Using Parsimony-Guided Tree Proposals to Accelerate Convergence in Bayesian Phylogenetic Inference

CHI ZHANG1,2, JOHN P. HUELSENBECK3, and FREDRIK RONQUIST4,*

1Key Laboratory of Vertebrate Evolution and Human Origins, Institute of Vertebrate Paleontology and Palaeoanthropology, Chinese Academy of Sciences, Beijing 100044, China; 2Department of Integrative Biology, University of California, Berkeley, CA 94720, USA; and 3Department of Bioinformatics and Genetics, Swedish Museum of Natural History, Box 50007, SE-10405 Stockholm, Sweden

*Correspondence to be sent to: Department of Bioinformatics and Genetics, Swedish Museum of Natural History, Box 50007, SE-10405 Stockholm, Sweden; E-mail: fredrik.ronquist@nrm.se

Received 22 February 2018; reviews returned 15 January 2020; accepted 17 January 2020

Abstract.—Sampling across tree space is one of the major challenges in Bayesian phylogenetic inference using Markov chain Monte Carlo (MCMC) algorithms. Standard MCMC tree moves consider small random perturbations of the topology, and select from candidate trees at random or based on the distance between the old and new topologies. MCMC algorithms using such moves tend to get trapped in tree space, making them slow in finding the globally most probable trees (known as “convergence”) and in estimating the correct proportions of the different types of them (known as “mixing”). Here, we introduce a new class of moves, which propose trees based on their parsimony scores. The proposal distribution derived from the parsimony scores is a quickly computable albeit rough approximation of the conditional posterior distribution over candidate trees. We demonstrate with simulations that parsimony-guided moves correctly sample the uniform distribution of topologies from the prior. We then evaluate their performance against standard moves using six challenging empirical data sets, for which we were able to obtain accurate reference estimates of the posterior using long MCMC runs, a mix of topology proposals, and Metropolis coupling. On these data sets, ranging in size from 357 to 5681 sites, we find that single chains using parsimony-guided moves usually converge an order of magnitude faster than chains using standard moves. They also exhibit better mixing, that is, they cover the most probable trees more quickly. Our results show that tree moves based on quick and dirty estimates of the posterior probability can significantly outperform standard moves. Future research will have to show to what extent the performance of such moves can be improved further by finding better ways of approximating the posterior probability, taking the trade-off between accuracy and speed into account. [Bayesian phylogenetic inference; MCMC; parsimony; tree proposal.]

Introduced to the field two decades ago (Rannala and Yang, 1996; Mau and Newton, 1997; Yang and Rannala, 1997; Li et al., 2008), Bayesian estimation of phylogeny has become widely used by evolutionary biologists (see Hueslenbeck et al., 2001; Holder and Lewis, 2003; Yang and Rannala, 2012; Yang, 2014; Nascimento et al., 2017, for reviews). The Bayesian framework is attractive for many reasons, including the simple interpretation of results, the ability to address interesting evolutionary questions, and the availability of efficient and easy-to-use computer programs that implement it (Ronquist and Huelsenbeck, 2003; Drummond and Rambaut, 2007; Ronquist et al., 2012b; Bouckaert et al., 2013; Höhna et al., 2016).

Bayesian estimation of phylogeny almost always relies on Markov chain Monte Carlo (MCMC: Metropolis et al., 1953; Hastings, 1970) to sample trees in proportion to their posterior probability. (For exceptions, see Bouchard-Côté et al., 2012; Wang et al., 2016). The MCMC procedure typically uses a mixture of different proposal mechanisms (also called moves or operators) that change one or a few of the parameters in the model. Unfortunately, devising MCMC proposals that sample well across the space of evolutionary trees is quite challenging. For any reasonable number of tips, a MCMC proposal has a huge space of tree topologies to explore. It also needs to tackle the complex dependencies between topology and branch lengths. Most MCMC tree proposals studied and used in current Bayesian MCMC phylogenetic software are stochastic versions of tree-perturbation methods that were originally used to find optimal parsimony or maximum likelihood trees in hill-climbing algorithms. The basic perturbation methods include nearest neighbor interchange (NNI), subtree pruning and regrafting (SPR), and (for non-clock trees) tree bisection and reconnection (TBR) (Larget and Simon, 1999; Lakner et al., 2008). Each of these mechanisms can generate a set of candidate trees from the current one by applying a particular type of tree modification.

In a hill-climbing algorithm, one would typically score all candidate trees generated by one of these mechanisms, and then choose the best one. In the MCMC context, a new tree is instead drawn from a suitable probability distribution over the candidate trees. The simplest choice is a uniform distribution, and this is still a common choice for stochastic NNI (sNNI) moves. Uniformly random SPR and TBR proposals, however, are inefficient because they tend to make such drastic changes that the new tree has negligible posterior probability and will be rejected. A better option is to bias SPR and TBR moves toward more modest changes as in the “extending” SPR (eSPR) and “extending” TBR (eTBR) moves introduced in MrBayes (Huelsenbeck and Ronquist, 2001; Ronquist and Huelsenbeck, 2003). These proposals move the regrafting point away from the pruning point in a stepwise fashion, applying a constant probability in each step to decide whether the
distance should be further extended. sNNI, eSPR, and eTBR are among the most efficient MCMC tree moves known for non-clock trees (Lakner et al., 2008), but given the frequent observation of empirical tree spaces that fail to yield unit root proposals, even when Metropolis-Hastings coupling (Geyer, 1991) is used, there is clearly room for improvement.

One possibility would be to choose among candidate trees according to their posterior probability. This is known as a “Gibbs” proposal (Liu, 2004), and it will always be accepted. The mixing rate of a Gibbs proposal can be further improved by “Metropolizing” the move, that is, by removing the probability of proposing the starting state as the new state (Peskun, 1973). Unfortunately, it is computationally costly to obtain the posterior probability of any set of alternative trees that is reasonably large. Therefore, Gibbs or Metropolized Gibbs proposals are quite difficult to implement except in some very unusual circumstances (Huelsenbeck et al., 2008). Gibbs sampling is a special case of the Metropolis–Hastings algorithm. In general, the proposal ratio (also known as the “Hastings ratio”) of the Metropolis–Hastings algorithm is used to adjust the acceptance ratio, correcting for any asymmetry that may arise between forward and backward proposals.

Instead of using posterior probabilities directly, we could guide tree proposals based on an approximation of the posterior distribution over topologies obtained in a previous analysis (Höhna and Drummond, 2012). Such approximations are best computed from the sampled split (taxon bipartition) frequencies, either alone or in combinations of two or three adjacent splits (Ronquist et al., 2004; Höhna and Drummond, 2012; Larget, 2013). Guiding tree proposals based on such approximations can significantly improve their performance (Höhna and Drummond, 2012). A potential concern with this approach is that the efficiency of the final run could be affected by any errors or biases in the approximation of the posterior distribution over topologies obtained in the preliminary run. In other words, this approach pushes the most difficult challenge for tree proposals to the preliminary analysis, when an approximation of tree space is not available.

In this article, we explore another idea, namely to guide tree proposals based on a quick and dirty approximation of the posterior probability of trees, computed on the fly. Specifically, we base the approximation on the parsimony score of topologies, inspired by the links that do exist between parsimony scores and probability (Huelsenbeck et al., 2008). To our knowledge, such parsimony-guided tree proposals were first introduced in MrBayes 3.2 (Ronquist et al., 2012b), where they were included in the default set of tree moves based on promising preliminary experiments. In the development of ExaBayes (Aberer et al., 2014), parsimony-guided tree proposals modeled after MrBayes were found to be essential for convergence on large data sets (Aberer A., personal communication). However, these proposals have never been described in the literature, nor have their properties been examined in detail.

Here, we describe a couple of variants of parsimony-guided pSPR (pSPR) and parsimony-guided TBR (pTBR) proposals, which have been implemented in MrBayes. We use simulations to verify the implementations, and we use analyses of empirical data sets to show that pSPR and pTBR moves significantly outperform eSPR and eTBR both in terms of convergence and mixing. Tree moves have received surprising little attention, given their importance in Bayesian MCMC phylogenetics (for exceptions, see Höhna et al., 2008; Lakner et al., 2008; Höhna and Drummond, 2012; Whidden and Matsen, 2015). We hope that this article will stimulate further work on guided MCMC proposals and other ideas for improving MCMC exploration of tree space.

Materials and Methods

In this section, we first describe the tree proposals used in the article. We then show the simulations we used to verify the implementation of the parsimony-guided proposals. Finally, we describe the empirical tests we used to evaluate the performance of the parsimony-guided tree proposals against standard extending tree proposals.

All algorithms were implemented in MrBayes version 3.2.7, available from GitHub (https://github.com/NBISweden/MrBayes). The MrBayes scripts used in the analyses are available as Supplementary Material available on Dryad at http://dx.doi.org/10.5061/dryad.98mp657. We focus the description here on the tree proposals; details of the other proposals and prior settings used in the analyses are available in the MrBayes scripts and in the manual.

Tree Proposals

We focus entirely on tree proposals for unrooted trees. Specifically, we explore two variants each of pSPR and pTBR, and compare their performance to eSPR and eTBR. In the reference runs, we also use sNNI. For a more complete description of the MCMC context of these proposals (see Appendix); here, we focus only on the essential details needed to describe them unambiguously. Note that the eSPR move used here is different from that examined in a previous study (Lakner et al., 2008) in that the initial branch is picked from all branches in the move, rather than only from internal branches.

Nearest Neighbor Interchange.—First, an internal branch is picked at random. Label the branches with subtrees that are incident to the chosen branch A, B, C, and D, such that the original topology is ((A,B),C,D). Without loss of generality, assume that D stays in its original position. Then the topology is modified by exchanging A and C, or B and C. In the stochastic Metropolized version...
of the move, denoted sNNI here, the two alternative topologies are chosen with equal probability. The move leaves the topology with the same taxon bipartitions except for the bipartition associated with the initially chosen internal branch. This gives a natural mapping of branches between the trees. The branch lengths of the old tree are applied unmodified to the new tree using this mapping. The proposal ratio is 1.0.

**Subtree Pruning and Rerooting**—The SPR move first picks a branch r (internal or external, with length \( v_r \)) at random. It then randomly selects one of the two subtrees incident to the branch; label this subtree A (Fig. 1a). If \( r \) is a terminal branch, \( A \) will be the tip node. A branch \( r \) in the remaining subtree, with length \( v_r \), is chosen for rerooting of A (Fig. 1a). The move is Metropolized, such that a topology with \( r \) is guaranteed. This means that branch \( b \) or \( q \) cannot be selected as \( r \). The exact mechanism for choosing \( r \) varies depending on the type of SPR move (see below). The branch connecting \( a \) in the moving direction (i.e., the branch closer to \( r \), \( q \) in this case, with length \( v_q \)) is also pruned away and moved together with \( A \), leaving the remaining subtree BCDE (Fig. 1b). Subtree A with branches \( a \) and \( q \) is inserted before branch \( r \) connecting subtree E, resulting in \( r \) connecting \( a \) and \( q \) (Fig. 1c). Note that NNI is a special case of SPR: If \( A \) is moved one node away to \( C \) (\( c \) is chosen for rerooting) or DE (\( p \) is chosen for rerooting), this is equivalent to a NNI move around branch \( q \).

The eSPR move applies an extension mechanism to choose \( r \); with probability 1/2, it moves the rerooting point one branch away in either direction (i.e., into the B or the CDE subtree). Then with probability \( p_r \) (the extension probability), the rerooting point is moved one branch further, and with probability \( 1 − p_r \) it stays at the current location (Fig. 6 in Lakner et al., 2008). If the branch is moved one branch further away, one of the two possible directions is chosen with probability 1/2, and the cycle is repeated. If the extension mechanism encounters a tip, it stops. We refer to a proposal that encounters a tip as a “constrained” proposal. The proposal ratio is 1.0 if the extension mechanism is unconstrained or constrained in both directions of the move. If the move is constrained in just one direction, the proposal ratio becomes \( 1/(2(1 − p_r)) \) (backward move constrained) or \( 2(1 − p_r) \) (forward move constrained) (Lakner et al., 2008).

The pSPR moves use a mechanism based on parsimony scores to choose \( r \). Let \( B \) be the set of all branches in subtree BCDE, that is, the tree remaining after subtree A is pruned away (Fig. 1b). The weight for proposing branch \( b \) for rerooting of A, \( w_b \), is given by:

\[
\omega_b = \rho_b e^{S_b},
\]

where \( \epsilon \) is the warp factor, \( \rho_b \) is the base factor, and \( S_b \) is the parsimony score of the tree after rerooting A at \( b \), minus the sum of the scores of the two subtrees A and BCDE. We call the tuning parameter \( \epsilon \) the warp factor because it determines how much the parsimony score will influence the weight, that is, the extent to which a uniform probability distribution over candidate trees is modified by the parsimony score. The larger the warp factor, the more heavily the parsimony score will influence the probability of candidate trees being proposed.

Assume that in the set \( B \), the branch at the pruning point is \( b \) and the branch at the rerooting point is \( r \). Then the proposal ratio of a pSPR move is

\[
\frac{\omega_b}{\sum_{b \in B \setminus \{r\}} \omega_b} \div \frac{\omega_r}{\sum_{b \in B \setminus \{b\}} \omega_b}
\]

where \( B \setminus \{b\} \) is the set of all branches in \( B \) except \( b \). The weights are constructed such that candidate trees with lower (better) parsimony scores receive higher weights and are proposed more often. The proposal ratio corrects for this preference: a proposed move to a more parsimonious tree is likely to be accepted only if the proposal bias is matched by a similar or larger increase in likelihood. Thus, a correctly implemented parsimony-guided proposal will not be affected by parsimony artifacts, such as long-branch attraction (LBA).

Here, we employ two schemes for converting parsimony scores to proposal weights. In the first scheme, pSPR1, we simply set all \( \rho_b = e^{-\epsilon S_b} \). In the second scheme, pSPR2, we try to accommodate the branch length effect: the higher the parsimony score for a branch, the longer the branch is likely to be in a probabilistic context, and the less its parsimony score should influence our preference among candidate trees. To justify the correction we use, it is helpful to consider the probability of one sequence evolving into another over some length of time \( \tau \) under the JC69 model (Jukes and Cantor, 1969). This will be a product of two factors: the probability of the ending state being the same as the starting state, \( p_0 \), and the probability of it being different, \( p_1 \), both functions of \( \tau \). Say that we observe that \( S \) of \( N \) sites are different. Then the overall probability will be \( p_1 \tau^S p_0^{N-S} \). Thus, the overall probability is proportional to the ratio \( p_1(\tau)/p_0(\tau) \) raised to \( S \), the number of sites that are different. The number of sites that are different is of course the same as the parsimony score. If \( \tau \) is small, \( p_1(\tau) \) should be sufficiently close to 1 that we can ignore it. Assuming that the JC69 model is a reasonable approximation of more complex substitution models, we set \( \rho_b = p_1(\tau) = \frac{3}{4}(1 - e^{-1.5\tau^S}) \), where \( \tau^S \) is the length of branch \( b \), as an estimate of \( \tau^S \), which can be computed easily, we use \( \tau^S = S_b/N + \eta \), where \( N \) is the total number of sites, and \( \eta \) is a small positive number (0.00001) to avoid \( \rho_b = 0 \) when \( S_b = 0 \). Table 1 gives some examples of pSPR1 and pSPR2 weights for different parsimony scores, branch lengths and choice of tuning parameters.
branch labeling used above (Fig. 1). Then the lengths of the picked branch \( (v) \) is picked at random from all branches in the tree. One of the subtrees incident to \( a \) is then picked at random and pruned away; this subtree is labeled A here.

When the branch \( q \) in the moving direction (the pendant branch) is moved together with \( a \), and is adjacent to \( r \), resulting in \( r \) connecting \( a \) and \( q \). The specific SPR variants differ in how they pick \( r \).

For all TBR moves, branch lengths are mapped from the old tree to the new tree as for the SPR moves, and then the lengths of the chosen internal branch and the two pendant branches moved with it, one on each side, are modified using independent scaler moves in a fashion analogous to that of the SPR moves.

### Tuning Parameters

The tuning parameter values used for the tree moves in the analyses are summarized in Table 2. Choosing \( p_e = 0.5 \) means that eSPR will propose NNI changes half the time (even more often if the extension mechanism is constrained by tips), and more radical changes the rest of the time. For eTBR, the same setting means that it will propose NNI changes 1/4 of the time. Choosing \( \beta = 5 \) for a pTBR move generates up to approximately \( 2^{6p_k} = 2^{12} \) = 4096 candidate trees when there are no tip constraints. A pSPR move on a tree with \( n \) tips will generate up to roughly \( 2n \) candidate trees, that is, from around 700 to around 1900 candidate trees for the empirical data sets analyzed here. The tuning parameter of the scaler moves used to modify branch lengths was set to \( 2 \ln 1.05 \), yielding proposed changes of maximally 5% up or down in branch lengths (Larget and Simon, 1999).

### Simulations

We performed simulations to verify the implementation of the parsimony-guided tree proposals. Importantly, we tested whether the proposal ratio was correctly implemented so that we could retrieve the uniform prior distribution on topologies when the likelihood was set to be constant (zero).

The tree of five taxa used has two long branches separated by a short one (Fig. 2a). The parsimony score...
Figure 2. Simulation used to study the effect of long-branch attraction on the parsimony-guided tree proposals. a) The true tree used in the simulation, with two long branches separated by a short one. b) The tree inferred using maximum parsimony, erroneously joining long branches.

Table 2. Tuning parameter values used here for the tree proposals

| Proposal | Tuning parameters |
|----------|------------------|
| eSPR    | \( p_c = 0.5 \) |
| eTBR    | \( p_c = 0.5 \) |
| pSPR1   | \( \epsilon_1 = 0.5 \) |
| pTBR1   | \( \epsilon_1 = 0.5, \lambda = 5 \) |
| pSPR2   | \( \epsilon_2 = 0.1 \) |
| pTBR2   | \( \epsilon_2 = 0.1, \lambda = 5 \) |

Note: In addition to the tuning parameters listed in the table, all proposals used scaler moves with a tuning parameter of \( 2\ln10.05 \) to change selected branch lengths.

Empirical Data

Evaluating the performance of tree proposals empirically is challenging. The problems need to be difficult enough to distinguish the performance of different topology proposals, while not being computationally too demanding to prohibit numerous repeated MCMC runs. Standard tree proposals under Metropolis-coupled MCMC are quite efficient in sampling from many small or simple tree spaces. Therefore, an obvious choice is to run empirical evaluations of tree proposals without Metropolis coupling, which makes the problem harder while bringing down the computational cost. The empirical data sets need to be reasonably large, with moderate to large number of taxa, as the parsimony-guided proposals should be particularly advantageous when sampling from large tree spaces. For this study, we chose to run the critical tests without Metropolis coupling but using reasonably large empirical data sets under a more realistic evolutionary model than JC69, namely the general time reversible model with gamma rate variation across sites (GTR + \( \Gamma_4 \); Yang, 1994). We assumed a single partition to avoid any confounding influence of the partitioning scheme.

For empirical data, we do not know the true posterior distribution over trees. However, it may be possible to generate a reference sample of the tree space, which can be used as a reasonably accurate approximation of the true posterior. Here, we chose to start from a number of empirical data sets of suitable size. We then ran an
We then ran 16 independent single-chain analyses starting from a random tree and was sampled every 1000 generations. The details are available in the MrBayes scripts provided as Supplementary Material available on Dryad.

To visualize the structure of the tree spaces, we used multidimensional scaling (MDS) based on the SPR distance between sampled trees (Whidden and Matsen, 2015). MDS methods compute a low-dimensional space (two-dimensional in our case) that represents a distance matrix (the minimum SPR distances between trees in our case) as accurately as possible. Calculating SPR distances among all sampled trees was prohibitively time consuming. Therefore, we randomly subsampled 4000 trees (~10%) and calculated their pairwise SPR distances using the rspr software (Whidden et al., 2010, 2013). The sampled trees were then plotted in the resulting space.

### RESULTS

#### Simulations

When the likelihood was constant (no data), and no sites were used to compute the parsimony guide weights, all tree proposals tested were able to retrieve the uniform prior distribution over topologies with good accuracy (Fig. 3). When 100 sites (Fig. 4a) or 1000 sites (Fig. 4b) were used to compute the parsimony weights, and likelihoods were artificially kept constant to test the ability of the proposal ratio to correct for the proposal bias, we still retrieved the uniform topology. However, when all 10,000 sites were included in computing the parsimony weights, some topologies were undersampled by the pSPR moves because of numerical inaccuracies in computing proposal ratios. Specifically, this affected the trees with the worst parsimony scores (topology 14 and 15 in Fig. 4), the trees which are unlikely to be sampled at all in a real analysis because they also have low likelihood scores. Examining the simulated data sets revealed that the likelihood scores of topology 14 and 15 were around 300 log units lower than the scores of the best tree when all 10,000 sites were considered. This means that a correct MCMC algorithm sampling from the posterior should visit them considerably less than a googolth ($10^{-100}$) as often as the best tree. Thus, for all practical purposes, the numerical errors affecting the sampling of these trees can be ignored. Importantly, the numerical errors did not cause oversampling of the LBA tree (topology 2) in relation to the best tree (topology 1) even under this extreme setting.

Interestingly, the pTBR moves were not affected even when all 10,000 sites were included in computing the parsimony weights (Fig. 4c). The reason is that they only pick interior branches, and therefore do not directly connect the worst trees (topology 14 and 15) with the best trees (topology 1 and 2) in a single move. This means that the pTBR proposal ratios are less extreme, making pTBR less sensitive to numerical errors in the computation of those ratios. However, this difference is of no consequence, as already the pSPR move is robust enough to numerical errors for all practical purposes.

---

**Table 3.** Empirical data sets used in this study

| Abbreviations | Taxa Sites | ID References |
|---------------|------------|---------------|
| SQ10          | 357 2925   | — Savolainen et al. (2000) |
| DA10          | 357 4493   | S1008 Davis and Anderson (2010) |
| LZ12          | 425 2468   | S12425 Lu et al. (2012) |
| CL12          | 459 1760   | S12988 Cardoso et al. (2012) |
| AZ12          | 545 5681   | S11973 Aliscioni et al. (2012) |
| NPII          | 934 3675   | S11260 Nagy et al. (2011) |

Notes: The abbreviation (initial of first and last authors and publication year), number of taxa, number of sites, TreeBase study ID, and reference for each data set are given. SQ18 was only found in GenBank (see Appendix of Savolainen et al., 2000).
FIGURE 3. Test of the ability of the tree proposals to retrieve a uniform prior over topologies without data. We used a tree with five tips and a constant likelihood. The estimated posterior probabilities are represented as the mean (dot) and range (bar) across 10 replicate analyses. The prior probability is $\frac{1}{15} \approx 0.0667$ represented as horizontal dashed line. Topology 1 is the true tree (Fig. 2a) and topology 2 is the LBA tree (Fig. 2b).

Empirical Data

For six out of the 20 data sets, we were able to obtain reasonably accurate estimates of the posterior distribution in the reference runs (ASDSF ≤ 0.02, Table 4). Of the remaining 14 data sets, ASDSF values reached below 0.05 for three, and another three had ASDSF values below 0.10 (Supplementary Table S2 available on Dryad). For the rest of the data sets, the tree samples obtained in the four independent analyses were even more heterogeneous at 20 million generations, and it seemed for a few of them that the runs would have had to be extended considerably to obtain good estimates of the posterior distribution.

Henceforth, we will focus on the six data sets for which we were able to generate reasonably accurate estimates of the posterior. Table 5 summarizes some characteristics of these data sets and the posteriors. The data sets span over a considerable range of tree shapes and percentages of missing data. Note that, for all six data sets, each tree topology in the posterior is sampled only once, so that the 95% credible sets all contain 45604 unique trees, that is, a unique tree for each sample from the posterior. This is because the posterior probability is spread out rather evenly over a very large number of topologies. Thus, our analyses only generate a limited subsample of the true credible set, which is likely to be much larger. Despite this shortcoming, the ASDSF diagnostic indicates that the reference runs have converged in the sense that the sampled trees are all representative of the true credible set in terms of split frequencies.

We analyzed distances and computed tree spaces for a subsample of 4000 trees from each of the posterior samples. The minimum SPR distance among the trees in these subsamples ranged from 20 to over 200. The visualizations show that the tree spaces are quite different (Fig. 5). There are distinct islands of similar trees for data sets DA10 (two islands) and LZ12 (six islands grouped into two larger clusters). There is also an indication of island structure for SQ 10, while the posterior tree spaces are more homogeneous for the remaining three data sets (Fig. 5).

Note that these tree spaces are based on minimum SPR distances. Trees that appear close in this space may be more distant, or vice versa, in a space based on the actual probability of moving between them using, say, eSPR or pTBR. Nevertheless, the minimum SPR distances are presumably fairly similar to the true distances of the SPR and TBR moves considered here.

The single-chain test runs on the six data sets revealed striking differences in the performance of tree proposals. Analyses using parsimony-guided tree proposals (pSPR1+pTBR1 or pSPR2+pTBR2) increased in likelihood considerably faster than those using the standard extending proposals (eSPR+eTBR) (Fig. 6). The increase in likelihood was about an order of magnitude faster (note the logarithmic scale of the x-axis). There was no clear difference between the two variants of parsimony-guided proposals.

Individual chains had a tendency to get stuck in suboptimal regions of tree space for a while before moving on toward more likely topologies, as indicated by local plateaus in the likelihood trace plots (Fig. 6). This phenomenon affected all tree proposals but was somewhat more pronounced in runs using standard moves, especially for data sets SQ10 and LZ12. For at least two of the data sets (SQ10 and NP11), the parsimony-guided chains reached higher likelihoods at the end of the test runs than all or most of the standard chains. The single exception was data set AZ12, where one of the 32 parsimony-guided chains remained stuck throughout the run, despite the last of the 16 standard chains reaching the final plateau in log likelihoods one fourth into the run (Fig. 6).

The convergence plots, which compare the topology samples from each of the test runs to the reference distribution using the ASDSF diagnostic, also reveal that the parsimony-guided proposals converged to the posterior much faster than the extending proposals, especially early on in the runs (Fig. 7; note the log scale on...
Test of the ability of the parsimony-guided tree proposals to retrieve a uniform prior over topologies when the first 100, 1000, or 10,000 sites were used to compute the parsimony weights (see legend to Fig. 3).

| Data set | R.20M | R.10M | 2 eSPR : 1 eTBR | 2 pSPR1 : 1 pTBR1 | 2 pSPR2 : 1 pTBR2 |
|---------|-------|-------|-----------------|------------------|------------------|
| SQ20    | 0.004 | 0.007 | 0.097           | 4.56%, 3.78%     | 0.037            | 8.52%, 12.1%     | 0.012            | 8.63%, 12.2%     |
| DA12    | 0.003 | 0.005 | 0.008           | 54.3%, 43.3%     | 0.006            | 48.2%, 43.1%     | 0.005            | 53.1%, 48.4%     |
| LZ12    | 0.004 | 0.006 | 0.009           | 18.0%, 12.7%     | 0.006            | 18.6%, 16.9%     | 0.006            | 18.8%, 18.8%     |
| CL12    | 0.004 | 0.007 | 0.031           | 4.88%, 4.18%     | 0.009            | 4.60%, 4.70%     | 0.037            | 7.14%, 10.0%     |
| AZ12    | 0.020 | 0.041 | 0.108           | 59.4%, 49.5%     | 0.039            | 46.9%, 44.1%     | 0.038            | 52.3%, 49.1%     |
| NP11    | 0.002 | 0.004 | 0.019           | 4.56%, 3.78%     | 0.037            | 8.52%, 12.1%     | 0.012            | 8.63%, 12.2%     |

Notes: We first give the average standard deviation of split frequencies (ASDSF) for the reference runs after 20 and 10 million generations (R.20M and R.10M, respectively). Then we give the ASDSF values and average acceptance proportion ($P_{\text{acpt}}$) for 16 test runs using three different combinations of tree proposals. The reference tree samples after 20 million generations had ASDSF ≤ 0.02 and were considered as ground truth in the detailed studies of the convergence and mixing behavior of tree proposals.
TABLE 5. The percentage of missing data (including gaps), estimated gamma shape parameter (mean and 95\% HPD interval), Sackin index (Mooers and Heard, 1997) measuring the balance of the consensus tree, and estimated size of the 95\% credible set, for each of the six data sets.

| Data set | Missing | Gamma shape ($\gamma$) | Sackin index | 95\% Credible set |
|----------|---------|------------------------|--------------|--------------------|
| SQ10     | 4.1\%   | 0.47 (0.46, 0.49)      | 0.83         | 45604              |
| DA10     | 18.0\%  | 0.46 (0.44, 0.49)      | 0.90         | 45604              |
| LZ12     | 0.05\%  | 0.27 (0.25, 0.29)      | 0.78         | 45604              |
| CL12     | 13.9\%  | 0.96 (0.89, 1.03)      | 1.15         | 45604              |
| AZ12     | 7.2\%   | 0.41 (0.39, 0.42)      | 0.75         | 45604              |
| NP11     | 83.9\%  | 0.78 (0.71, 0.85)      | 0.94         | 45604              |

Note: The Sackin index is normalized under the uniform model (Blum et al., 2006). Larger Sackin index indicates less balanced tree. The true credible sets are likely to be much larger; the run settings constrain the estimates to a maximum value. See text for further discussion.

FIGURE 5. Multidimensional scaling to the SPR distance matrix of 4000 randomly subsampled trees from the posterior sample for each of the six data sets.
A few runs got stuck in local regions of tree space and did not cover the posterior distribution well; these runs generally corresponded to those that did not reach the final plateau in the likelihood trace plots (Fig. 6).

To look specifically at mixing behavior, we analyzed the last 5 million generations (the second half) of the runs. Note that they correspond to only the last 10% of the x-axis in Figure 6 because of the log scale (see also Supplementary Fig. S1 available on Dryad). All of the chains had reached the highest likelihood plateau at this point for LZ12 and CL12, and some extending and most (if not all) of the parsimony-guided chains for the remaining four data sets. We used both all chains (Fig. 8) and chains reaching the highest likelihood plateau (Supplementary Fig. S2 available on Dryad) to display the mixing behavior. The mixing plots show that the parsimony-guided proposals usually covered the set of likely trees significantly faster than the extending proposals, even for samples drawn after the run was assumed to have converged to the posterior. This pattern is particularly clear for the chains that had reached the highest likelihood plateau (Supplementary Fig. S2 available on Dryad).

**Figure 6.** Performance of three different tree proposal combinations—eSPR + eTBR (black), pSPR1 + pTBR1 (blue), and pSPR2 + pTBR2 (red)—on the six data sets (a–f) for which we had good reference samples from the posterior. We show the likelihood trace plots for 16 individual single-chain runs for each proposal combination. Note that all axes are in log scale. The trace plots corresponding to the last 5 million generations are shown in Supplementary Figure S1 available on Dryad.
The mixing plots also reveal that tree spaces that were difficult to sample from using extending proposals (SQ10, DA10, and AZ12), causing individual chains to get stuck for a very long time, were also difficult to sample from using parsimony-guided proposals. In general, however, it appeared that chains using parsimony-guided proposals were less likely to get stuck. The sampling difficulty is only partly correlated with the peakiness of the tree landscape indicated by the MDS plots (Fig. 5). Distinct islands in tree space can apparently cause chains to get stuck (DA10) but there are also tree spaces with distinct islands that are easy to sample from (LZ12) and spaces without obvious island structure that are difficult to sample from (AZ12).

**DISCUSSION**

Fast approximations of the posterior probability of different trees or tree topologies have the potential of making MCMC tree proposals “aware” of the structure of the local tree space around the current tree, allowing them to make smarter proposals with a higher chance of being accepted. The faster the approximation, the larger the region of tree space that a MCMC tree proposal could
 Parsimony scores are interesting in this context because they can be computed so rapidly. Evaluating the likelihood of an alignment of $c$ discrete characters with $s$ states over a tree with $n$ tips has time complexity roughly proportional to $cn^2$. Evaluating the parsimony score can be done in time proportional to $cn$, which is small in comparison. The constants involved in the time complexity equations are also considerably smaller for parsimony implementations, particularly with appropriate optimizations (Ronquist, 1998). Importantly, the parsimony scores of SPR and TBR candidate trees can be obtained in time that is dependent on only $c$ for each candidate evaluated, once a $cn$ order computation of parsimony sets has been completed. The parsimony sets need to be computed only once for the entire tree; after that they can be updated in time only dependent on $c$, under some reasonable assumptions about tree perturbations that are likely to be accepted in a hill-climbing or an MCMC algorithm (Ronquist, 1998). In this study, we used a naive implementation to compute parsimony scores, without employing any of the advanced optimizations described above. Nevertheless, we found that the pSPR and pTBR moves were only slightly slower than eSPR and eTBR. This means that there is considerable room for expansion of the region of tree space evaluated with parsimony scores beyond what we explored in the moves presented here.
The fact that parsimony scores may favor the wrong tree in some cases due to LBA has been widely discussed in the literature (Felsenstein, 1978). Therefore, one might be concerned that parsimony-guided proposals may introduce biases in the MCMC sample of trees. However, our simulations show that the parsimony-guided proposals are implemented correctly and that the proposal ratio adequately corrects for parsimony biases, such as the LBA effect. Even under extreme settings, when numerical inaccuracies start affecting the sampling of the least likely trees (the ones with the lowest parsimony weights but also with the lowest likelihood scores), we detect no oversampling of the LBA tree in relation to the best tree in our simulations.

Even though posterior distributions are sampled correctly, mismatches between the parsimony scores and posterior probabilities are likely to reduce the acceptance proportion and therefore the effectiveness of parsimony-guided proposals. It is possible that some of the variation in the success of parsimony-guided proposals that we observed is explained by differences in the effectiveness of the parsimony scores in approximating posterior probabilities for different data sets. However, such effects were not immediately apparent. One might expect data sets with more rate heterogeneity across sites to be more challenging than data sets with less variation in site rates, because rate heterogeneity would increase the mismatch between parsimony and likelihood scores. However, there were not the least challenging for parsimony-guided proposals included both the two data sets that had the least rate heterogeneity (CL12 and NP11) and the data set with the most pronounced rate variation across sites (LZ12; Table 5). One might also expect data sets with more variation in branch lengths to be more challenging for parsimony-guided proposals. However, there appeared to be no relation between variation in branch lengths (Fig. 9) and the difficulty of sampling the posterior correctly using parsimony-guided proposals. Of course, these conclusions are only based on six data sets, so they remain tentative at best.

In general, the tree spaces that were difficult to sample from using extending proposals were also comparatively more difficult to sample from using parsimony-guided proposals. Surprisingly, the peakness of the tree space did not seem to be the only factor determining the sampling difficulty, as the more difficult data sets included both tree spaces with distinct tree islands (DA10; Fig. 5) and more homogeneous ones (AZ12). A potential explanation for this is that the tree space based on minimum SPR distances is different from the tree space based on the overall probability of moving between trees with high posterior probability using a particular tree proposal. It is the latter distance that actually determines the extent to which there are local peaks in the tree space for that proposal. Unfortunately, there is no method that can visualize large tree spaces based on move probabilities as far as we know. Besides, it is also possible that the 2D projection using MDS may hide cryptic structure in higher-dimensional tree space.

Intuitively, one would expect that parsimony-guided proposals should be particularly helpful in the burn-in phase of an MCMC run, when moving from a poor starting tree to a tree with high posterior probability. It seems less clear whether parsimony-guided proposals would also help chains mix well over the posterior once the best trees have been reached. Our results do show that parsimony-guided proposals can dramatically shorten the burn-in phase of an MCMC run, but they also indicate that parsimony-guided proposals can improve mixing after the chains have converged. Presumably, the ability of the parsimony-guided proposals to see promising peaks beyond the local region in tree space is important enough that it compensates for the mismatches between parsimony scores and posterior probabilities even during the later phases of an MCMC run. However, it should be borne in mind that the mixing conclusions are based on only six data sets, and the pattern was less clear for one of them (DA10), so it is possible that parsimony-guided proposals do not improve mixing on all data sets.

As shown in Table 4, the ASDSF was always lower in the reference runs than in the test runs at 10 million generations, indicating that convergence and mixing were faster when using Metropolis coupling and combining the extending and parsimony-guided tree proposals. A question that arose during the study was whether this improvement in convergence was due largely to Metropolis coupling or to the combination of moves. To answer this question, we repeated some of the reference runs using the same mix of tree proposals, but without Metropolis coupling. The results (Supplementary Table S3 available on Dryad) indicate that the mix of proposals is responsible for a large proportion of the improvement in convergence. This supports the recommendation of using mixes of tree proposals in empirical analyses, which is also suggested by our observation that the relative performance of different tree proposals depends strongly on the data set analyzed. These results also show that it is more computationally efficient to run long single runs than it is to use Metropolis coupling for these six data sets. This conclusion, of course, is not likely to hold for all data sets (see Whidden and Matsen, 2015).

The fact that we used a more realistic evolutionary model than the JC69 evolutionary model suggests that our results should be relevant for many empirical analyses, even though there is no guarantee that this is the case. Topological convergence may become either easier or harder for more realistic and complex evolutionary models, and the relative performance of different types of moves could change both with the model and the data set analyzed.

Although we focused entirely on unrooted trees, our conclusions regarding the pSPR moves should apply also to rooted and clock-constrained trees. In fact, parsimony-guided moves have an interesting property that could be particularly useful when estimating divergence times in clock trees with fossils using total-evidence dating.
Figure 9. Histograms of the median branch lengths in the consensus tree of the reference tree samples for each of the six data sets.

Figure 9. Histograms of the median branch lengths in the consensus tree of the reference tree samples for each of the six data sets.

(also referred to as tip dating or integrative dating) (Ronquist et al., 2012a). In such analyses, fossil taxa tend to have an extremely large proportion of missing characters because they lack sequence data and usually have incomplete coding for morphology as well, while extant taxa can potentially be coded for all characters. A parsimony-guided proposal would then generate a much more diffuse proposal distribution when moving fossil taxa than when moving extant taxa, as would be appropriate.

Despite a large computational budget, our experiments were not able to distinguish clearly between the performance of the two different pSPR and pTBR variants. Clearly, there is a need for further experimentation with tuning parameters of the moves described here, and exploration of other types of approximations of the posterior tree probabilities. We also want to emphasize that, although our results were consistent across all the six data sets for which we were able to obtain reasonably accurate estimates of the posterior, the sample size is still small and it would be highly desirable to extend the experiments to more data sets. In either case, other experimental approaches than the one used here may be needed to make this practical. Although finding such approaches could prove challenging, our results indicate that efforts in this direction might be rewarding.

Supplementary Material

Data available from the Dryad Digital Repository: http://dx.doi.org/10.5061/dryad.98mp657.

Funding

This research was supported by Swedish Research Council (VR) (2014-05901 to F.R.). C.Z. was supported by the 100 Young Talents Program of Chinese Academy of Sciences and the Strategic Priority Research Program of Chinese Academy of Sciences (XDB26000000). Computations were performed using the Swedish National Infrastructure for Computing under projects SNIC 2014/1-323 and SNIC 2015/1-394.

Appendix

Metropolis–Hastings Tree Proposals

All tree proposals described in the article are examples of the Metropolis–Hastings algorithm. It generates successive samples from the posterior distribution
by proposing a new state in parameter space, \( \theta' \), conditional on the current state \( \theta \), according to a proposal distribution that we will denote \( q(\theta'|\theta) \). The new state is then accepted with a probability \( R \), determined by the equation

\[
R = \min\left( 1, \frac{q(\theta|\theta')}{q(\theta'|\theta)} \right)
\]

where the observations (data) are denoted \( y \). The acceptance probability involves the product of three factors—the likelihood ratio \( (f(\theta'|y)/f(y|\theta)) \), the prior ratio \( (q(\theta)/q(\theta'|\theta)) \), and the proposal ratio \( (q(\theta'|\theta)/q(\theta'|\theta')) \), also known as the Hastings ratio.

Note that the proposal ratio is computed as the probability of the backward move, returning to \( \theta \) from \( \theta' \), divided by the probability of the forward move, proposing \( \theta' \) when starting from \( \theta \). Assume we bias the move so that it is more probable to propose \( \theta' \) from \( \theta \) than the other way around. Then the proposal ratio for a move from \( \theta \) to \( \theta' \) will be smaller than one, making it less likely to accept the move. However, if the likelihood of \( \theta' \) is higher than that of \( \theta \), this preference can still pay off. A special case occurs if the preference for \( \theta' \) over \( \theta \) is exactly the same as the ratio of the posterior probabilities of these states (a Gibbs proposal). Then we have the Hastings ratio

\[
\frac{q(\theta|\theta')}{q(\theta'|\theta)} = \frac{f(\theta'|y)}{f(y|\theta)} \cdot \frac{q(\theta)}{q(\theta|\theta')}
\]

This is the reverse of the likelihood ratio times the prior ratio, which results in these terms canceling the proposal ratio. Thus, the acceptance ratio becomes 1.0, which means that the proposal is always accepted, showing that a Gibbs proposal is a special case of the Metropolis–Hastings algorithm.

Computing the posterior probability ratio (the likelihood ratio times the prior ratio) for a set of candidate states is often computationally expensive. The goal we pursue in the current paper is to find proposals based on approximations of the posterior probability ratio that are easy to compute. For good approximations, the proposal ratio should come close to canceling the true posterior probability ratio, thereby increasing the average acceptance probability and improving mixing.

An unrooted phylogenetic tree \( T \) can be described as consisting of a topology \( T \) and a set of branch lengths \( v \). Typically, a phylogenetic model also includes substitution model parameters \( \omega \). Thus, we can write a state in parameter space as \( \hat{\theta} = (T, v, \omega) \). All tree proposals considered here leave \( \omega \) unchanged, but propose a new topology \( \hat{\theta}' \) and (usually) a new set of branch lengths \( v' \).

The acceptance ratio is then obtained as

\[
R = \min\left( 1, \frac{f(\hat{\theta}'|T, v', \omega)}{f(\hat{\theta}|T, v, \omega)} \cdot \frac{q(\hat{\theta}|\hat{\theta}')}{q(\hat{\theta}'|\hat{\theta})} \right)
\]

The computation of the likelihood ratio and the prior ratio in this equation is well-known (e.g., see Felsenstein, 2003). Therefore, to describe a tree proposal, it is sufficient to specify how to generate a proposed state (\( \hat{\theta}' \), \( v' \)) from the current state (\( \hat{\theta} \), \( v \)) and how to compute the associated proposal ratio \( q(\hat{\theta}|\hat{\theta}')/q(\hat{\theta}'|\hat{\theta}) \).

Implementation Details

For pSPR moves, calculating the proposal weight \( \omega_0 \) (Equation 1) involves calculating the parsimony score \( S_i \) of branch \( i \). This score can be obtained easily by first computing the most parsimonious states at the root node of \( A \) in Fig. 1b and at all nodes in subtree BCDE, then computing the number of most parsimonious changes among the three nodes incident to the potential regrafting point (\( x, y, \) and \( z \) in Fig. 1b) using well-known algorithms (e.g., see Felsenstein, 2003). This procedure ensures that \( S_i \) is the parsimony score of the tree after regrafting \( A \) at \( i \) minus the sum of the parsimony scores of the two subtrees A and BCDE. For pTBR moves, \( S_i \) scores are computed by first obtaining the most parsimonious states of all nodes in both subtrees, and then obtaining the number of most parsimonious changes among the four nodes, two in each subtree, adjacent to the proposed reconnection point.

Computationally, we do not obtain the proposal ratio (Equation 2) by directly calculating the \( \omega_0 \) or their sum because this could result in severe numerical problems. Instead, to obtain the ratio associated with the backward or the forward move (the numerator or denominator in Equation 2), we first find the maximum value of \( \omega_0 \) (assuming that \( 0 < p_i < 1 \)) denoted as \( \omega_{\text{max}} = p_0 S_0 \). In pSPR1 or pTBR1, since \( p_i \)'s are all equal to \( e^{-\epsilon} \), this is equivalent to finding the \( \omega \) value for the minimal parsimony score. This may not be the case for pSPR2 or pTBR2, since the \( p_i \)'s vary and \( S_i \) may not be the minimal value among the \( S_j \) values.

After finding \( \omega_{\text{max}} \), we divide each \( \omega_i \) by \( \omega_{\text{max}} \) and convert \( \omega_i / \omega_{\text{max}} \) to

\[
\exp(\log(\omega_i) - \log(\omega_{\text{max}})) = \exp(S_i - S_{\text{max}} - \log(S_{\text{max}})).
\]

Using these transformed weights for the forward move and backward move allows us to obtain good numerical stability in computing the proposal ratio.

To further improve the precision of calculating the sum for \( \epsilon = B_1 \), we adopted the Kahan summation algorithm (Kahan, 1965). Even so, there might still be biases because of numerical inaccuracies in adding extremely unequal numbers. Take pSPR1 for example. Computing the proposal ratio for this move depends on weight terms of the form \( e^{(S_0 - S_{\text{max}})/2} \) by default (\( \epsilon = 0.5 \)); see Equation A4. The maximum value of these terms is 1.0 when \( S_{\text{max}} = S_j \) (the most parsimonious tree) but the value can be extremely small when \( S_{\text{max}} \ll S_j \) (e.g., \( \epsilon = 0.30 \) when \( S_{\text{max}} = 10 \) and \( S_j = 70 \)), making the small terms neglected during the summation (cf. Equation 2). This may affect moves considering both trees with extremely high and extremely low guide weights. The likely effect is undersampling of trees with extremely low guide weights, and distorted balance among trees with extremely low guide weights. In general, trees with
extremely low guide weights also have extremely low likelihood scores. Therefore, the effect of these numerical errors should be negligible when analyzing empirical data.

REFERENCES

Aberer A.J., Kobert K., Stamatakis A. 2014. ExaBayes: massively parallel bayesian tree inference for the whole-genome era. Mol. Biol. Evol. 31:2553–2556.

Alisenti S., Bell H.L., Besnard G., Christin-Paix P.-A., Columbus J.T., Duval M.R., Edwards E.J., Giussani L., Hasenstab-Lehman K., Hilu K.W., Hodkinson T.R., Ingram A.L., Kellogg E.A., Mashayekhi S., Morroni O., Osborne C.P., Salamin N., Schaefer H., Spriugs E., Smith S.A., Zuloga F., Grp C.P. 2012. New grass phylogeny resolves deep evolutionary relationships and discovers C4 origins. New Phytol. 193:304–312.

Blum M.C.B., François O., Janson S. 2006. The mean, variance and limiting distribution of two statistics sensitive to phylogenetic tree balance. Ann. Appl. Probab. 16:2195–2214.

Bouchard-Côté A., Sankaranarayanan S., Jordan M.J. 2012. Phylogenetic inference via sequential Monte Carlo. Syst. Biol. 61:579–593.

Bouckert R., Heled J., Huelsenbeck J.P., Larget B., Drummond A.J. 2014. BEAST 2: a software platform for Bayesian evolutionary analysis. PLoS Comput. Biol. 10:e1003537.

Cardoso D., de Queiroz L.P., Pennington R.T., de Lima H.C., Fonty G. 2010. ExaBayes: massively parallel inference of phylogeny and its impact on evolutionary biology. Syst. Biol. 61:1–11.

Felsenstein J. 1978. Cases in which parsimony or compatibility methods will be positively misleading. Syst. Biol. 27:401–410.

Felsenstein J. 2003. Inferring phylogenies. Sunderland (MA): Sinauer Associates.

Geyer C.J. 1991. Markov chain Monte Carlo maximum likelihood. J. Comput. Graph. Stat. 6:156–173.

Jukes T., Cantor C.R. 1969. Evolution of protein molecules. In: Munro H.N., editor. Mammalian protein metabolism. New York: Academic Press. p. 21–132.

Kahan W. 1965. Pracniques: further remarks on reducing truncation errors. Commun. ACM. 8:40.

Kimura M. 1980. A simple method for estimating evolutionary rates. base substitutions through comparative studies of nucleotide sequences. J. Mol. Evol. 16:111–120.

Lakner C., van der Mark P., Huelsenbeck J.P., Larget B., Ronquist F. 2008. Efficiency of Markov chain Monte Carlo tree proposals in Bayesian phylogenetics. Syst. Biol. 57:86–103.

Larget B. 2013. The estimation of tree posterior probabilities using conditional clade probability distributions. Syst. Biol. 62:501–511.

Larget B., Simon D. 1999. Markov chain Monte Carlo algorithms for the Bayesian analysis of phylogenetic trees. Mol. Biol. Evol. 16:750–759.

Li S., Pearl D.K., Dose H. 2000. Phylogenetic tree construction using Markov chain Monte Carlo. J. Am. Stat. Assoc. 95:508.

Liu J.S. 2004. Monte Carlo strategies in computational science. Springer Series in Statistics. Springer New York. New York.

Li B., Zheng Y., Murphy R.W., Zeng X. 2012. Coalescence patterns of endemic Tibetan species of stream salamanders (Hylobiidae: Batischephrus). Mol. Ecol. 21:3306–3324.

Maas B. and Newton M.A. 1997. Phylogenetic inference for binary data on dendrograms using Markov chain Monte Carlo. J. Comput. Graph. Stat. 6:122–131.

Metropolis N., Rosenbluth A.W., Rosenbluth M.N., Teller A.H., Teller E. 1953. Equation of state calculations by fast computing machines. J. Chem. Phys. 21:1087–1092.

Moores A.O., Heard S.B. 1997. Inferring evolutionary process from phylogenetic tree shape. Q. Rev. Biol. 72:31–54.

Nagy L.G., Pelikovits T., Kovacs G.M., Voigt K., Vargaesegyi C., Papp T. 2011. Where is the unseen fungal diversity hidden? A study of Mortierella reveals a large contribution of reference collections to the identification of fungal environmental sequences. New Phytol. 191:789–794.

Nascimento F.F., dos Reis M., Yang Z. 2012. A biologist’s guide to Bayesian phylogenetic analysis. Nat. Ecol. Evol. 1:1446–1454.

Peskan P.H. 1973. Optimum Monte Carlo sampling using Markov chains. Biometrika. 60:607–612.

Rambaut A., Grassly N.C. 1997. Seq-Gen: an application for the Monte Carlo simulation of DNA sequence evolution along phylogenetic trees. Comput. Appl. Biosci. 13:235–238.

Rannala B., Yang Z. 1996. Probability distribution of molecular evolutionary trees: a new method of phylogenetic inference. Journal of Molecular Evolution. 43:304–311.

Ronquist F. 1998. Fitch-parsimony algorithms for large data sets. Cladistics. 14:387–400.

Ronquist F., Huelsenbeck J.P. 2003. MrBayes 3: Bayesian phylogenetic inference under mixed models. Bioinformatics. 19:1572–1574.

Ronquist F., Huelsenbeck J.P., Britton T. 2004. Bayesian supertrees. In: Phylogenetic supertrees. Dordrecht, Dordrecht: Springer. p. 193–224.

Ronquist F., Klopfstein S., Viellhelm L., Schalmesteiner S., Murray D.L., Råshnyn A.P. 2012a. A total-evidence approach to dating with fossils, applied to the early radiation of the Hymenoptera. Syst. Biol. 61:973–999.

Ronquist F., Teslenko M., van der Mark P., Ayres D.L., Darling A., Hohna S., Larget B., Liu L., Suchard M.A., Huelsenbeck J.P. 2012b. MrBayes 3.2: efficient Bayesian phylogenetic inference and model choice across a large model space. Syst. Biol. 61:539–542.

Savoilainen V., Chase M.W., Hoot S.B., Morton C.M., Soltis D.E., Bayer E., Mau B. and Newton M.A. 1997. Phylogenetic inference for binary data on dendrograms using Markov chain Monte Carlo. J. Comput. Graph. Stat. 6:122–131.

Swofford D.L. 2003. PAUP*. Phylogenetic analysis using parsimony and evolutionary distances. Sunderland (MA): Sinauer Associates.

Takahata Y. 1988. The origin of endemism of Tibetan species of stream salamanders (Hynobiidae: Batrachuperus). Mol. Ecol. 7:189–200.

Zhong B., Rambaut A., Grassly N.C. 1997. Seq-Gen: an application for the Monte Carlo simulation of DNA sequence evolution along phylogenetic trees. Comput. Appl. Biosci. 13:235–238.
Whidden C., Beiko R.G., Zeh N. 2013. Fixed-parameter algorithms for maximum agreement forests. SIAM J. Comput. 42:1431–1466.
Whidden C., Matsen F.A. 2015. Quantifying MCMC exploration of phylogenetic tree space. Syst. Biol. 64:472–491.
Yang Z. 1994. Maximum likelihood phylogenetic estimation from DNA sequences with variable rates over sites: approximate methods. J. Mol. Evol. 39:306–324.
Yang Z. 2014. Molecular evolution: a statistical approach. Oxford, UK: Oxford University Press.
Yang Z., Rannala B. 1997. Bayesian phylogenetic inference using DNA sequences: a Markov Chain Monte Carlo Method. Mol. Biol. Evol. 14:717–724.
Yang Z., Rannala B. 2012. Molecular phylogenetics: principles and practice. Nat. Rev. Genet. 13:303–314.
Zhang C., Rannala B., Yang Z. 2012. Robustness of compound Dirichlet priors for Bayesian inference of branch lengths. Syst. Biol. 61:779–784.