- A Standardized Effect Size for Evaluating the Strength of Phylo-
- 2 genetic Signal, and Why Lambda is not Appropriate

5 Abstract

### 6 Introduction

Investigating macroevolutionary patterns of trait variation requires a phylogenetic perspective, because the shared ancestry among species generates statistical non-independence (Felsenstein 1985; Harvey and Pagel 1991). Accounting for this evolutionary non-independence is the purview of phylogenetic comparative methods (PCMs); a suite of analytical tools that condition the data on the phylogeny through the course 10 of statistical evaluations of phenotypic trends (e.g., Grafen 1989; Garland and Ives 2000; Rohlf 2001; 11 Butler and King 2004). The past several decades have witnessed a rapid expansion in the development 12 of PCMs to address an ever-growing set of macroevolutionary hypotheses (Martins and Hansen 1997; 13 O'Meara et al. 2006; Revell and Harmon 2008; Beaulieu et al. 2012; Adams 2014b,a; Adams and Collyer 2018). These methods are predicated on the notion that phylogenetic signal – the tendancy for 15 closely related species to display similar trait values – is present in cross-species datasets (Felsenstein 1985; Pagel 1999; Blomberg et al. 2003). Indeed, under numerous evolutionary models, phylogenetic 17 signal is to be expected, as stochastic character change along the hierarchical structure of the tree of life generates trait covaration among related taxa (see Felsenstein 1985; Blomberg et al. 2003; Revell et al. 2008). 19

Several analytical tools have been developed to quantify phylogenetic signal in phenotypic datasets, including measures of serial independence (C: Abouheif 1999), autocorrelation estimates (I: Gittleman and Kot 1990), statistical ratios of trait variation relative to what is expected given the phylogeny (Kappa: Blomberg et al. 2003; Adams 2014a), and scaling parameters used in maximum likelihood fitting of the data to the phylogeny ( $\lambda$ : Pagel 1999), among others (e.g., Klingenberg and Gidaszewski 2010). The statistical properties of these methods – namely type I error rates and power – have also been investigated to determine when phylogenetic signal can be detected and under what conditions (e.g., Munkemuller et al. 2012; Pavoine and Ricotta 2012; Diniz-Filho et al. 2012; Adams 2014a; Molina-Venegas and Rodriguez 2017; see also Revell et al. 2008; Revell 2010). One of the most widely used methods for characterizing phylogenetic signal in macroevolutionary studies is Pagel's  $\lambda$  (Pagel 1999). Here, maximum likelihood is used to fit the data to the phylogeny under a Brownian motion model of evolution. A parameter ( $\lambda$ ) is included, which transforms the lengths of the internal branches of the phylogeny to improve the fit (Pagel 1999; Freckleton et al. 2002). Pagel's  $\lambda$  ranges from  $0 \to 1$ , with larger values signifying a greater dependence of observed trait variation on the phylogeny. Pagel's  $\lambda$  also has the appeal that it may be included in phylogenetic regression (PGLS) to account for the degree of phylogenetic signal in comparative analyses (see Freckleton et al. 2002).

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Evolutionary biologists commonly seek to describe the relative strength of phylogenetic signal in phenotypic traits, to determine the extent to which shared evolutionary history has influenced trait covariation among taxa. This is often accomplished by interpreting empirical estimates of  $\lambda$ ; with smaller values signifying weak' phylogenetic signal, while larger values are interpreted as 'strong' phylogenetic signal (e.g., De Meester et al. 2019; Pintanel et al. 2019; Su et al. 2019). Other approaches for interpreting  $\lambda$  are more statistical. 41 For instance, some have evaluated whether the observed  $\lambda$  differs from some expected value through the use of confidence intervals (Vandelook et al. 2019) or by performing likelihood ratio tests that compare the observed model fit to that obtained when  $\lambda = 0$  or  $\lambda = 1$  (Freckleton et al. 2002; Cooper et al. 2010; Bose et al. 2019). Additionally, qualitative comparisons of  $\lambda$  estimates obtained from multiple phenotypic traits have been used to infer whether the strength of phylogenetic signal is greater in one trait as compared to another (e.g., Liu et al. 2019; Bai et al. 2019). Indeed, statements regarding the strength of phylogenetic signal based on  $\lambda$  are rather common in the evolutionary literature. For instance, of the 204 papers published in 2019 that estimated and reported Pagel's  $\lambda$  (found in Google.scholar), 40% interpreted the strength of phylogenetic signal for at least one phenotypic trait. Additionally, because nearly half of the 1,572  $\lambda$  values reported were near the limits of the parameter (Figure 1), this percentage is even higher, as biological interpretation of 51 phylogenetic signal at the limits of  $\lambda$  are known.

54 [insert Figure 1 here]

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It seems intuitive to interpret the strength of phylogenetic signal based on the value of  $\lambda$ , as  $\lambda$  is a parameter on a bounded scale  $(0 \to 1)$  for which interpretation of its extremal points are understood. Specifically,  $\lambda = 0$  represents no phylogenetic signal, while  $\lambda = 1$  is phylogenetic signal as expected under Brownian motion. However, equating values of  $\lambda$  directly to the strength of phylogenetic signal presumes two important statistical properties that have not been fully explored. First, it presumes that values of  $\lambda$  can be precisely estimated, as biological inferences regarding the strength of phylogenetic signal depend on high accuracy in its estimation. Therefore, understanding the precision in estimating  $\lambda$  is paramount. One study (Boettiger et al. 2012) found that estimates of Pagel's  $\lambda$  displayed less variation (i.e., greater precision) when data were simulated on a large phylogeny (N = 281) as compared to a small one (N = 13). From this observation it was concluded that insufficient data (i.e., the number of species) was the underlying cause of the increased variation across parameter estimates (Boettiger et al. 2012). Indeed, such a pattern is common with statistical estimators, as summary statistics and parameters are often more precise at greater sample sizes (Cohen 1988). However, this conclusion also assumes that the precision of  $\lambda$  remains constant across its range

 $(\lambda = 0 \rightarrow 1)$ ; an assumption that to date, has not been verified. Thus, despite widespread use of Pagel's (1999)  $\lambda$  in macroevolutionary studies, at present, we still lack a general understanding of the precision with which  $\lambda$  can estimate levels of phylogenetic signal in phenotypic datasets.

Second, while estimates of  $\lambda$  are within a bounded scale  $(0 \to 1)$ , this does not *de-facto* imply that the estimated values of this parameter correspond to the actual strength of the underlying input signal in the data. For this to be the case,  $\lambda$  must be a statistical effect size. Effect sizes are a measure the magnitude of a statistical effect in data, represented on a common scale (Glass 1976; Cohen 1988). Effect sizes have widespread use in many areas of the quantiative sciences, as they represent measures that may be readily summarized across datasets as in meta-analysis (Glass 1976; Hedges and Olkin 1985; Arnqvist and Wooster 1995), or compared among datasets (e.g., Adams and Collyer 2016, 2019a). Unfortunatley, not all model parameters and test statistics are effect sizes, and thus many summary measures must first be converted to standardized units (i.e., an effect size) for meaningful comparison (see Rosenthal 1994). As a consequence, it follows that only if  $\lambda$  is a statistical effect size can comparisons of estimates across datasets be interpretable. For the case of  $\lambda$ , this has not yet been explored.

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In this study, we evaluate the precision of Pagel's  $\lambda$  for estimating known levels of phylogenetic signal in phenotypic data. We use computer simulations with differing numbers of species, differently shaped phylogenies, and differing input levels of phylogenetic signal, to explore the degree to which  $\lambda$  correctly identifies known levels of phylogenetic signal, and under what circumstances. We find that estimates of  $\lambda$  vary widely for a given input value of phylogenetic signal, and that the precision in estimating  $\lambda$  is not constant across its range. Rather, there is decreased precision when input levels of phylogenetic signal are of intermediate strength. Additionally, the same estimated values of  $\lambda$  may be obtained from datasets containing vastly different input levels of phylogenetic signal. Thus,  $\lambda$  is not a reliable estimate of the strength of phylogenetic signal in phenotypic data. We then describe a standardized effect size for measuring the strength of phylogenetic signal in phenotypic datasets, and apply the concept to two common measures of phylogenetic signal:  $\lambda$  and Kappa. Through simulations across a wide range of conditions, we find that the precision of effect sizes based on  $\lambda$  ( $Z_{\lambda}$ ) are less reliable than that those based on Kappa ( $Z_{K}$ ), implying that  $Z_{K}$  is a more robust effect size measure. We also propose a two-sample test statistic that may be used to compare the strength of phylogenetic signal among datasets, and provide an empirical example to demonstrate its use. We conclude that estimates of phylogenetic signal using Pagel's  $\lambda$  are often inaccurate, and thus interpreting strength of phylogenetic signal in phenotypic datasets based on this measure is compromised. By contrast, effect sizes obtained from *Kappa* hold promise for characterizing phylogenetic signal, and for comparing the strength of phylogenetic signal across datasets.

### Methods and Results

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### $^{104}$ The Precision of $\lambda$ is Variable

We conducted a series of computer simulations to evaluate the precision of Pagel's  $\lambda$ . Our primary simulations 105 were based on pure-birth phylogenies; however, we also evaluated patterns on both balanced and pectinate trees to determine whether tree shape affected our findings (see Supporting Information). First we generated 107 50 pure-birth phylogenies at each of six different tree sizes, ranging from 32 to 1024 taxa  $(n = 2^5 - 2^{10})$ . Next, we rescaled the simulated phylogenies by multiplying the internal branches by  $\lambda_{in}$ , using 21 intervals 109 of 0.05 units across its range ( $\lambda_{in} = 0.0 \rightarrow 1.0$ ), resulting in 1050 scaled phylogenies at each level of 110 species richness (n). Continuous traits were then simulated on each phylogeny under a Brownian motion 111 model of evolution to obtain datasets with differing levels of phylogenetic signal, that ranged from no 112 phylogenetic signal (when  $\lambda_{in} = 0$ ), to phylogenetic signal corresponding reflecting Brownian motion (when 113  $\lambda_{in} = 1$ ). For each dataset we then estimated phylogenetic signal  $(\lambda_{est})$ , and calculated the precision of 114  $\lambda$  using the variance  $(\sigma_{\lambda}^2)$  across datasets at each input level of phylogenetic signal and level of species richness. 115

We also evaluated the precision of  $\lambda$  when estimated in PGLS regression and ANOVA (i.e.,  $Y \sim X$ ). Here, 117 an independent variable X was simulated on each phylogeny under a Brownian motion model of evolution 118 (for PGLS regression). For phylogenetic ANOVA, random groups (X) were obtained by simulating a 119 discrete (binary) character on each phylogeny. Next, the dependent variable was simulated in such a manner as to contain a known relationship with X plus random error containing phylogenetic signal. This was 121 accomplished as:  $Y = \beta X + \epsilon$ . Here, the association between Y and X was modeled using a range of values:  $\beta = (0.0, 0.25, 0.5, 0.75, 1.0)$ , and the residual error was modeled to contain phylogenetic signal simulated 123 under a Brownian motion model of evolution:  $\epsilon = \mathcal{N}(\mu = 0, \sigma = \mathbf{C})$ : (see Revell 2010 for a similar simulation design). The fit of the phylogenetic regression was estimated using maximum likelihood, and parameter 125 estimates ( $\beta_{est}$  and  $\lambda_{est}$ ) were obtained. Precision estimates ( $\sigma_{\lambda}^2$ ) at each input level of phylogenetic signal and level of species richness were then observed. 127

9 All analyses were performed in R v3.6.0 (R Core Team 2019) using the packages geiger (Harmon et al.

2008), caper (Orme et al. 2013), phytools (Revell 2012), and geomorph 3.2.1 (Adams and Otárola-Castillo 2013; Adams et al. 2020). R-scripts are found in the Supporting Information.

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Results. We found that the precision of  $\lambda_{est}$  varied widely across simulation conditions. Predictably, precision improved as the number of species increased (Figure 2). This confirmed earlier findings of 134 Boettiger et al. (2012), and adhered to parametric statistical theory. However, in many cases the set of  $\lambda_{est}$  spanned nearly the entire range of possible values (e.g., n=32;  $\lambda_{in}=0.5$ :  $\lambda_{est}=0.0 \rightarrow 0.985$ ), 136 revealing that estimates of  $\lambda$  were not a reliable indicator of input phylogenetic signal. Importantly, the 137 precision of  $\lambda_{est}$  was not uniform across all levels of phylogenetic signal. The worst precision was observed at 138 intermediate levels of phylogenetic signal ( $\lambda_{in} \approx 0.5$ ), while precision improved as input levels approached 139 the extremes of  $\lambda$ 's range (i.e.,  $\lambda_{in} \to 0 \& \lambda_{in} \to 1$ ). Thus, estimates of  $\lambda$  were least reflective of the true input signal at intermediate values. Additionally, even at large levels of species richness, we found 141 that the range of  $\lambda_{est}$  still encompassed a substantial portion of possible values (e.g.,  $n=512; \lambda_{in}=0.5$ :  $\lambda_{est} = 0.32 \rightarrow 0.68$ ). Likewise, the same  $\lambda_{est}$  could be obtained from datasets containing vastly different input 143 levels of phylogenetic signal (e.g., n = 512;  $\lambda_{est} = 0.5$ ;  $\lambda_{in} = 0.25 \rightarrow 0.65$ ). These findings were particularly unsettling when considered in light of our literature survey. Over one quarter of the  $\lambda$  estimates obtained 145 in empirical studies (421 of 1,572) were between  $\lambda = 0.25$  and  $\lambda = 0.75$  (Figure 1). This range reflected the region that our simulations identified as being the least reliable in terms of accurately characterizing 147 levels of phylogenetic signal, yet 30% of these empirical estimates were explicitly interpreted in terms of 148 the strength of phylogenetic signal that they represented (i.e., weak, intermediate, strong phylogenetic signal).

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Finally, when  $\lambda$  was co-estimated with regression parameters in PGLS regression, the results of our simulations were quite similar. Here, regression parameters ( $\beta$ ) were accurately estimated, confirming earlier findings of Revell (2010) (see Supporting Information). However, estimates of phylogenetic signal ( $\lambda$ ) were less precise (Figure 3), and the spread of  $\lambda_{est}$  was similar to that observed when  $\lambda$  was estimated for only the dependent variable, as in Figure 2. Taken together, these findings reveal that  $\lambda_{est}$  does not precisely characterize observed levels of phylogenetic signal in phenotypic datasets, and that biological interpretations of the strength of phylogenetic signal based on  $\lambda$  may be highly inaccurate.

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[insert Figure 2 here]

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i [insert Figure 3 here]

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## $^{_{13}}$ A Standardized Effect Size for Phylogenetic Signal

The results above demonstate that  $\lambda$  is not a reliable estimate of the phylogenetic signal in phenotypic data.

As such, biological interpretations of the strength of phylogenetic signal, and comparisons of the magnitude of such effects across datasets are severely compromised when based on this parameter. As an alternative, we propose that summary estimates of phylogenetic signal be converted to effect sizes for interpretation and comparison of the relative strength of phylogenetic signal in phenotypic datasets. Statistically, a standardized effect size may be found as:

$$Z_{\theta} = \frac{\theta_{obs} - E(\theta)}{\sigma_{\theta}} \tag{1}$$

where  $\theta_{obs}$  is the observed test statistic,  $E(\theta)$  is its expected value under the null hypothesis, and  $\sigma_{\theta}$  is its standard error (Glass 1976; Cohen 1988; Rosenthal 1994).  $Z_{\theta}$  expresses the magnitude of the effect in  $\theta_{obs}$  by transforming the original test statistic to its standard normal deviate (Glass 1976; Kelley and Preacher 2012). Typically,  $\theta_{obs}$  and  $\sigma_{\theta}$  are estimated from the data, while  $E(\theta)$  is obtained from the distribution of  $\theta$  derived from parametric theory. However, recent advances in resampling theory (Collyer et al. 2015; Adams and Collyer 2016, 2019a) have shown that  $E(\theta)$  and  $\sigma_{\theta}$  may also be obtained from an empirical sampling distribution of  $\theta$  obtained from permutation procedures.

Adams and Collyer (2019b) suggested that the strength of phylogenetic signal could be represented as an effect size, based on the *Kappa* statistic and its empirical sampling distribution from permutation. Here we

180 formalize that suggestion, resulting in an effect size of:

$$Z_K = \frac{K_{obs} - \hat{\mu}_K}{\hat{\sigma}_K} \tag{2}$$

where  $K_{obs}$  is the observed phylogenetic signal, and  $\hat{\mu}_K$  and  $\hat{\sigma}_K$  are the mean and standard deviation of the empirical sampling distribution of Kappa obtained via permutation. Similarly, an effect size based on  $\lambda$  could be envisioned as:

$$Z_{\lambda} = \frac{\lambda_{obs} - 0}{\hat{\sigma}_{\lambda}}.\tag{3}$$

In this case,  $\lambda_{obs}$  and  $\hat{\sigma}_{\lambda}$  are empirically derived using maximum likelihood. Note that under the null hypothesis, no phylogenetic signal is expected (Freckleton et al. 2002), and thus  $E(\lambda) = 0$  under this condition.

To evaluate the utility of  $Z_K$  and  $Z_{\lambda}$  we calculated both effect sizes for the simulated datasets generated 187 above, and summarized the precision of each using its variance  $(\sigma_{Z_K}^2)$  and  $\sigma_{Z_\lambda}^2$ . Results are found in Figure 4 (additional results are found in the Supporting Information). Here two things are evident. First, estimates of 189  $Z_K$  track the input phylogenetic signal in a more linear fashion than do estimates of  $Z_{\lambda}$  (Figure 4A,B). Thus, actual changes in the strength of phylogenetic signal are reflected more evenly in the corresponding values of 191 the effect size  $Z_K$ . Second, the precision of  $Z_K$  is considerably more stable as compared with  $Z_{\lambda}$ . This may be seen by calculating the coefficients of variation for the set of precision estimates (i.e.,  $\sigma_{Z_K}^2$  and  $\sigma_{Z_\lambda}^2$ ) across 193 input levels of phylogenetic signal. Here coefficients of variation in the precision of  $Z_K$  were an order of 194 magnitude smaller for than for  $Z_{\lambda}$  (Figure 4C,D). This implied that estimates of the strength of phylogenetic 195 signal were more reliable and robust when using  $Z_K$  as compared with  $Z_{\lambda}$ . 196

98 [insert Figure 4 here]

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### $Statistical \ Comparisons \ of \ Phylogenetic \ Signal$

Once the magnitude of phylogenetic signal is characterized using  $Z_K$ , one may wish to compare such measures across datasets, to determine whether the strength of phylogenetic signal is significantly greater in one phenotypic trait as compared with another. As with other effect sizes derived from permutation distributions (e.g., Adams and Collyer 2016, 2019a), a two-sample test statistic may be found as:

$$\hat{Z}_{12} = \frac{|(K_1 - \hat{\mu}_{K_1}) - (K_2 - \hat{\mu}_{K_2})|}{\sqrt{\hat{\sigma}_{K_1}^2 + \hat{\sigma}_{K_2}^2}} \tag{4}$$

where  $K_1$ ,  $K_2$ ,  $\hat{\mu}_{K_1}$ ,  $\hat{\mu}_{K_2}$ ,  $\hat{\sigma}_{K_1}$ , and  $\hat{\sigma}_{K_2}$  are as defined above for equation 2. Estimates of significance of  $\hat{Z}_{12}$  may be obtained from a standard normal distribution. Typically,  $\hat{Z}_{12}$  is considered a two-tailed test,

however directional (one-tailed) tests may be specified should the empirical situation require it (see Adams and Collyer 2016, 2019a).

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One important advantage of the approach advocated here is that the resulting effect sizes  $(Z_K)$  are dimen-209 sionless, as the units of measurement cancel out during the calculation of Z (Sokal and Rohlf 2012). As a consequence, the effect sizes represent the strength of phylogenetic signal on a common and comparable 211 scale, measured in standard deviation units, regardless of the initial units and scale of the original phenotypic variables under investigation. This means that conceivably, the strength of phylogenetic signal may be 213 compared across datsets for phenotypic traits in different units and scale (e.g., behavioral and morphological 214 traits), so long as they are continuous variables. Note that this is directly related to the common practice of 215 converting data to their standard normal deviates prior to their use in multivariate analysis, if the original 216 variables are expressed in different units or scales (see Everitt and Hothorn 2011; Legendre and Legendre 217 2012; Adams and Collyer 2019b). 218

### 219 Empirical Example

To demonstrate the utility of  $\hat{Z}_{12}$  we performed an analysis of the strength of phylogenetic signal in two phenotypic datasets from species of plethodontid salamander. The data were part of a series of studies 221 examining macroevolutionary trends in phenotypic diversification in this group (Baken and Adams 2019; Baken et al. 2020). Our dataset contained surface area to volume ratios (SA:V) and relative body width 223  $(\frac{BW}{SVL})$  for 305 species (Figure 5A). For SA:V, 11 linear body measurements from 2,781 individuals were taken, from which estimates of the surface area and volume of the head, body, and tail were calculated and 225 subsequently combined to arrive at the SA:V for each individual (for mathematical details see Baken et al. 2020). Species means were then obtained. Likewise, body size (SVL) and body width (BW) measurements 227 were taken from 3,371 individuals, and species means of relative body width  $(\frac{BW}{SVL})$  were calculated (data 228 from Baken and Adams 2019). A time-dated molecular phylogeny for the group (Bonett and Blair 2017) was then pruned to match the species in the dataset, resulting in a phylogeny and corresponding phenotypic 230 dataset containing 305 species. The phylogenetic signal in each trait was then characterized using Kappa, 231 which was converted to its effect size  $(Z_K)$  using geomorph 3.2.1 (Adams and Otárola-Castillo 2013; Adams 232 et al. 2020). Finally, the strength of phylogenetic signal was compared across traits using  $\hat{Z}_{12}$  as described above (to be incorporated in geomorph upon manuscript acceptance). 234

Results. Both SA:V and relative body width displayed significant phylogenetic signal ( $Kappa_{SA:V} = 0.7608$ ; P = 0.001;  $Kappa_{BW/SVL} = 0.2515$ ; P = 0.001). For both phenotypic traits,  $K_{obs}$  differed markedly from their corresponding permutation distributions, which were found to overlap almost perfectly (Figure 5B). 238 However, while both traits displayed significant phylogenetic signal, there was nearly a four-fold difference in the magnitude of their effect sizes, with SA:V displaying the greater phylogenetic signal (Figure 5C). 240 Using the two-sample test statistic above, this difference was found to be highly significant ( $\hat{Z}_{12} = 4.13$ ; P = 0.000036). Thus it may be concluded that SA:V displays significantly stronger phylogenetic signal 242 than does relative body width, and that shared evolutionary history has strongly influenced trait covariation 243 among taxa for SA:V. Biologically, this observation corresponds with the fact that tropical species – which form a monophyletic group within plethodontids – display greater variation in SA:V which covaries with 245 disparity in their climatic niches (Baken et al. 2020). Thus, because of this macroevolutionary association, strong phylogenetic signal in SA:V is to be expected. 247

# <sup>248</sup> Conclusions and Implications

249 1: summary paragraph

250 2: expand on Lambda.. lambda innacurate, not precise, level of precision varies with input physig (worse in mid-range). NEW RESULT. We are first to show this. NOTE: pattern is obvious with reflection. Since it is a 'bounded' parameter estimation should be best at the extremes... (state this?).. hmm.

Patterns worse with PGLS, though beta still estimated properly. Conclusion, lambda not overly useful.

3: By contrast, effect size Z-K useful, equally precise across range of values. Can be used to characterize the strength of physignal, and because robust to input levels, etc. may be used to compare across datasets.

Somewhere, recognize that this is somewhat 'backwards' from prior recommendations where Kappa had somewhat lower performance in terms of type I and type II error (which?? I forget). However, recall that those studies did not examine the precision of the estimates. Nor was Z-k included, because it was not yet invented. So Use of Z-k should make good sense here.

260 Closing paragraph.

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More discussion paragraphs

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# Figure Legends

Figure 1. Frequency distribution of  $\lambda$  estimates published in 2019. The majority of these values were close to 0 or 1, and from phylogenies with fewer than 200 taxa.

Figure 2. Precision of Pagel's  $\lambda$  across known levels of input phylogenetic signal  $(\lambda_{in})$  on phylogenies of various sizes. As phylogenies increase in size, variation in  $\lambda_{in}$  decreases; however the precision is not constant across the range of input levels  $(\lambda_{in}:0\to1)$ , and is highest at intermediate levels of phylogenetic signal.

Figure 3. Precision of Pagel's  $\lambda$  when incorporated in phylogenetic regression  $(Y \sim X)$ , across known levels of input phylogenetic signal  $(\lambda_{in})$  on phylogenies of various sizes. As phylogenies increase in size, variation in  $\lambda_{in}$  decreases; however the precision is not constant across the range of input levels  $(\lambda_{in}:0\to1)$ , and is highest at intermediate levels of phylogenetic signal.

Figure 4. Variation in estimates of phylogenetic signal across input levels of phylogenetic signal. (A) Estimates of Pagel's  $\lambda$  for data simulated on phylogenies with 128 taxa (n = 128), (B) Estimates of  $Z_K$  for data simulated on phylogenies with 128 taxa (n = 128), (C) Variance in the variation of  $\lambda_{est}$  across input levels of phylogenetic signal, estimated on phylogenies containing differing numbers of species. (D) Variance in the variation of  $Z_K$  across input levels of phylogenetic signal, estimated on phylogenies containing differing numbers of species.

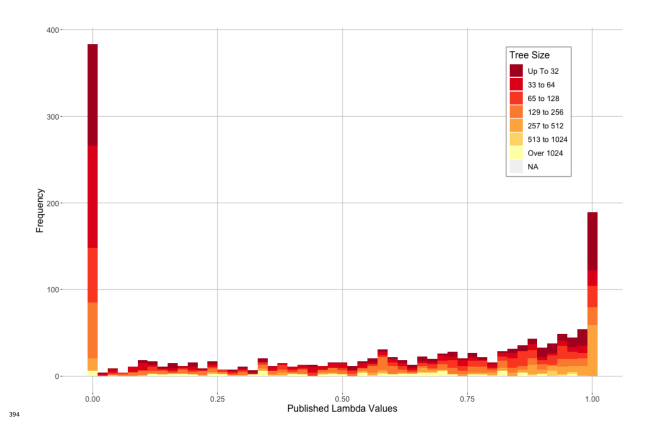


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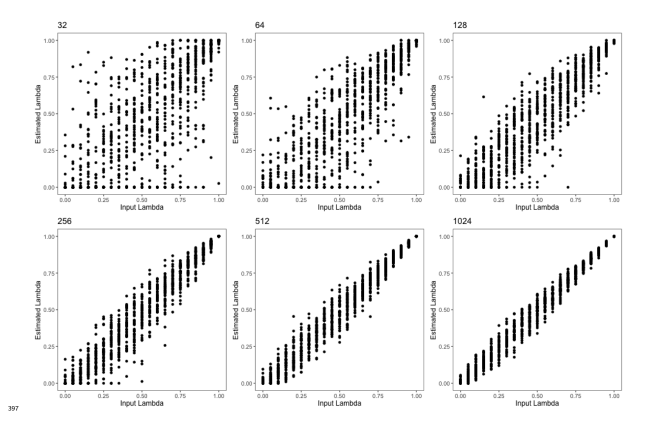


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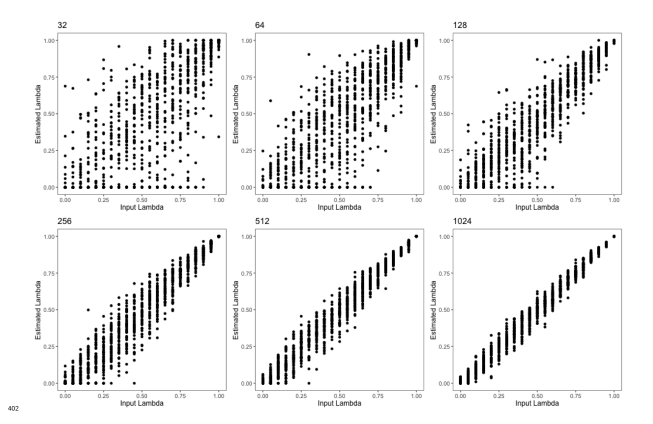


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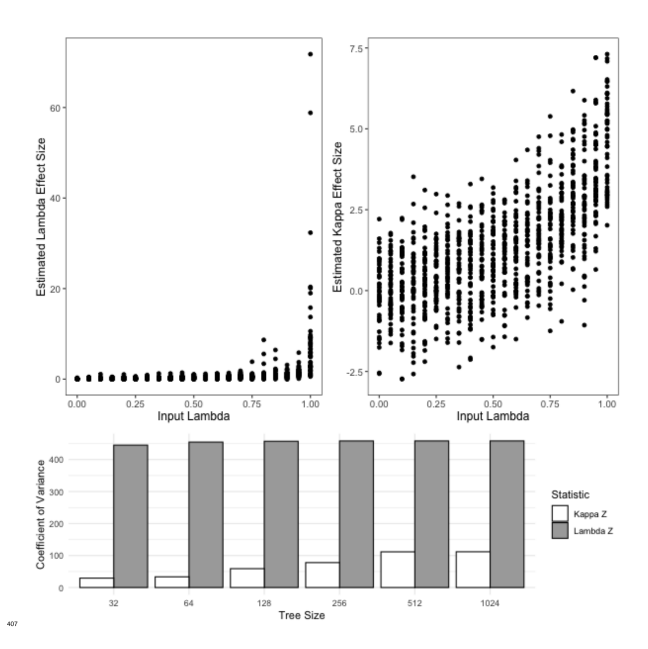


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