma_p^2$).

Fig. 4a shows the coverage rates of the confidence interval for μ for models 9 and 11 as the size of the non-phylogenetic species-level variance component (i.e., σ_n^2) was increased. While model 9 provided rates close to or somewhat below the nominal level, the rates for model 11 were often equal to 100% and hence the confidence interval tended to be too wide. Furthermore, Fig. 4b demonstrates that the bias in the phylogenetic variance component of model 11 inflated rapidly as the value of σ_n^2 increased (the value of σ_p^2 had no noteworthy



Figure 4: Comparison of models 9 and 11 as the size of the non-phylogenetic species-level variance component (i.e., σ_n^2) was systematically increased. (a) Coverage rates of the 95% confidence intervals for μ , (b) bias in the non-phylogenetic and phylogenetic variance components.

influence on the bias and hence we averaged these results over the three possible values of σ_p^2 . In contrast, model 9 estimated these two variance components essentially without bias under these scenarios.

Model fitting times differed between the various models (Table 2), with model 9 requiring the most amount of time on average, regardless of the true data generating mechanism. The most challenging conditions for the more complex models were those scenarios where model 3 corresponded to the true data generating mechanism. In this case, a single fit of model 9 took around 26 seconds on average when $(N_{studies}, N_{species}) = (50, 100)$. In these conditions, convergence rates were also the lowest, although even model 9 then converged in approximately 96% of the iterations.

353 3.2 Illustrative Example

We use the data from the meta-analysis by Rios Moura et al. (2021) on size-assortative mating (SAM) to illustrate an application of the models. Each study included in the meta-analysis

	(a)	$(N_{studies}, N_S)$	$p_{pecies}) = (20,$	40)		(b) $(N_{studies}, N_{Species}) = (50, 100)$					
	True Model					True Model					
Model Fit	Model 3	Model 5	Model 7	Model 9	Model Fit	Model 3	Model 5	Model 7	Model 9		
Model 3	0.939	0.668	0.700	0.700	Model 3	1.589 1.313 1.307	1.294				
Model 3	(100.00%)	(100.00%)	(100.00%)	(100.00%)	Model 3	(100.00%)	(100.00%)	(100.00%)	(100.00%)		
Model 5	2.653	1.104	1.151	1.162	Madal 5	3.934	1.986	1.999	1.959		
Model 9	(99.81%)	(100.00%)	(100.00%)	(100.00%)	Model 5	(99.78%)	(100.00%)	(100.00%)	(100.00%)		
Model 7	2.484	1.876	0.868	0.858	Model 7	19.823	14.752	7.364	7.393		
Model /	(97.53%)	(100.00%)	(100.00%)	(100.00%)	Model /	(96.86%)	(100.00%)	(100.00%)	(100.00%)		
Model 0	3.316	3.053	2.288	1.463	163 25.980 23.540	18.641	11.005				
Model 9	(96.56%)	(99.73%)	(99.99%)	(99.99%)	woder 9	(95.63%)	(99.60%)	(100.00%)	(100.00%)		

Table 2: Average model fitting times in seconds and convergence rates (in parentheses) of all models under the different data generating mechanisms.

provided one or multiple correlation coefficients describing the similarity in some measure of 356 body size in mating couples. For the analysis, the correlation coefficients were transformed 357 with Fisher's r-to-z transformation. We focus here on the estimate of the overall mean 358 (transformed) correlation coefficient, leaving aside the issue of differences between studies 359 where correlations were computed with or without pooling of data across different timepoints 360 or areas (i.e., temporal/spatial pooling). Also, using the method by Grafen (1989), we turned 361 the phylogenetic tree used by Rios Moura et al. (2021) into an ultrametric tree before fitting 362 models 9 and 11, to bring these analyses more in line with how our simulation study was 363 conducted. The dataset includes 1828 effect size estimates (i.e., transformed correlations) 364 collected from 457 studies and 341 species. 365

Table 3 presents the results obtained from each model. Interestingly, the estimate of the overall mean tended to be somewhat larger in the more complex models, although differences between models 7, 9, and 11 were relatively small. More importantly, we see a substantial increase in the standard error of the estimated overall mean for the more complex models. As a result, the confidence intervals become wider, the values of the test statistics smaller, while the respective *p*-values increase. Although each model suggests that the overall mean significantly differs from 0 (at the conventional 0.05 level of significance), the *p*-value for

	$\hat{\mu}$	$SE[\hat{\mu}]$	95% CI	Z	p	$\hat{\sigma}_u^2$	$\hat{\sigma}_s^2$	$\hat{\sigma}_n^2$	$\hat{\sigma}_p^2$	AIC
Model 3	0.24	0.007	0.23, 0.25	34.15	< 0.0001	0.0641	_	_	_	1082.8
Model 5	0.30	0.015	0.27, 0.33	20.42	< 0.0001	0.0149	0.0806	—	—	429.0
Model 7	0.34	0.020	0.30, 0.38	17.37	< 0.0001	0.0143	0.0195	0.0815	—	386.3
Model 9	0.37	0.130	0.11, 0.62	2.83	0.0046	0.0145	0.0192	0.0555	0.0512	344.7
Model 11	0.36	0.172	0.02, 0.70	2.07	0.0383	0.0149	0.0557	_	0.0914	367.2

Table 3: Results derived from fitting the various models to the example dataset. The first five columns show the estimated overall mean, its standard error, the 95% confidence interval, the test statistic, and the *p*-value for testing H_0 : $\mu = 0$, respectively. The next four columns show the estimates of the variance components in the respective models. The last column shows the Akaike Information Criteria (AIC) values.

³⁷³ model 11 was approaching the rejection threshold.

The estimates of the variance components also show some interesting patterns. While the 374 simple random-effects model 3 cannot distinguish between different sources of variability and 375 attributes all of the heterogeneity to differences between the individual effect size estimates, 376 model 5 suggests that the variance in the effects is more related to differences between 377 studies than particular estimates within studies. However, once species-level variability is 378 considered, it becomes apparent that this is actually the dominant source of heterogeneity. 379 Moreover, model 9 shows that this variability is to approximately equal parts attributable to 380 non-phylogenetic and phylogenetic species-level differences. In contrast, when ignoring the 381 non-phylogenetic variance component in the simplified model 11, part of the variance from 382 that component is forced back into the study-level variance component. Furthermore, $\hat{\sigma}_p^2$ in 383 the simplified model is substantially inflated compared to model 9 which may be an example 384 of the inflation in this component when σ_n^2 is excluded (see Fig. 4b). Based on these findings 385 and the Akaike Information Criteria (AIC) values of the various models, we would strongly 386 favor model 9 in this comparison, illustrating that both non-phylogenetic and phylogenetic 387 variance components should be considered in the analysis. 388

389 4 Discussion

Meta-analyses in the fields of ecology and evolution typically need to address the fact that 390 multiple effect size estimates can be extracted from at least some of the studies and that 391 the estimates are based on various species that are related to each other due to their shared 392 evolutionary history. In this paper, we investigated the performance of the phylogenetic 393 multilevel meta-analytic model by Hadfield and Nakagawa (2010) and Nakagawa and Santos 394 (2012) that captures these intricacies along with some simpler models. Despite the concerns 395 raised in the introduction, the model can successfully estimate the overall mean and its 396 uncertainty. It also provides approximately unbiased estimates of all variance components, 397 including the non-phylogenetic and phylogenetic species-level variances, as long as there are 398 at least moderately strong phylogenetic relationships among the species. In addition, despite 399 its complexity, the model does not appear to suffer from convergence problems and model 400 fitting does not require excessive computational times. 401

402 4.1 Estimating the Overall Mean and its Uncertainty

Not only the phylogenetic multilevel meta-analytic model, but also the simpler models that 403 leave out certain variance components provide essentially unbiased estimates of the overall 404 mean, regardless of the nature of the true model that underlies the data (Fig. 2a). However, 405 the uncertainty in the overall mean will only be estimated accurately when the fitted model 406 includes the variance components that contribute to the heterogeneity and the dependencies 407 among the underlying true effects. Fitting underspecified models typically led to severe 408 undercoverage of the confidence interval for the overall mean and hence anticonservative 409 inferences. In fact, subtracting the coverage rates shown in Fig. 2b from 1 yields the Type 410 I error rates for the test of the overall mean, which could go as high as 91% when using 411 a simple random-effects model that ignores the multilevel structure and the species-level 412 variance components. 413

These findings are in line with those by Chamberlain et al. (2012), who demonstrated, based on 30 published meta-analyses, that the inclusion of phylogeny into a random-effects model usually only led to minor changes in the pooled effect size, but had a more substantial impact on the statistical significance of the finding (turning significant findings into nonsignificant ones in the majority of cases where changes occurred).

Our findings can also be used to alleviate concerns with using the phylogenetic multilevel 419 meta-analytic model when it is actually an overspecified model (i.e., when the actual data 420 generating mechanism is simpler). In those cases, the mean confidence interval width of the 421 model was just barely wider than that of the simpler models, indicating little to no loss in 422 efficiency by fitting an overly complex model (Fig. 2c). The superfluous variance components 423 then converge towards 0 (or close to it), which appears to be slightly more challenging for 424 the optimization algorithm, leading to longer model fitting times and occasional convergence 425 problems, but not to any worrisome degree (Table 2). Moreover, in practice, for any particu-426 lar dataset, convergence problems can typically be resolved by selecting a different optimizer 427 or making changes to the settings for the optimization routine, so the convergence rates as 428 given only apply to the default settings. 429

At the same time, we should point out that the coverage rate of the model did fall slightly 430 below the nominal 95% level in the majority of conditions when all variance components 431 were in fact non-zero (see Fig. 2b, rightmost panel). This undercoverage stems from using 432 an overly simple Wald-type confidence interval using critical values based on a standard 433 normal distribution that ignores the uncertainty in the estimates of the variance components 434 (especially in the study and the two species-level components when $N_{studies}$ and $N_{species}$ are 435 low). A similar issue, but for a simpler model with only between- and within-study variance 436 components (i.e., model 5 in our simulation) was also recently pointed out by Song et al. 437 (2020). Improved methods based on the t-distribution, with various approximations for 438 the degrees of freedom, have been proposed and studied extensively in the context of the 439 standard random-effects model (e.g., Sanchez-Meca and Marin-Martinez, 2008) and mixed-440

effects models in general (e.g., Luke, 2017), but these methods have not been generalized to the present context. As a simple approximation, using the smaller of $N_{studies} - 1$ and $N_{species} - 1$ as the degrees of freedom for a confidence interval based on a t-distribution is likely to bring the coverage rate quite close to the nominal rates in the majority of conditions.

445 4.2 Including and Testing the Phylogenetic Effect

Phylogenies play a central role in the context of phylogenetic comparative studies (Freckleton 446 et al., 2002; Blomberg et al., 2003; Ives et al., 2007). An important step in such studies is 447 testing the significance of the 'phylogenetic signal' in some trait of interest. This test is 448 often performed through a statistic such as λ (Pagel, 1999) or K (Blomberg et al., 2003). 449 Although model 9 does not parameterize the phylogenetic effect in this manner, one can 450 derive information from its output that shows its relationship to the λ statistic. In particular, 451 Pagel's λ is a multiplicative factor that is applied to the off-diagonal values of the correlation 452 matrix that represents the phylogenetic relationships (i.e., the A matrix). For example, the 453 variance-covariance matrix for three species would be given by 454

$$\sigma^2 \begin{bmatrix} 1 & \lambda a_{12} & \lambda a_{13} \\ & 1 & \lambda a_{23} \\ & & 1 \end{bmatrix}$$

while the decomposition of the species-level heterogeneity in model 9 implies the variance covariance matrix

$$\sigma_n^2 \begin{bmatrix} 1 & & \\ & 1 & \\ & & 1 \end{bmatrix} + \sigma_p^2 \begin{bmatrix} 1 & a_{12} & a_{13} \\ & 1 & a_{23} \\ & & & 1 \end{bmatrix} = (\sigma_n^2 + \sigma_p^2) \begin{bmatrix} 1 & \left(\frac{\sigma_p^2}{\sigma_n^2 + \sigma_p^2}\right) a_{12} & \left(\frac{\sigma_p^2}{\sigma_n^2 + \sigma_p^2}\right) a_{13} \\ & 1 & \left(\frac{\sigma_p^2}{\sigma_n^2 + \sigma_p^2}\right) a_{23} \\ & & & 1 \end{bmatrix}$$

and hence $\sigma^2 = \sigma_n^2 + \sigma_p^2$ and $\lambda = \sigma_p^2/(\sigma_n^2 + \sigma_p^2)$ (see also Lynch, 1991; Freckleton et al., 2002). Hence, $\sigma_p^2/(\sigma_n^2 + \sigma_p^2)$ indicates the degree of the phylogenetic signal in the overall variance sourced from the species. A likelihood ratio test of $H_0: \sigma_p^2 = 0$ can be easily performed by comparing $X^2 = -2(ll_7 - ll_9)$ against a chi-squared distribution with one degree of freedom, where ll_7 and ll_9 are the (restricted) log likelihoods of models 7 and 9, respectively.

462 4.3 Estimating the Non-Phylogenetic and Phylogenetic Variance

Given the informative nature of these two variance components, it is essential to estimate 463 their true values accurately. We found that model 9 was usually able to estimate these compo-464 nents unbiasedly, but should note that the model struggles to separate the non-phylogenetic 465 and phylogenetic species effects when phylogenetic relationships are weak. In essence, the 466 two sources of variability then start to collapse into one, with a total variance of $\sigma_n^2 + \sigma_p^2$. This 467 total variance is then distributed in approximately equal parts into the two estimates, which 468 explains the apparent low bias when (coincidentally) $\sigma_n^2 = \sigma_p^2$ (Fig. 3a and d). However, 469 when $\sigma_n^2 \neq \sigma_p^2$, the bias in the two estimates becomes apparent (Fig. 3b and c). Therefore, 470 we would caution against the use of model 9 when phylogenetic relationships are weak. As a 471 rough guideline, for $\alpha = 0.5$, the mean correlation in the A matrix (excluding the diagonal) 472 is around 0.2 and hence a lower mean correlation would call into question the trustworthiness 473 of the estimates of σ_n^2 and σ_p^2 . 474

Some meta-analyses in ecology and evolution have used model 11 to reduce model com-475 plexity (e.g., Garamszegi et al., 2012; Moore et al., 2016). Our results indicate that this 476 approach cannot be recommended. As we increased the value of σ_n^2 , the bias in the phyloge-477 netic variance component inflated massively in this simplified model (Fig. 4b). As a result, 478 the relevance of the phylogeny could be greatly overestimated. In addition, the confidence 479 interval for the overall mean then becomes extremely conservative with coverage rates at 480 or very close to 100%. This in turn implies a loss of efficiency for estimating the overall 481 mean and a loss of power for testing $H_0: \mu = 0$. The illustrative example also shows this 482

483 phenomenon.

484 4.4 Caveats and Conclusions

For the simulation study, we used a 'generic' effect size measure, that is, we directly simulated 485 the sampling errors from a normal distribution and treated the sampling variances (i.e., 486 the v_{ij} values) as known. These conditions only apply asymptotically to measures typically 487 used in practice (e.g., standardized mean differences, response ratios, correlation coefficients, 488 risk/odds ratios). The present results therefore reflect the performance of the various models 489 under idealized conditions (i.e., when the sample sizes of the individual studies are sufficiently 490 large, such that the sampling distributions of the estimates are indeed approximately normal 491 and any inaccuracies in the estimated sampling variances are negligible). The advantage of 492 using a generic measure is that we were able to identify problems that are inherent to certain 493 models and not (potentially) a consequence of violations to the model assumptions. On the 494 other hand, it remains to be determined how well the phylogenetic multilevel model performs 495 when the effect sizes are generated based on the exact distributional assumptions underlying 496 specific measures. 497

Therefore, at least for the moment, the present results suggest that model 9 is the most appropriate tool for conducting a multi-species meta-analysis in ecology and evolution. For the vast majority of conditions examined, it provides approximately unbiased estimates of the variance components and the overall mean and a confidence interval for the latter with a close to nominal coverage rate. Therefore, we recommend that meta-analysts in ecology and evolution use the phylogenetic multilevel model as the de facto standard when analyzing multi-species datasets.

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506 Conflict of interest statement: The authors declare that they have no competing 507 interests.

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Author contributions: SN provided contextual and literature review support, WV provided the code to run the simulation, all authors contributed to the manuscript.

⁵¹² **Data accessibility statement:** No new data were used in this study. The material to ⁵¹³ reproduce the results are available at: https://osf.io/ms8eq/.

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