Reconstructing Phylogenies Using Branch-Variable Substitution Models and Unaligned Biomolecular Sequences: A Performance Study and New Resampling Method

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ABSTRACT

In many clades in the Tree of Life, nucleotide substitution rates and base frequencies are hypothesized to have changed as genome evolution unfolded over time. Rigorous testing of this hypothesis relies on accurate phylogenetic reconstruction under suitable models of biomolecular sequence evolution. By far the most common approach for phylogenetic reconstruction is a "two-phase" analysis, where unaligned biomolecular sequence data are first aligned, and the resulting multiple sequence alignment (MSA) is used as input to downstream phylogenetic reconstruction. For a traditional "homogeneous" substitution model that is fixed across a species phylogeny, it has long been established that accurate phylogenetic inference and learning requires accurate upstream multiple sequence alignments. But the same question has not been carefully studied for "heterogeneous" models of substitution processes that can vary across the branches of a phylogeny.

We therefore conducted a comprehensive performance study to quantify the impact of upstream MSA estimation error on downstream phylogenetic inference and learning under branch-variable models of nucleotide substitution. Across model conditions with either 10 or 20 taxa and spanning a range of evolutionary divergence, we find a consistent and significantly positive association between upstream and downstream estimation error. The relationship is robust to the choice of MSA estimation method as well as substitution model mis-specification. We further quantify the relatively large contribution of upstream MSA estimation error to downstream phylogenetic reconstruction quality, compared to other experimental factors. We also conducted an empirical study of flowering monocots. Phylogenetic analyses of orthologous genes in the clade confirm the simulation study findings, and species tree estimation using branch-variable substitution models reveals new insights into sequence evolution heterogeneity. Our findings underscore several key gaps in the state of the art, including the

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need for MSA-aware phylogenetic inference and learning methods under heterogeneous models of sequence evolution.

To this end, we introduce a new computational method, NoHTS ("Non-Homogeneous Tree Support"), to directly assess phylogenetic estimation uncertainty due to MSA estimation error and other factors. The new method uses sequence-aware statistical resampling to place confidence intervals on a phylogeny estimated under a branch-variable substitution model. We demonstrate its superior type I and type II error versus a de facto standard in phylogenetic and phylogenomic studies – the phylogenetic bootstrap method.

CCS CONCEPTS

 Applied computing → Computational genomics; Computational biology; Molecular sequence analysis; Molecular evolution; Computational genomics; Bioinformatics; Population genetics.

KEYWORDS

multiple sequence alignment, maximum likelihood estimation, branchvariable substitution model, phylogenetic tree, simulation study, Poales, phylogenetic support, bootstrap, resampling

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1 INTRODUCTION

Throughout the Tree of Life, interspecific genomic variation in base composition and other features are thought to signify heterotachy and other shifts in biomolecular sequence evolution over time. Important models of this phenomenon include grasses [5], insects [4], and birds [26]. Knowing when and where these compositional biases arise in the evolutionary history of these organisms will help shed light into its functional significance. For example, nucleotide composition changes are hypothesized to result in concomitant downstream changes in transcription and translation (e.g., codon usage bias and cellular distributions of different amino acids) [18].

Rigorous statistical testing of this evolutionary hypothesis relies on statistical models and computational methods to reconstruct phylogenies using biomolecular sequence data, where the latter

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include models of sequence evolution that can vary across the branches of a phylogeny. By far the most widely used approach is a "two-phase" method: (1) a multiple sequence alignment (MSA) is first estimated on the input set of unaligned biomolecular sequences, and (2) a phylogeny is then reconstructed using the estimated MSA as input.

Many methods have been developed for addressing the initial phase of multiple sequence alignment, which is a classical and well-studied computational problem in computational biology and bioinformatics. The problem is also known to be NP-Complete [37]. For this reason, a variety of MSA heuristics have been developed. One such heuristic is progressive multiple sequence alignment, where an input guide tree is used to successively perform pairwise alignment of alignments. Among the most accurate and most popular MSA methods are MAFFT [16], MUSCLE [8], Clustal Omega [34] and its predecessor Clustal W [20], and FSA [2].

The second phase in a two-phase analysis consists of phylogenetic reconstruction using an estimated MSA as input. Statistical inference and learning methods for this task typically model sequence evolution as a branch-homogeneous substitution process (i.e., a substitution process that does not vary across the branches of a phylogeny). More general branch-variable substitution models have been proposed, and phylogenetic software packages for performing statistical inference and learning under these models have also been developed. These software are broadly classified by their statistical optimization criteria. One class consists of maximum likelihood estimation (MLE) methods. PAML [43] is a widely used method in this class, as well as nhPhyML [1]. Other popular phylogenetic MLE methods include limited support for branch-variable substitution models (e.g., RAxML [35] supports a leaf-versus-internal-edge substitution model). Another class of statistical methods utilize Bayesian inference and learning. BEAST [7] is widely used for Bayesian phylogenetic estimation, and it supports relaxed molecular clock models of rate heterogeneity among branches. MrBayes [33] is another option in this class, but it only offers limited support for covarion models [14]. We focus on PAML as a representative state-of-the-art method that is scalable to relatively large datasets.

A variety of factors contribute to phylogenetic reconstruction accuracy. Beyond the question of branch-variable sequence evolution, a large body of studies has demonstrated the central importance of estimated MSA quality in traditional phylogenetic analyses using branch-homogeneous substitution models [21, 22, 25]. But this question has not been well studied for phylogenetic estimation under branch-variable substitution models. The same question arises in the context of statistical estimation problems that are unique to branch-variable rate heterogeneity: these include shift edge inference, rooting phylogenetic trees under non-stationary models, and continuous parameter estimation (i.e. substitution rates, branch lengths, and base frequencies) for more complex substitution models. Another practical consideration is whether methodological guidance from earlier studies is applicable to inference and learning under more complex models.

Critically, new tools are needed to perform data-driven assessment of the relationship between upstream MSA estimation error and downstream phylogenetic estimation error. Such path-breaking tools promise to convert an "unknown unknown" – to paraphrase Donald Rumsfeld's infamous quote – into a quantifiable and surmountable challenge; a sufficient critical mass of evidence, as provided by such tools, can set the stage for further research progress.

In this study, we directly address both gaps. A comprehensive performance study using simulated and empirical datasets is conducted to assess the effect of multiple sequence alignment quality on phylogenetic estimation when evolution is non-homogeneous and non-stationary. We then apply RAWR ("RAndom Walk Resampling") [38], our recently introduced sequence-aware statistical resampling technique, to a new task: confidence interval estimation for phylogenetic tree reconstruction under branch-variable substitution models when unaligned biomolecular sequence data are used as input.

2 MATERIALS AND METHODS

We begin with the following notation and definitions. Let T = (V, E) be a rooted tree with labeled leaves $X \subset V$ and root $\rho \in V$. An unrooted version of a tree T can be obtained by "omitting" the root ρ (i.e., deleting ρ and "connecting" its incident edges). Each edge $e = (u, v) \in E$ where $u, v \in V$ has a length d(e). An edge (u, v) is a leaf edge if either u or v is a leaf, otherwise it is an internal edge. Deleting an edge e from a tree T gives two subtrees $T_1 = (V_1, E_1)$ and $T_2 = (V_2, E_2)$. The vertex sets V_1 and V_2 are disjoint, and $V_1 \cup V_2 = V$. The same can be said for their respective leaf sets, so $\{X_1, X_2\}$ is a bipartition of X. Let this be denoted as $b(e) = \{X_1, X_2\}$.

2.1 Methods under study

Multiple sequence alignment. Our study included a range of the most commonly used and/or most accurate multiple sequence alignment methods. We aligned simulated and empirical datasets using MAFFT [16] version 7.475, MUSCLE [8] version 5.0.1428, Clustal Omega [34] version 1.2.4, Clustal W [20] version 2.1, and FSA [2] version 1.15.9. Each method was run using their respective default settings.

Phylogenetic estimation. We use the General Time Reversible (GTR) model of finite-sites nucleotide substitution for phylogenetic estimation. The GTR model is parameterized by base frequencies $\pi_T, \pi_C, \pi_A, \pi_G$ and substitution rate parameters a, b, c, d, e, f. We use the same conventions as used by [42], where *a* corresponds to $T \leftrightarrow C$, *b* corresponds to $T \leftrightarrow A$, *c* to $T \leftrightarrow G$, *d* to $C \leftrightarrow A$, *e* to $C \leftrightarrow G$, and *f* to $G \leftrightarrow A$. *f* is canonically fixed to 1 and the remaining free rate parameters are specified relative to *f*. The substitution rate matrix *Q* is then defined as follows:

$$Q = \begin{bmatrix} \cdot & a\pi_C & b\pi_A & c\pi_G \\ a\pi_T & \cdot & d\pi_A & e\pi_G \\ b\pi_T & d\pi_C & \cdot & f\pi_G \\ c\pi_T & e\pi_C & f\pi_A & \cdot \end{bmatrix}$$

with the diagonals set to $Q_{ii} = -\sum_{i \neq j} Q_{ij}$. The transition probability matrix is given by $P(t) = \exp(-Qt)$ and is used to calculate model likelihood for a phylogenetic tree. Typically in phylogenetic estimation using Markov models of substitution, the rate matrix is assumed to be constant over the whole tree. We refer to models under this assumption as homogeneous or "no-shift" models.

More general models are needed to account for substitution process variation across a phylogeny. To this end, the homogeneity assumption can be relaxed by associating each edge e with a set of parameters $\theta(e)$ that defines the rate matrix for that edge. We use a GTR model for the branch models. The traditional homogeneous model is the case where $\theta(e)$ is fixed, i.e. $\theta(e_i) = \theta(e_i)$ for all $e_i, e_i \in E$. For heterogeneous models, we considered two different classes that we refer to as "single-shift" and "all-shift". For all-shift, $\theta(e)$ is independent for each edge. For single-shift, there are exactly two sets of parameters, θ_{shift} and $\theta_{\text{background}}$ and some restrictions on which edges they apply to. There is a shift edge, $e_{\text{shift}} \in E$, and all edges descending from it all have $\theta(e) = \theta_{\text{shift}}$. Any remaining edges are $\theta_{\rm background}.$ In nonhomogeneous models, rooting can impact likelihood values since these models are not time-reversible, so rooted trees are used. We note that the no-shift, single-shift, and all-shift models are nested in order of increasing model complexity. The all-shift model approaches the no common mechanism model, under which MLE is known to be statistically inconsistent [36].

RAxML [35] version 8.2.12 was used to perform maximum likelihood estimation under a homogeneous GTR model. PAML [43] version 4.9j was used to perform maximum likelihood estimation under fixed tree topologies using a branch model. PAML supports maximum likelihood fixed-tree-topology optimization of continuous parameters under nonhomogeneous substitution models, but does not support full tree search under nonhomogeneous models. We therefore implemented custom software to perform tree search and thereby maximize likelihood under the all-shift substitution model, where PAML was used to evaluate model likelihood of a tree topology during search. MLE under the single-shift model was a special case of all-shift model-based search, where the shift and background parameters are estimated and all tree edges in a given topology are evaluated to find a shift edge that maximizes single-shift model likelihood.

NoHTS, a new phylogenetic support estimation method. RAWR (or "RAndom Walk Resampling") [38] is a recently introduced method for sequence-aware statistical resampling of biomolecular sequence data. RAWR resampling takes the form of a random walk conducted directly on biomolecular sequences (see Supplementary Methods for pseudocode). The first application of RAWR resampling was to the task of phylogenetic support estimation under homogeneous substitution models [38]. In this study, we apply RAWR to a new task: phylogenetic support estimation under branch-variable substitution models (Figure 1). The new application naturally generalizes the original application since the branch-variable substitution models under study are a superset of traditional homogeneous substitution models. We refer to the resulting phylogenetic support estimation method as NoHTS (or "Non-Homogeneous Tree Support"). Each resampled replicate dataset consists of a set of unaligned sequences and is used to perform MSA and phylogenetic re-estimation. The resulting set of re-estimated trees is then used to calculate support for the annotation estimated tree, where the support for an edge in the annotation tree is the proportion of re-estimated trees that also display that edge. We note that other support estimation tasks are possible (e.g., support calculations for substitution process shifts along a phylogeny, tree rooting, and others), although we leave these to be explored in future research.

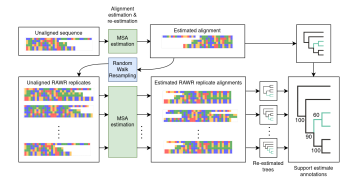


Figure 1: Graphical overview of NoHTS (or "Non-Homogeneous Tree Support"), the new phylogenetic support estimation method. RAWR [38] is used to perform sequence-aware resampling of an input set of biomolecular sequences. Whereas Wang et al. [38] originally applied RAWR resampling to phylogenetic support estimation under homogeneous substitution models, our new application of RAWR resampling focuses on phylogenetic support estimation under branchvariable substitution models. In the new application, each RAWR replicate consists of a set of unaligned sequences, and is used to perform MSA and phylogenetic tree re-estimation where the latter utilizes branch-variable substitution modelbased MLE. The resulting re-estimates are used to calculate branch support for the annotation tree (i.e., the tree estimated on the original input dataset): the support value for a branch in the annotation tree is the proportion of re-estimated trees that also display that branch.

For phylogenetic support estimation, we compared NoHTS versus the phylogenetic bootstrap method [11] on the 10-taxon simulation study model conditions. NoHTS utilized RAWR resampling with reversal probability parameter $\gamma = 0.1$ and either 10 or 100 resampled replicates. The phylogenetic bootstrap analyses were run using 100 bootstrap replicates. All replicates were aligned using MAFFT, and alignment re-estimation was performed with MAFFT as well. Phylogenetic estimation and re-estimation were performed using single-shift maximum likelihood estimation.

Performance assessments. We use Robinson-Foulds distance [32] to assess topological difference between trees. Let $S(T) = \{b(e) | e \in E, e \text{ is an internal edge}\}$. The Robinson-Foulds distance between two unrooted trees *T* and *T'* is the symmetric difference of S(T) and S(T'). For identifying root placement, we say two trees *T* and *T'* have identical roots, ρ and ρ' respectively, if the leaf sets of the subtrees induced by deleting the respective root nodes are identical.

We take the L1 norm of the relative errors for substitution model parameters to assess model parameter estimation performance in the simulation study. For the base frequencies, this would be $\sum_{i \in ACGT} \left| \frac{\pi_i - \hat{\pi}_i}{\pi_i} \right|$.

To assess how well the shift subtree is being predicted in the single-shift model, we use the size of the maximum agreement subtree (MAST) between the true and estimated shift subtrees. The MAST problem is to find a subtree given a set of trees \mathscr{T} with the largest subset of leaves that also agrees with all the trees in \mathscr{T} .

For evaluating alignment quality, we use sum-of-pairs false positive and false negative proportions, denoted SP-FP and SP-FN respectively. SP-FP is calculated as the proportion of homologous nucleotide pairs in the estimated alignment and not in the true alignment. SP-FN is defined vice versa.

We performed linear regression analyses to quantify the relationship between upstream MSA estimation error and downstream phylogenetic estimation error. Python was used with the package scikit-learn [29] to perform the linear regression analyses.

For evaluating phylogenetic support estimation methods (i.e., NoHTS and the phylogenetic bootstrap method), we used precision-recall (PR) and receiver operating characteristic (ROC) curves and the area under curves (PR-AUC and ROC-AUC, respectively). Confusion matrix entries used to construct the PR and ROC curves were calculated using the same procedures as in [38].

To quantify the relative importance of each factor in the simulation study, we utilized the random forest approach in the study of Lanier and Knowles [19]. Likewise, we performed random forest analysis in R using the package randomForest [6]. We fit 1000 regression trees with the following factors: model condition (indel probability and model tree height), MLE method, alignment type (true or estimated), and number of taxa. Relative importance of a factor was measured as the increase in mean squared error (MSE) when excluding that factor over the highest increase in MSE.

2.2 Simulated datasets

Supplementary Figure S1 provides a graphical overview of our study's simulation procedures.

Model tree generation. Model trees were sampled using INDELible [12] under a random birth-death process. Non-ultrametricity was introduced using the procedure described in [27] with deviation factor c = 2. First, a rooted model tree is generated using INDELible under a birth-death process. For every branch, sample x from a uniform distribution $x \sim U(-\ln(2), \ln(2))$ and scale the branch length by x. The tree height is then rescaled so that the maximum root-to-tip distance is the height specified by the model condition. Finally, a subtree containing as close to half of the leaves is selected to evolve under the shift substitution model.

Simulating sequence evolution. Our simulations utilized model conditions that were based on the study of Wang et al. [38] (Table 2). Following the rationale of Wang et al. [38], the model condition parameter settings reflect a range of evolutionary divergence that are often encountered in modern-day phylogenetic systematics and related research topics. To simulate sequence evolution on the model trees, INDELible was used to perform finite-sites simulations under a GTR-based branch model and the indel model of [12]. Based on Nabholz et al. [26]'s report of GC content variation in the avian phylogeny, the GTR model parameters were empirically estimated using single-copy orthologs from [15] for the subset of species (Calypte anna, Alligator mississippiensis, Melopsittacus undulatus, Corvus brachyrhynchos, and Manacus vitellinus) included in Nabholz et al. [26]'s study. To estimate these parameters, we aligned the single-copy orthologs using MAFFT with the default settings. MLE under the single-shift model was used to estimate parameters on each individual aligned sequence. Then, we looked at the two sets of estimated substitution rates, and we observed that the ratio

Table 1: Simulation study: GTR model parameters used for the single-shift substitution model in our simulations. The single-shift substitution model is comprised of two sets of model parameters: one corresponding to a "background" substitution process, and the other to a "shift" substitution process. Settings for each set of base frequency parameters π_T , π_C , π_A , π_G are listed, followed by the settings for each set of substitution rate parameters a, b, c, d, e, f. (See Methods section for details.)

Parameter	π_T	π_C	π_A	π_G	$C \leftrightarrow T$	A⇔T	$G \leftrightarrow T$	A⇔C	C⇔G	A⇔G
Shift	0.216	0.237	0.317	0.230	5.847	3.186	1.214	3.437	1.307	1.0
Background	0.183	0.226	0.058	0.534	1.505	0.367	0.141	0.412	0.094	1.0

Table 2: Simulation study model conditions. The 10-taxon model conditions are named 10.A through 10.E in order of generally increasing evolutionary divergence, and the 20taxon model conditions are named 20.A through 20.E similarly. As noted in the Methods section, model condition parameter settings were based on the study of Wang et al. [38] and reflect a range of evolutionary divergence. The model tree height and indel model parameter are listed for each model condition. Each model condition consists of settings for these two parameters and the single-shift substitution model parameters. (See Methods section for details.)

Model	Number	Tree	Indel
condition	of taxa	height	probability
10.A	10	0.47	0.13
10.B	10	0.7	0.1
10.C	10	1.2	0.06
10.D	10	2	0.031
10.E	10	4.4	0.013
20.A	20	0.47	0.13
20.B	20	0.7	0.1
20.C	20	1.2	0.06
20.D	20	2	0.031
20.E	20	4.4	0.013

between them was bimodal. We chose GTR model parameters based on the estimated parameters in the lower rate mode (Table 1). The simulation outputs consisted of a true multiple sequence alignment, the corresponding set of unaligned sequences, a model tree with branch lengths, and the true substitution model instance. Table 3 lists summary statistics for true and estimated MSAs.

Experimental replication. For each model condition, the simulation procedure was repeated to obtain 30 experimental replicates. Results are reported on average (along with standard errors) across all experimental replicates in each model condition.

2.3 Empirical datasets

Flowering monocot dataset. The distribution of GC content in the Poales, an order of flowering monocots, is bimodal [5]. This pattern is notably strong in rice. We applied nonhomogeneous substitution model-based phylogenetic tree estimation to a set of 7 taxa from the Poales – Oryza sativa japonica [28], Sorghum bicolor [23], *Carex cristatella, C. scoparia, Juncus effusus, Juncus inflexus* [30], and *Ananas comosus* [24] – and one additional taxon from the order Zingiberales – *Musa balbisiana* [39]. We identified 1900 single-copy orthologs using OrthoFinder [10] with default settings. Average sequence length across all taxa and single-copy orthologs was 1377.

The Impact of MSA Error on Branch-Variable Phylogenetic MLE

Table 3: Simulation study: summary statistics for ground truth and estimated MSAs. MAFFT, Muscle, Clustal W, Clustal Omega, and FSA were used to estimated MSAs on each simulation study dataset. For each model condition, each MSA method's average alignment SP-FN and SP-FP error ("SP-FN" and "SP-FP", respectively) are reported across all replicate datasets (n = 30). Alignment length ("length"), average normalized Hamming distance ("ANHD") for aligned sequence pairs, and proportion of MSA cells that consist of indels ("Gappiness") are reported for each method as an average across all replicates in a model condition (n = 30).

Statistic	Alignment		Model condition									
Statistic	0	10.A	10.B	10.C	10.D	10.E	20.A	20.B	20.C	20.D	20.E	
	MAFFT	0.531	0.667	0.751	0.831	0.890	0.392	0.539	0.775	0.873	0.948	
	MUSCLE	0.526	0.637	0.716	0.788	0.851	0.355	0.480	0.702	0.812	0.908	
SP-FN	CLUSTALW	0.715	0.765	0.806	0.855	0.892	0.630	0.736	0.846	0.890	0.943	
	CLUSTALO	0.710	0.769	0.813	0.854	0.884	0.640	0.728	0.843	0.889	0.937	
	FSA	0.680	0.751	0.820	0.886	0.927	0.550	0.706	0.864	0.923	0.961	
	MAFFT	0.526	0.663	0.750	0.833	0.893	0.364	0.519	0.770	0.872	0.949	
	MUSCLE	0.513	0.633	0.715	0.792	0.858	0.331	0.465	0.700	0.814	0.913	
SP-FP	CLUSTALW	0.702	0.759	0.803	0.856	0.896	0.595	0.715	0.843	0.891	0.945	
	CLUSTALO	0.667	0.741	0.795	0.847	0.886	0.566	0.679	0.827	0.884	0.939	
	FSA	0.394	0.501	0.599	0.695	0.763	0.181	0.313	0.535	0.529	0.782	
	TRUE	2123.8	2315.8	2315.8	2313.2	2063.0	2410.3	2585.1	2895.8	2696.2	2723.6	
	MAFFT	1478.6	1477.1	1484.5	1461.8	1529.2	1643.7	1670.7	1683.0	1691.4	1804.3	
Length	MUSCLE	1518.5	1570.1	1573.1	1561.9	1590.5	1790.2	1863.0	1907.7	1890.0	1979.5	
Lengui	CLUSTALW	1191.4	1186.0	1170.6	1143.0	1146.5	1278.6	1261.2	1227.8	1181.6	1162.8	
	CLUSTALO	1247.1	1248.7	1237.6	1222.9	1234.0	1306.3	1318.2	1303.2	1287.4	1283.8	
	FSA	3609.3	4471.2	4992.4	5729.6	6196.2	4394.5	5959.0	8231.8	10177.9	11566.5	
	TRUE	0.306	0.364	0.364	0.465	0.649	0.301	0.374	0.484	0.581	0.667	
	MAFFT	0.389	0.435	0.484	0.528	0.569	0.368	0.433	0.516	0.569	0.608	
ANHD	MUSCLE	0.413	0.449	0.499	0.542	0.589	0.380	0.445	0.530	0.585	0.630	
ANHD	CLUSTALW	0.466	0.496	0.537	0.573	0.614	0.449	0.509	0.573	0.615	0.652	
	CLUSTALO	0.484	0.504	0.541	0.565	0.602	0.471	0.518	0.570	0.602	0.637	
	FSA	0.280	0.319	0.377	0.420	0.477	0.289	0.349	0.429	0.486	0.525	
	TRUE	0.528	0.564	0.564	0.566	0.510	0.581	0.606	0.649	0.622	0.629	
	MAFFT	0.326	0.321	0.328	0.310	0.342	0.390	0.394	0.400	0.403	0.444	
Gappiness	MUSCLE	0.342	0.361	0.364	0.353	0.366	0.439	0.456	0.470	0.466	0.493	
Gappiness	CLUSTALW	0.165	0.155	0.148	0.119	0.124	0.216	0.198	0.178	0.148	0.140	
	CLUSTALO	0.202	0.198	0.194	0.177	0.187	0.233	0.232	0.226	0.218	0.221	
	FSA	0.715	0.770	0.795	0.820	0.836	0.763	0.819	0.874	0.899	0.913	

We aligned the sequences individually using MAFFT, MUSCLE, Clustal Omega, Clustal W, and FSA using the same settings as those used in the simulation study. We performed phylogenetic estimation under a single-shift model for every individual gene.

2.4 Data availability statement

Data and scripts used are available under an open copyleft license at https://gitlab.msu.edu/liulab/nonhomogeneous-substitution-model-study-data-scripts.

3 RESULTS

3.1 Simulation study

MSA estimation error and topological error of single-shift MLE. We focus first on phylogenetic reconstruction with no model misspecification. On the least divergent 10.A model condition, MLE under the single-shift model returned the lowest average topological error when the true MSA was provided as input, followed by estimated MSAs excluding FSA (i.e., ClustalOmega, MAFFT, MUS-CLE, and ClustalW), and FSA-estimated MSAs returned the worst accuracy overall. As evolutionary divergence increased across the 10-taxon model conditions – with 10.A being the least divergent and 10.E the most divergent – topological error returned by the different methods also increased to differing extents. The smallest effect was seen on true alignments; a relatively stronger effect seen on estimated alignments, with FSA exhibiting the greatest effect among all MSA estimation methods. Across the 10-taxon model conditions,

MLE(TrueAln), MLE(ClustalOmega), MLE(MAFFT), MLE(Muscle), MLE(ClustalW, and MLE(FSA) returned average topological error of 0.116, 0.232, 0.245, 0.277, 0.253, and 0.386, respectively; the corresponding estimated MSAs had average SP-FN (SP-FP) error of 0.778 (0.816), 0.706 (0.743), 0.677 (0.723), 0.776 (0.816), and 0.782 (0.321), respectively.

All MSA estimation methods returned some degree of alignment error, and comparison of phylogenetic MLE using estimated MSAs vs. true MSAs demonstrates the clear impact of upstream estimation error on downstream phylogenetic reconstruction. However, relative comparisons among MSA methods were not a perfect predictor of resulting topological error, at least at the coarse granularity of per-model-condition averages. Muscle and MAFFT were generally among the more accurate MSA methods in terms of SP-FN and SP-FP error – with Muscle outperforming MAFFT slightly, FSA consistently returned lower SP-FP error compared to all other methods, and ClustalW-estimated alignments were generally least accurate. We note that our findings are consistent with other related studies involving traditional homogeneous model-based MLE [21, 22].

We performed linear regression analyses to examine the relationship between upstream MSA error and downstream topological error at a finer granularity. Per-replicate scatterplots and fitted linear regression models are shown in Figure 3. The correlation between alignment SP-FN error and topological error was observed to be positive and statistically significant across all model conditions. A similar outcome was observed between alignment SP-FP error and topological error, except on 2 of the least divergent 10-taxon model conditions (Supplementary Figure S4 in the SOM Appendix). Correlation coefficients tended to increase as evolutionary divergence increased, with the strongest associations observed on the most divergent model conditions.

Similar outcomes were observed on the 20-taxon model conditions. Topological error on the 20-taxon model conditions were somewhat higher than on the 10-taxon model conditions, as expected due to the combinatorially larger solution spaces required by the former compared to the latter. Across the 20-taxon model conditions, MLE(TrueAln), MLE(ClustalOmega), MLE(MAFFT), MLE(Muscle), MLE(ClustalW, and MLE(FSA) returned average topological error of 0.121, 0.329, 0.352, 0.310, 0.325, and 0.408, respectively; the corresponding MSAs had average SP-FN (SP-FP) error of 0.792 (0.814), 0.696 (0.712), 0.643 (0.675), 0.791 (0.822), and 0.778 (0.255), respectively. (Per-method regression analyses also yielded consistent results, as shown in Supplementary Figures S11 through S15.)

MSA estimation error and topological error: role of model mis-specification. Across the different model conditions in our study, single-shift MLE using a given MSA as input (i.e., the true MSA or one of the estimated MSAs) returned the best (or among the best) average topological error, compared to the zero-shift and all-shift MLE methods (Figure 2). In fact, on the most divergent 10.E and 20.E model conditions, zero-shift model-based MLE on the true MSA returned topological error that was comparable to estimated MSA-based MLE under any of the substitution models. For MLE on the true, MUSCLE, Clustal Omega, ClustalW, MAFFT, and FSA alignments, the respective average topological error for single-shift phylogenetic estimation was 0.094, 0.258, 0.261, 0.269, 0.289, and 0.380; the respective average topological error for all-shift phylogenetic estimation was 0.120,

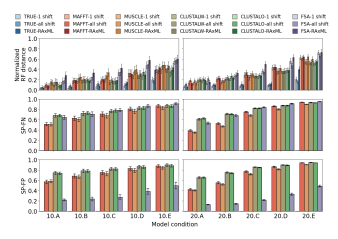


Figure 2: Simulation study: estimated MSA and tree error returned by each method on each model condition. Phylogenetic trees were estimated using the true and estimated MSAs as input, where the latter were estimated using a range of different MSA methods: MAFFT, MUSCLE, Clustal W, Clustal Omega, and FSA. Phylogenetic trees were estimated using MLE under one of three nested models: no-shift, single-shift, and all-shift. Topological error was measured using normalized Robinson-Foulds distance between a model tree and estimated tree. An estimated MSA was compared against the true MSA based on alignment SP-FN and SP-FP error. For each model condition and method, average and standard error of each performance assessment are reported (n = 30).

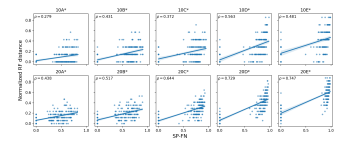


Figure 3: Simulation study: the relationship between upstream MSA estimation error and downstream phylogenetic estimation error. The former was assessed based on sum-of-pairs false negative rate, and the latter was assessed based on normalized Robinson-Foulds distance. We performed linear regression analyses to quantify the relationship between the two assessments on each model condition. The fitted linear regression model is shown as a blue line along with a confidence interval shown in a light blue shade. MSA and tree estimation methods are as described in Figure 2. Results are aggregated for all alignment types (n = 180).

0.312, 0.292, 0.290, 0.306, and 0.386; finally, the respective average topological error for zero-shift phylogenetic estimation was 0.149, 0.333, 0.286, 0.301, 0.297, and 0.445. On both true and estimated MSAs, the difference in average topological error between MLE

Table 4: Simulation study: AUC for the precision-recall and ROC curves of NoHTS versus bootstrap support estimation methods on the 10-taxon model conditions. For the bootstrap method, 100 resampled replicates were utilized for each analysis. For NoHTS, either 100 or 10 resampled replicates were utilized for each analysis.

		PR-AUC		ROC-AUC				
Model	Bootstrap	No	HTS	Bootstrap	NoHTS			
condition	100 reps	10 reps	100 reps	100 reps	10 reps	100 reps		
10.A	0.988	0.992	0.996	0.875	0.916	0.957		
10.B	0.972	0.984	0.988	0.876	0.926	0.944		
10.C	0.945	0.978	0.983	0.748	0.899	0.926		
10.D	0.889	0.978	0.977	0.749	0.950	0.947		
10.E	0.874	0.945	0.957	0.791	0.910	0.922		

under single-shift vs. all-shift models was smaller than single-shift vs. zero-shift comparison, with the exception of Clustal Omega on the 10-taxon model conditions and MAFFT and MUSCLE on the 20-taxon model conditions. We hypothesize that extraneous model parameters and possible overfitting is not appreciably degrading topological accuracy of phylogenetic reconstruction. Finally, the impact of model mis-specification on resulting topological error was less apparent on estimated MSAs than on the true MSA.

In our random forest analysis, relative importance of the evolutionary divergence, alignment type, number of taxa, and maximum likelihood method used were 1.0, 0.48, 0.09, and 0.03 for predicting normalized RF-distance.

Performance evaluation of NoHTS versus bootstrap support estimates. For phylogenetic support estimation, NoHTS with both 100 and 10 resampled replicates consistently yielded an improvement over the phylogenetic bootstrap method based on both PR-AUC and ROC-AUC assessments (Table 4). Model conditions with increased evolutionary divergence, and consequently increased MSA and phylogenetic tree estimation error, trended towards lower PR-AUC and ROC-AUC. However, both methods were not equally affected by evolutionary divergence. Based on AUC assessments, the performance advantage returned by NoHTS over the phylogenetic bootstrap method grew larger as model conditions became more divergent, suggesting that NoHTS is especially well suited to sequence inputs that pose a greater challenge in terms of MSA and phylogenetic estimation. This pattern is also visible in the curves (Supplementary Figure S16). Additional simulation experiments indicated that NoHTS was largely robust to the effects of varying biomolecular sequence length (Supplementary Figure S8).

MSA estimation error and phylogenetic tree rooting. Overall, singleshift-model-based phylogenetic estimation on true MSAs returned the most accurate rooting (Table 5). Among estimated MSAs, MAFFT and Muscle analyses under the single-shift model returned the most accurate rooting, followed by Clustal Omega and Clustal W; FSA returned the least accurate rooting. As evolutionary divergence increased across the 10-taxon model conditions, rooting error generally increased as well; smaller increases were seen on true MSAs and comparatively larger increases were seen on estimated MSAs, with the largest increases seen on FSA alignments. A similar effect concerning evolutionary divergence was seen on the 20-taxon model conditions. The Impact of MSA Error on Branch-Variable Phylogenetic MLE

Table 5: Simulation study: proportion of correct root placements by MSA, MLE method, and model condition. Results are reported as an average across all replicates in each model condition (n = 30).

		Correct root rate								
Model condition	MLE method	TRUE	MAFFT	MUSCLE	CLUSTALW	CLUSTALO	FSA			
10A	single shift	56.7%	23.3%	43.3%	26.7%	26.7%	23.3%			
10A	all shift	23.3%	20.0%	20.0%	23.3%	6.7%	20.0%			
10B	single shift	60.0%	16.7%	23.3%	13.3%	6.7%	20.0%			
10D	all shift	30.0%	6.7%	10.0%	3.3%	6.7%	0.0%			
10C	single shift	60.0%	16.7%	16.7%	10.0%	10.0%	16.7%			
100	all shift	30.0%	13.3%	16.7%	3.3%	0.0%	6.7%			
10D	single shift	40.0%	20.0%	26.7%	6.7%	16.7%	6.7%			
1012	all shift	23.3%	3.3%	16.7%	0.0%	6.7%	3.3%			
10E	single shift	30.0%	10.0%	10.0%	6.7%	10.0%	3.3%			
IUE	all shift	6.7%	3.3%	3.3%	3.3%	6.7%	0.0%			
20A	single shift	33.3%	20.0%	30.0%	13.3%	16.7%	36.7%			
20A	all shift	31.0%	6.9%	7.1%	6.7%	16.7%	17.2%			
20B	single shift	37.9%	17.2%	20.7%	6.9%	10.3%	24.1%			
200	all shift	24.1%	13.8%	10.3%	0.0%	7.1%	10.3%			
20C	single shift	46.7%	20.0%	13.3%	13.3%	3.3%	0.0%			
200	all shift	13.3%	6.7%	0.0%	6.7%	0.0%	3.3%			
20D	single shift	50.0%	13.3%	6.7%	13.3%	6.7%	3.3%			
2015	all shift	10.3%	3.3%	0.0%	3.3%	0.0%	0.0%			
20E	single shift	46.7%	16.7%	3.3%	6.7%	6.7%	0.0%			
20E	all shift	6.7%	0.0%	3.3%	3.3%	0.0%	0.0%			

Model mis-specification also served to increase rooting error. Still, the relative comparison among different MSAs was largely similar under the all-shift model, as compared to the single-shift model.

For each model and method on each simulation condition, rooting error is generally higher than overall topological error. This finding suggests that rooting is more difficult than estimation of other aspects of topological reconstruction, which is consistent with experimental outcomes for traditional approaches to phylogenetic rooting (e.g., outgroup rooting, midpoint rooting, etc.) and phylogenetic estimation under homogeneous substitution models (cf. section 2.2.6 in [41]).

MSA estimation error and continuous parameter estimation. Looking beyond topological error, we next examined the consequences of upstream MSA estimation error on downstream estimation of substitution rates. Results for MLE under the single-shift model are shown in Figure 4. Single-shift MLE on the true MSA consistently returned the lowest average error across the 10- and 20-taxon model conditions, compared to single-shift MLE on estimated MSAs. We found that single-shift MLE on FSA-estimated MSAs was among the most accurate relative to the other estimated MSA-based methods. We also found that the other estimated MSA-based methods (excluding single-shift MLE on FSA-estimated MSAs) often returned relatively high substitution rate estimation error on background edges and lower substitution rate estimation error on shift edges, or vice versa, on each model condition. Model mis-specification also played a role. Compared to single-shift MLE, zero-shift MLE returned higher substitution rate estimation error (Figure 4), and all-shift MLE had higher error still (Supplementary Figure S4). The differences were stark - amounting to as much as an order of magnitude or more when comparing single-shift vs. all-shift MLE. Relatively high rate estimation errors were seen under mis-specified zero-shift and all-shift models, and the role of MSA quality became more difficult to discern.

Results for base frequency estimation are reported for each model condition and method in Figure 5. Similar to substitution rate estimation outcomes, single-shift MLE returned the most accurate base

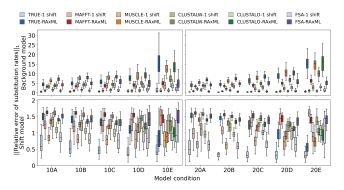


Figure 4: Simulation study: substitution rate estimation error returned by the methods under study. Substitution rate estimation error is measured by L1-norm of the relative errors for rate parameter estimates. Results for all replicates in a model condition are reported as a box-and-whisker plot (n = 30).

frequency estimates on true MSAs, compared to estimated MSAs. Single-shift MLE on FSA MSAs was typically least accurate, and single-shift MLE on the other estimated MSAs were intermediate in accuracy. The error difference between neighboring rankings became more pronounced as evolutionary divergence increased, whereas the different methods returned generally comparable base frequency estimation error on the least divergent model conditions. Also mirroring the substitution rate estimation outcomes, model mis-specification tended to inflate base frequency estimation error. Compared to single-shift MLE, zero- and all-shift MLE returned higher error by as much as several factors. Finally, all-shift MLE consistently returned higher error variance compared to zero- and single-shift MLE.

We also assessed branch length estimation error in our experiments. Leaf branch lengths were consistently over-estimated across our study, regardless of substitution model and input MSA (Supplementary Figure S5 in SOM Appendix). A more comprehensive examination of both leaf and internal branch lengths would be more revealing; while the branch-score metric used was proposed for this purpose [17], assessing internal branch lengths in isolation from other factors - particularly topological error - remains a challenge. MSA error and shift edge estimation. For all single-shift methods, we evaluated shift clade accuracy based on maximal MAST size. Shift clade estimation was most accurate using MLE on the true MSA (Figure 6). Shift clade estimation using estimated MSAs was less accurate, with FSA MSAs resulting in comparable or lower accuracy versus the other MSA methods. Greater evolutionary divergence served to enhance these relative rankings, with the greatest differences between methods occurring on the most divergent 10and 20-taxon model conditions.

3.2 Empirical study

Gene trees were reconstructed on each locus using the MSA and phylogenetic MLE methods under study. Per-locus topological disagreement among the different methods averaged $\sim 5\%$ or so based on traditional R-F distance and $\sim 20\%$ for the rooted version of

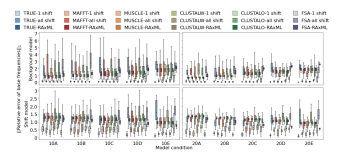


Figure 5: Simulation study: base frequency estimation error returned by the methods under study. Base frequency estimation error is measured by the L1 norm of the relative errors for base frequency parameter estimates. Results for all replicates in a model condition are reported as a box-and-whisker plot (n = 30).

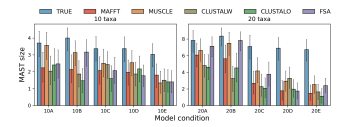


Figure 6: Simulation study: shift edge assessment of singleshift MLE. Shift edge accuracy is assessed based on the size of the MAST of the predicted shift subtree and the true subtree. Average and standard error bars are shown for all replicates in each model condition (n = 30).

R-F distance. On average across loci, the largest topological disagreements were observed among phylogenies reconstructed using ClustalW-estimated MSAs versus any other estimated MSA; all other pairwise comparisons returned relatively less topological disagreement (Table 6).

For continuous parameter estimates, the median absolute difference of estimated base frequencies under the background vs. shift models was in agreement across different single-shift model-based analyses and loci; observed differences were at most 0.10, 0.12, 0.11, and 0.10, respectively (Supplementary Figure S18). A similar outcome was observed for estimated substitution rates, with observed differences of at most 0.41. On the other hand, a relatively large amount of variation around the median was observed across loci for these estimates.

4 DISCUSSION

Simulation study. We observed a consistent impact of upstream MSA error on topological error of downstream phylogenetic MLE under variable-across-phylogeny substitution models. Regression analyses consistently returned positive and significant correlation between upstream and downstream estimation error across all of the model conditions in our simulation study. Upstream MSA error was also observed to have major consequences for continuous

Table 6: Empirical study: topological disagreement among gene trees estimated using different MSA methods and singleshift MLE. For each gene, topological discordance was calculated based on pairwise Robinson-Foulds distance between estimated gene trees returned by different methods. Average ("Avg") and standard error ("SE") are reported across all genes (n = 1377).

	Muscle		Clustal Omega		Clust	al W	FSA	
nRF	Avg	SE	Avg	SE	Avg	SE	Avg	SE
MAFFT	0.044	0.088	0.052	0.097	0.059	0.106	0.041	0.087
Muscle			0.053	0.100	0.058	0.108	0.044	0.090
Clustal Omega					0.059	0.105	0.052	0.095
Clustal W							0.061	0.103

parameter estimation. The outcomes observed for discrete phylogenetic relationships were qualitatively different from those of continuous parameter estimates, which suggests that the latter is more difficult to reconstruct versus the former. Our random forest analysis attributed MSA type to have the second highest relative contribution to both topological error and branch length error, after evolutionary divergence.

We found that NoHTS support estimates consistently performed better than traditional bootstrap support estimates, even with as few as 10 resampled replicates. We attribute this performance advantage to a key difference between the two methods: NoHTS support estimation accounts for both MSA and tree estimation uncertainty, while bootstrap support only accounts for tree estimation uncertainty. The former is possible due to RAWR's ability to preserve sequential ordering information of resampled sites, which is necessary for meaningful MSA re-estimation. The experimental outcome in the NoHTS experiment reinforces our study finding concerning the impact of upstream MSA error on downstream phylogenetic reconstruction, and new computational methods like NoHTS will allow direct quantification of these effects.

Phylogenetic estimation using a branch model matching the number of shifts during sequence evolution returned the best estimation accuracy, as expected. Furthermore, model mis-specification effects were attenuated when estimated MSAs were used as input rather than true MSAs. We attribute the difference to the higher topological error observed on the former versus the latter, which may swamp effects of other methodological factors like model mis-specification. Interestingly, phylogenetic inference under the all-shift model returned topological error that was comparable to the no-shift model for topology estimation, except on the 20.E model condition, where it performs slightly better than the no-shift model. Overall, the impact of MSA error on downstream topological error was qualitatively larger than that of substitution model mis-specification.

Generally speaking, evolutionary divergence served to amplify upstream and downstream estimation error and their association. Increasing dataset size had similar effects, which we anticipate will become even more pronounced on datasets with hundreds or thousands of taxa.

Empirical study. Different MSA methods with varying MSA estimation error resulted in observable differences among downstream The Impact of MSA Error on Branch-Variable Phylogenetic MLE

single-shift MLE phylogenetic estimates. The genome-wide distributions of substitution rate estimates were also variable between alignment methods. We caution that direct comparison between the simulation study and empirical genomic sequence analyses are complicated by differences in data type: the former involves single-locus data and all loci were simulated i.i.d., whereas the latter utilized multi-locus data where different loci may well evolved under more complex and non-i.i.d. evolutionary processes (e.g., genetic recombination). A new generation of species-tree-aware phylogenomic inference methods can help better account for complex evolution of multi-locus biomolecular sequences [3, 9], but we anticipate that the influence of upstream MSA estimation error will become even greater under more complex evolutionary models.

5 CONCLUSIONS

Our performance study assessed the impact of upstream MSA estimation error on downstream phylogenetic MLE under branchheterogeneous substitution models. Throughout the simulation study, MSA estimation error was consistently and significantly associated with downstream topological error for single-shift MLE. Greater correlations were observed as evolutionary divergence increased across model conditions. We introduced NoHTS, a random walk resampling method that assesses phylogenetic support for branch-variable model-based estimation using unaligned biomolecular sequence inputs. For all simulated model conditions, we found that NoHTS support estimation performed better for phylogenetic support estimation, compared to the phylogenetic bootstrap method. Estimation of continuous parameters, tree rooting, and shift edge were also impacted by MSA estimation error. Additional MLE experiments with under- and over-parameterized models indicate that our study findings are robust to model mis-specification. An empirical study of the order Poales revealed findings consistent with the simulation study experiments.

Several recommendations follow from our study's findings. First, NoHTS can and should be used to directly quantify the effects of upstream MSA estimation error on downstream phylogenetic MLE under branch-variable substitution models. We also recommend that NoHTS-estimated phylogenetic confidence intervals be reported alongside any phylogenies that are reconstructed using heterogeneous model-based MLE. Second, our study reinforces a through-line in computational phylogenetics: there is a great need for alignment-aware phylogenetic inference and learning - both under homogeneous [21, 22, 25] and branch-heterogeneous substitution models. Ideally, an MSA and phylogenetic tree would be co-estimated under the same evolutionary model that generated sequence observations. Such co-estimation methods exist for homogeneous models [22, 25, 31] but not for branch-heterogeneous models. In the interim, mitigation via best practices when reconstructing species trees offers a stop-gap workaround. We recommend that species trees be reconstructed using multi-locus data and the latest statistical methods for phylogenomic inference and learning, such as ASTRAL [44]. We caution that this mitigation measure is no silver bullet. More deliberate decision making is needed during upstream phylogenetic study and analysis design [3, 9]. For example, denser taxon sampling may help ameliorate long branch attraction [13], but our experiments suggest that the impact of upstream

MSA error on downstream phylogenetic inference becomes more pronounced as both dataset sizes and divergence increase.

We conclude with thoughts on future work. Our study examined the relationship between upstream MSA estimation error and downstream phylogenetic reconstruction where exactly one substitution process shift occurred along a phylogeny. Even stronger effects are anticipated where multiple evolutionary shifts occur along a phylogeny, as is expected to be the case where larger and more divergent clades are sampled within Tree of Life. This hypothesis merits future study with an expanded set of multiple-shift simulations. Finally, a fundamental issue is model mis-specification due to the use of substitution-only model-based analysis of aligned biomolecular sequence data. This simplifying assumption is pervasive throughout phylogenetics and phylogenomics. Phylogenetic MLE under traditional homogeneous substitution models is known to be statistically inconsistent where sequences evolved under insertion and deletion processes [40]; a similar theoretical limitation is also expected for heterogeneous substitution models. As noted above, statistical co-estimation methods are needed to reconstruct MSAs and phylogenetic trees under a variable-across-phylogeny model of substitutions, insertions, and deletions.

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