

1 Missing the point (estimate): Bayesian and likelihood phylogenetic
2 reconstructions of morphological characters produce generally
3 concordant inferences. A comment on Puttick *et al.* (2017)

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9 **Abstract**

10 Puttick *et al.* (2017) performed a simulation study to compare accuracy between methods
11 inferring phylogeny from discrete morphological characters. They report that a Bayesian
12 implementation of the Mk model (Lewis, 2001) was most accurate (but with low resolution),
13 while a maximum likelihood (ML) implementation of the same model was least accurate. They
14 conclude by strongly advocating that Bayesian implementations of the Mk model should be the
15 default method of analysis for such data. While we applaud investigations into accuracy and
16 alternative methods of analysis, this conclusion is based on an inappropriate comparison of the
17 ML point estimate with the Bayesian consensus. We revisit these issues through simulation by
18 considering uncertainty in ML reconstructions, and demonstrate that Bayesian and ML estimates
19 are generally concordant when conventional edge support thresholds are considered. We therefore
20 disagree with the conclusions of Puttick *et al.* (2017), and consider their prescription of *any*
21 default method to be unfounded. Instead, we recommend caution and thoughtful consideration of
22 the model or method being applied to a morphological dataset.

23 **Key words:** phylogeny, morphology, paleontology, Bayesian, likelihood

24 **Comparing point estimates to consensus summaries**

25 Puttick *et al.* (2017) report that ML tree inference under the Mk model results in higher
26 topological error than Bayesian implementations. However, this result is driven precisely by the
27 comparison of maximum likelihood point estimates (MLE) to Bayesian majority-rule (BMR)
28 consensus trees. MLE topologies are fully resolved, but this stems from the standard binary tree
29 searching algorithms employed and not from an explicit statistical rejection of unresolved nodes.
30 Therefore, individual MLE estimates may contain edges with negligible statistical support. On
31 the other hand, consensus summaries, independent of phylogenetic method, may have reduced

32 resolution as a product of uncertainty arising by summarization across conflicting sampled
33 topologies. Thus, a direct comparison between a consensus tree (i.e., BMR) and a point estimate
34 (i.e., MLE) is inappropriate. BMR topologies of Puttick *et al.* (2017) are more accurate because
35 poorly supported conflicted edges were collapsed, while MLE topologies were fully resolved,
36 even if poorly supported. While contrasting MLE and Bayesian maximum *a posteriori* (MAP)
37 trees would be a more appropriate comparison of optimal point estimates, the incorporation of
38 uncertainty is an integral part of all phylogenetic analysis. Therefore, comparison of consensus
39 trees from Bayesian and ML analyses hold more practical utility for systematists. For these
40 reasons, we argue that the results of Puttick *et al.* (2017) are an artifact of their comparison
41 between fundamentally incomparable sets of trees.

42 **Support metrics are available for morphological characters**

43 To avoid drawing untenable conclusions, it is *de rigueur* of any phylogenetic analysis to explicitly
44 assess edge support. Systematists often accomplish this via non-parametric bootstrap sampling
45 (Felsenstein, 1985), though other measures exist (see below). Puttick *et al.* (2017) did not assess
46 edge support in their ML estimates, stating that morphological (but not genetic) data do not meet
47 an underlying assumption of the bootstrap statistical procedure that phylogenetic signal is
48 distributed randomly among characters. The authors do not explain the meaning of this statement,
49 and no references are provided to support the assertion. Non-parametric bootstrapping has been a
50 staple of phylogenetic reconstruction for decades, including for the analysis of discrete
51 morphological characters. Bootstrapping works via the assumption that the observed characters
52 are a representative sample from a population of possible characters evolving under the same
53 process, and thus can be resampled to assess confidence in parameters (Felsenstein, 1985). While
54 morphological matrices typically include only variable characters (i.e., an ascertainment bias),
55 this is an informative subset of the possible characters, and should not be thought of as misleading
56 calculations. Were this otherwise, the original sample would be likewise suspect, as the use of
57 model-based phylogenetic inference (such as Mk) explicitly assumes characters evolve according
58 to the same process. Concerns about the interpretation and use of the bootstrap exist (Sanderson,
59 1995), the primary of which involves the assumption that individual characters are statistically
60 independent. However, it is reasonable to assume that individual sites in a morphological matrix
61 would be more independent than adjacent sites from the same gene, and genetic datasets are
62 routinely bootstrapped. We therefore disagree with the claims of Puttick *et al.* (2017) that
63 bootstrapping is inappropriate for morphological data, or at least any *more* inappropriate than for
64 genetic data.

65 There are also other methods researchers can use to assess edge support in a likelihood
66 framework. Jackknifing, unlike bootstrapping, samples without replacement, conditioning on
67 strict subsets of the observed data. More recently, the SH-like test (Guindon *et al.*, 2010)
68 computes support for each internal edge in the MLE tree by considering all nearest neighbour
69 interchanges (NNIs). This test is implemented in several software packages including RAxML
70 (Stamatakis, 2014), one of the programs used by Puttick *et al.* (2017). Alternatively, ML
71 programs frequently offer an option to collapse edges on a MLE tree that fall below some
72 minimum threshold length. Use of any of these options would enable a fairer comparison of
73 likelihood and Bayesian reconstructions.

74 **ML and Bayesian comparisons incorporating uncertainty**

75 To measure the effect of comparing BMR and MLE trees, we used the simulation code from
76 Puttick *et al.* (2017) to generate 1000 character matrices, each of 100 characters on a fully
77 pectinate tree of 32 taxa, as these settings generated the most discordant results in Puttick *et al.*
78 (2017). Each matrix was analyzed in both Bayesian and ML frameworks using the Mk+G model
79 (Lewis, 2001). Bayesian reconstructions were performed using MrBayes v3.2.6 (Ronquist *et al.*,
80 2012), using the same settings as Puttick *et al.* (2017): 2 runs, each with 5×10^5 generations,
81 sampling every 50 generations, and discarding the first 25% samples as burnin. As in Puttick
82 *et al.* (2017), we summarized each analysis with a BMR consensus tree (i.e. only edges with \geq
83 0.5 posterior probability are represented). Likelihood analyses were performed in RAxML v8.2.9
84 (Stamatakis, 2014). For each simulated matrix we inferred both the MLE tree and 200
85 nonparametric bootstrap trees. Accuracy in topological reconstruction was assessed using the
86 Robinson-Foulds (RF) distance (Robinson and Foulds, 1981), which counts the number of
87 unshared bipartitions between trees. We measured the following distances from the true simulated
88 tree: d_{BMR} , the distance to the Bayesian majority-rule consensus; d_{MLE} , the distance to the MLE
89 tree; d_{ML50} , the distance to the MLE tree which has had all edges with $<50\%$ bootstrap support
90 collapsed. Finally, for each matrix we calculate $D_{\text{MLE}} = d_{\text{MLE}} - d_{\text{BMR}}$, and $D_{\text{ML50}} = d_{\text{ML50}} - d_{\text{BMR}}$.
91 These paired distances measure the relative efficacy of ML and Bayesian reconstructions: values
92 of D greater than 0 indicate that ML produces less accurate estimates (that is, with a greater RF
93 distance from the true generating tree).

94 As demonstrated by Puttick *et al.* (2017), MLE trees are indeed less accurate than BMR trees
95 (Figure 1; D_{MLE}), with MLE trees on average having an RF distance 17.6 units greater than the
96 analogous Bayesian consensus distance. However, when collapsing MLE edges with less than
97 50% bootstrap support, Bayesian and ML differences are normally distributed around 0 (Figure 1;
98 D_{ML50}), indicating that when standardizing the degree of uncertainty in tree summaries there is no
99 difference in topology reconstruction accuracy. These results support the argument that the
100 original comparisons made in Puttick *et al.* (2017) of MLE and BMR trees are inappropriate.
101 Depending on the level of uncertainty involved, an optimal point estimate from a distribution
102 (e.g., MLE or MAP) may be arbitrarily distant from a summary of the same distribution. And so,
103 the differences in MLE vs. BMR are not expected to be consistent.

104 **The expected concordance of Bayesian and ML results**

105 Our results reveal much greater congruence between Bayesian and ML estimates than suggested
106 by Puttick *et al.* (2017). This is to be expected. ML and Bayesian tree construction methods
107 should yield similar results under the conditions in which they are often employed. While
108 Bayesian tree reconstruction differs from ML by incorporating prior distributions, the methods
109 share likelihood functions. In phylogenetics, researchers typically adopt non-informative priors,
110 with a few exceptions (e.g., priors on divergence time parameters). Arguments can be made for
111 pseudo-Bayesian approaches when care is taken to ensure that priors used are truly uninformative,
112 which result in posterior probabilities that mirror the likelihood and are therefore congruent with
113 ML (Alfaro and Holder 2006; Gelman *et al.* 2014). If prior distributions are formulated
114 thoughtfully, as with Wright *et al.* (2016), in shaping the Mk model using hyperpriors to
115 accommodate character change heterogeneity, Bayesian methods can outperform ML.

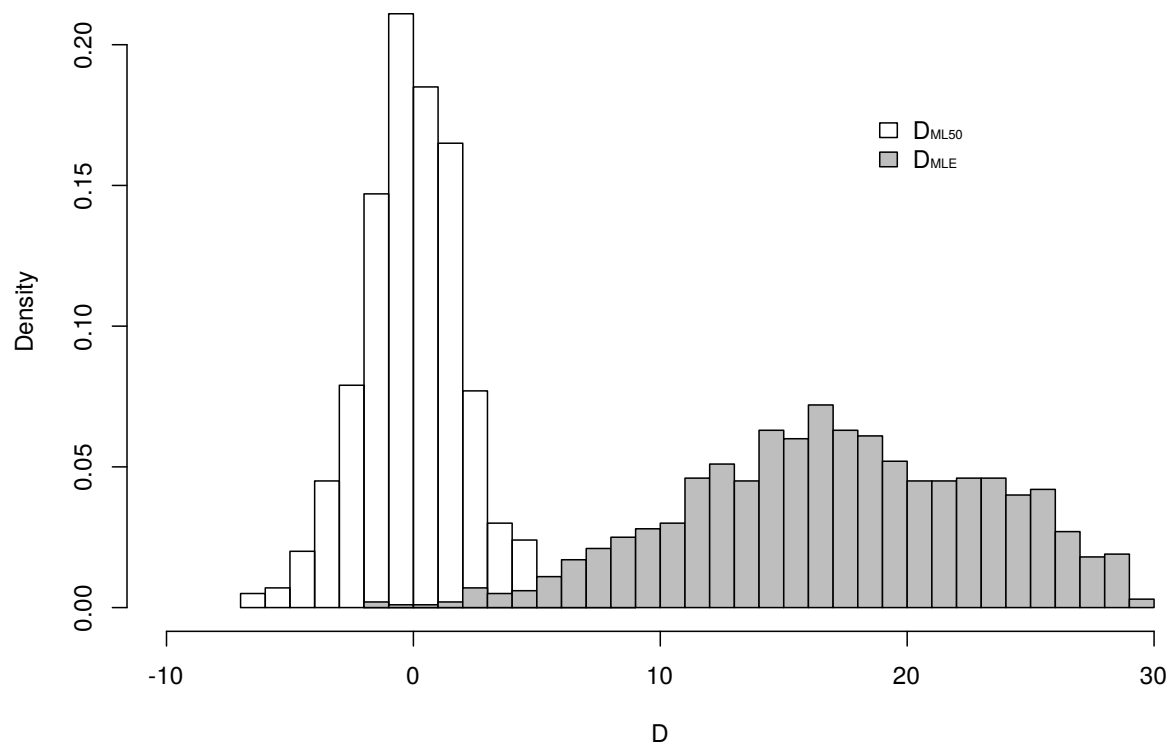


Figure 1: Topological accuracy of ML vs. Bayesian reconstructions. D measures how much larger ML distances are from the true tree (d_{ML}) than are Bayesian distances (d_{BMR}). MLE trees are indeed less accurate than BMRs (D_{MLE}), but when conventional bootstrap thresholds are employed (D_{ML50}) the difference in efficacy disappears.

116 Alternatively, inappropriate priors can positively mislead (Gelman *et al.*, 2014). Generally, when
117 informative prior distributions are known or can be estimated using hierarchical approaches,
118 Bayesian reconstruction methods may be strongly favored over ML. It is unclear whether Puttick
119 *et al.* (2017) intend to draw the comparisons discussed above as they do not describe any reasons
120 to prefer Bayesian over ML in principle.

121 Although our results demonstrate general concordance between ML and Bayesian approaches
122 when uncertainty is represented, further simulation work is needed to determine the extent and
123 conditions of this concordance. Issues surrounding the application of Bayesian methods are
124 particularly important in paleontology, where researchers often conduct inference upon very
125 limited data. In these cases, it may be desirable to construct informative prior distributions when
126 conducting Bayesian analyses (Gelman *et al.*, 2014). The questions posed by Puttick *et al.* (2017)
127 are critically important as statistical morphological phylogenetics moves forward. However, their
128 inappropriate comparison between ML and Bayesian approaches leaves the relative performance
129 of the two implementations of the Mk model unresolved.

130 We conclude by stating that we are not advocating one method over another for morphological

131 phylogenetic reconstruction. Methods differ in model (Mk vs. parsimony), inferential paradigm
132 (parsimony vs. ML/Bayesian), assumptions (prior distributions, model adequacy), interpretation,
133 and means to incorporate uncertainty (ML/parsimony vs. Bayesian). We therefore recommend
134 caution and thoughtful consideration of the biological question being addressed and then
135 choosing the method that will best address that question. All inferential approaches possess
136 strengths and weaknesses, and it is the task of researchers to determine the most appropriate given
137 available data and the questions under investigation. The excitement of new morphological data
138 sources and new means for analyzing these data should not overshadow the obligation to apply
139 methods thoughtfully.

140 **Authors' contributions**

141 J.W.B. conceived the design of the study and performed the analyses; J.W.B. and C.F.-P. drafted
142 the manuscript; all authors contributed to the interpretation of results and the writing of the
143 manuscript.

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