

Prospects for inferring very large phylogenies by using the neighbor-joining method

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Current efforts to reconstruct the tree of life and histories of multigene families demand the inference of phylogenies consisting of thousands of gene sequences. However, for such large data sets even a moderate exploration of the tree space needed to identify the optimal tree is virtually impossible. For these cases the neighbor-joining (NJ) method is frequently used because of its demonstrated accuracy for smaller data sets and its computational speed. As data sets grow, however, the fraction of the tree space examined by the NJ algorithm becomes minuscule. Here, we report the results of our computer simulation for examining the accuracy of NJ trees for inferring very large phylogenies. First we present a likelihood method for the simultaneous estimation of all pairwise distances by using biologically realistic models of nucleotide substitution. Use of this method corrects up to 60% of NJ tree errors. Our simulation results show that the accuracy of NJ trees decline only by $\approx 5\%$ when the number of sequences used increases from 32 to 4,096 (128 times) even in the presence of extensive variation in the evolutionary rate among lineages or significant biases in the nucleotide composition and transition/transversion ratio. Our results encourage the use of complex models of nucleotide substitution for estimating evolutionary distances and hint at bright prospects for the application of the NJ and related methods in inferring large phylogenies.

phylogenetics | molecular evolution | distance estimation | tree of life | maximum likelihood

Inference of phylogenetic trees is becoming increasingly important in the study of molecular evolution and functional genomics. However, with the enormous increase in the size of data sets for orthologous genes from diverse species and homologous sequences from multigene families, the probability of finding the optimal tree(s) diminishes rapidly with an astronomical increase in the number of possible topologies to be examined (Fig. 1A) (1, 4). Even a moderate exploration of topological (tree) space is not practical because of the enormous amount of computational time required. These circumstances have led to the extensive use of the NJ method (2), which quickly generates a final tree for large phylogenies under the principle of minimum evolution. This method is especially useful when the number of sequences to be analyzed is in the order of hundreds or thousands (e.g., 5–7). Furthermore, the accuracy of NJ trees is similar to other more time-consuming methods for relatively small data sets (<200 sequences) (1, 4, 8–13).

The NJ method constructs trees by clustering neighboring sequences in a stepwise manner. In each step of sequence clustering, it minimizes the sum of branch lengths (2) and thus examines multiple topologies. For large data sets, however, NJ examines only a minuscule fraction of the total number of possible topologies. For instance, it will examine all three unrooted trees in the case of four sequences but only 10^{10} of $>10^{13,867}$ possible trees when 4,000 sequences are used (Fig. 1B). The effect of this property on the accuracy of NJ trees has been unclear. Therefore, we have done computer simulations to study

the accuracy of NJ trees when tens to thousands of sequences are used.

NJ is known to be statistically consistent in the sense that, if correct pairwise distances with no statistical errors are used, it reconstructs the true tree (2). In practice, however, estimates of all distances are subject to statistical errors, so it may produce erroneous trees. At present, all distances are estimated independently for each pair of sequences [independent estimation (IE) method] either by analytical formulas (1, 14, 15) or by likelihood methods (4, 15). The standard errors of the estimates obtained in this way are rather high unless very long sequences are used. However, these standard errors can be reduced considerably if all distances for a given set of aligned sequences are estimated simultaneously [simultaneous estimation (SE) method]. Recently, we developed an SE method based on the maximum likelihood principle and found that the use of this method substantially improves the accuracy of NJ trees. In this article, we first present this SE method and then discuss the accuracy of NJ trees obtained for large and small phylogenies.

Methods

Simultaneously Estimating All Pairwise Distances. Pairwise distances used for constructing NJ trees are currently estimated by the IE method for a variety of mathematical models to incorporate varying degrees of complexity of nucleotide or amino acid substitution (reviewed in refs. 1, 4, 14, 15). The estimates obtained by the IE method are expected to have larger standard errors than those obtained by the SE method. These larger errors are partly because the parameters, such as the transition/transversion rate ratio in the Kimura model (16), are estimated independently for each pair of sequences in the IE method, whereas they are estimated by considering all pairs of sequences simultaneously in the SE method. There is obviously a greater advantage of using the SE method for a sophisticated model of substitution than for a simple model, because in the former model we have to estimate a larger number of parameters. Furthermore, the accuracy of parameter estimates obtained by the SE method is expected to increase as the number of sequences increases.

In the following we consider the evolutionary distance based on the TN model (3), which is one of the most sophisticated models of nucleotide substitution. For this model, the evolutionary distance (d) is given by

$$d = 4(g_{AG}a_1 + g_{TC}a_2 + g_{RY}b), \quad [1]$$

where a_1 , a_2 , and b are the numbers of transitional substitutions between purines (a_1), transitional substitutions between pyrimidines (a_2), and transversional substitutions (b) per site, respec-

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Abbreviations: NJ, neighbor-joining; SE, simultaneous estimation; IE, independent estimation; TN, Tamura–Nei.

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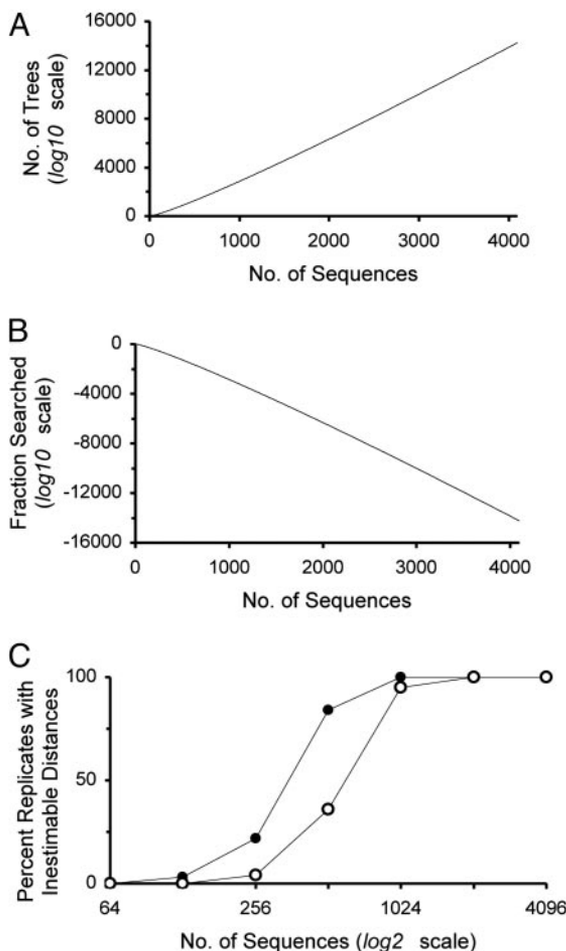


Fig. 1. (A) Total number of possible bifurcating trees for different number of sequences. Computed by equation 5.1 of ref. 1. (B) Fraction of all topologies that are examined by the neighbor-joining (NJ) (2) method in producing a final tree. For a given number of sequences (m), the number of topologies explored by the NJ algorithm can be given by $[m(m^2 - 1)/6] - 7$. This formula was derived from the observation that at each step of sequence clustering the NJ method examines $m_i(m_i - 1)/2$ trees for $m_i \geq 5$ (assuming that each sequence pairing in the NJ algorithm examines a distinct tree), where m_i is the number of sequences at step i . For $m_i = 4$, NJ examines all three possible trees. Because the number of sequences decreases by one in each step of sequence clustering, the total number of topologies examined is given by $\sum_{m_i=5}^m m_i(m_i - 1)/2 + 3 = m(m^2 - 1)/6 - 7$. (C) Proportion of data sets in which the original Tamura-Nei (TN) (3) distance was not applicable for one or more pairwise comparisons for type B model trees (filled symbols) and type C model trees (open symbols) in Fig. 2. The expected maximum distance was 0.47 for 32-sequence trees and 0.61 for 4,096-sequence trees. The simulation procedure was as follows. For a given model tree, a data set of extant nucleotide sequences of $n = 1,000$ was generated by using the TN model with $k_1 = k_2 = 20$, $g_A = g_T = 0.4$ and $g_C = g_G = 0.1$. For each model tree, 100 data sets were generated, and the proportion of data sets in which unestimable distances occurred was computed.

tively, and g_A , g_T , g_C , and g_G each represent the frequencies of nucleotides A, T, C, and G. For the distance (d_{ij}) between sequences i and j , Eq. 1 can be rewritten as follows.

$$d_{ij} = 4(g_A g_G k_1 + g_T g_C k_2 + g_R g_Y) b_{ij}, \quad [2]$$

where $k_1 = a_{1ij}/b_{ij}$ and $k_2 = a_{2ij}/b_{ij}$. In Eq. 2, the transition/transversion rate ratios (k_1 and k_2) are independent of evolutionary time and are the same for all pairs of sequences, whereas b_{ij} depends on evolutionary time, varying with sequence pair i and j . When there are m sequences, the total number of b_{ij} 's is $m(m - 1)/2$. To estimate d_{ij} in Eq. 2, we need to know the

estimates of k_1 , k_2 , and b_{ij} . The maximum likelihood estimates of k_1 , k_2 , and b_{ij} in Eq. 2 can be obtained by maximizing the following log likelihood function (IE method):

$$L_{ij} = \hat{P}_{1ij} \ln(P_{1ij}) + \hat{P}_{2ij} \ln(P_{2ij}) + \hat{Q}_{ij} \ln(Q_{ij}) + (1 - \hat{P}_{1ij} - \hat{P}_{2ij} - \hat{Q}_{ij}) \ln(1 - P_{1ij} - P_{2ij} - Q_{ij}), \quad [3]$$

where \hat{P}_{1ij} , \hat{P}_{2ij} , and \hat{Q}_{ij} are the observed proportions of nucleotide sites showing transitional differences between purines and between pyrimidines and sites showing transversional differences when sequences i and j are compared, respectively. P_{1ij} , P_{2ij} , and Q_{ij} are the theoretical values of \hat{P}_{1ij} , \hat{P}_{2ij} , and \hat{Q}_{ij} , respectively, and are given by

$$P_{1ij} = \frac{2g_A g_G}{g_R} \{g_R - \exp[-2(g_R k_1 + g_Y) b_{ij}] + g_Y \exp(-2b_{ij})\}, \quad [4]$$

$$P_{2ij} = \frac{2g_T g_C}{g_Y} \{g_Y - \exp[-2(g_Y k_2 + g_R) b_{ij}] + g_R \exp(-2b_{ij})\}, \quad [5]$$

$$Q_{ij} = 2g_R g_Y [1 - \exp(-2b_{ij})], \quad [6]$$

where $g_R = g_A + g_G$ and $g_Y = g_C + g_T$.

However, because the parameters k_1 and k_2 are shared by the log likelihood functions for all pairs of i and j , they should be estimated by maximizing the sum of all likelihood functions (SL), which is

$$SL = \sum_i \sum_{j < i} L_{ij}. \quad [7]$$

Theoretically, this is not a likelihood function, because d_{ij} 's are not necessarily independent. However, by using the method of Taylor's expansion (as in ref. 17), one can show that approximately unbiased estimates of k_1 and k_2 can be obtained by maximizing SL . We therefore suggest the following procedure for estimating k_1 , k_2 , and b_{ij} 's. We first compute the initial estimates of k_1 , k_2 , and b_{ij} 's by using uncorrected estimates, $k_1 = (\hat{P}_{1ij}/g_A g_T)/(Q_{ij}/g_R g_Y)$, $k_2 = (\hat{P}_{2ij}/g_T g_C)/(Q_{ij}/g_R g_Y)$, and $b_{ij} = Q_{ij}/(4g_R g_Y)$. This method is computationally efficient and gives estimates with smaller standard errors. We then compute the averages of estimates of k_1 and k_2 separately and use them for obtaining improved estimates of b_{ij} 's by maximizing L_{ij} in Eq. 3. We can now obtain improved estimates of k_1 and k_2 by maximizing SL in Eq. 7. These estimates are then used for further improvement of the estimates of b_{ij} 's by Eq. 3. The last two processes are repeated until stable estimates of k_1 , k_2 , and b_{ij} 's are obtained. The final estimates are asymptotically unbiased, although they may not be maximum likelihood estimates.

The SE method above can be used for many other substitution models. For example, the Hasegawa-Kishino-Yano (HKY) model (18) is a special case of the TN model, in which k_1 and k_2 are assumed to be the same. Therefore, the HKY distance for sequences i and j can be estimated by using Eq. 3 under the assumption of $k_1 = k_2 = k$. Similarly, the Tamura (19) and the Kimura (16) models are a special case of the TN model (see ref. 1). In the case of the Tamura model the $G + C$ content ($\theta = g_G + g_C$) instead of nucleotide frequencies is considered with a single k parameter. In the Kimura model all nucleotide frequencies are assumed to be equal (0.25) with a single k . The SE method can also be used when the substitution rate varies with site, following the gamma or other distribution (e.g., refs. 20 and 21).

In Eqs. 4–6, the use of average nucleotide frequencies for the entire set of sequences is recommended because of the smaller sampling errors. However, when the distance methods that relax the assumption of the equality of nucleotide frequencies among lineages (heterogeneity of substitution pattern over the phylogeny) are used, the sequence-specific nucleotide frequencies should be used for each comparison (22). Note that these methods do not take into account the variation of transition/transversion ratio among lineages and are not guaranteed to generate asymptotically unbiased estimates of evolutionary distances.

The SE approach easily produces estimates of the transition/transversion rate ratios (k_1 and k_2) and pairwise distances (d_{ij}) for all sequence pairs simultaneously. Computation of the variances of these estimates by analytical formulas is somewhat complicated, but they can be obtained by the bootstrap method with site resampling (1, 23).

Methods of Computer Simulation. In our study of the accuracy of NJ trees obtained by the IE and SE methods of distance estimation, we used three different sets of model trees. The first model tree consisted of 66 sequences (Fig. 2A), of which the relative branch lengths were derived from the mammalian DNA sequence data (figure 1 in ref. 24). For this model tree, we considered 448 different sets of evolutionary parameters (substitution parameters and sequence lengths), in which the number of nucleotides per sequence (n) varied from 147 to 9,359, the evolutionary rate varied 10 times, the G + C content (θ) varied from 0.3 to 0.9, and the transition/transversion rate ratio ($k_1 = k_2 = k$) varied from 2.1 to 26.6 (see ref. 25). Using this model tree and the set of evolutionary parameters with the TN model, we evaluated the relative performance of the SE and IE methods in estimating evolutionary parameters and inferring phylogenies by the NJ method.

The second and third sets of model trees were used to compare the accuracy of NJ trees of different sizes. The second set was based on two predefined model trees in Fig. 2, where the tree in B represents a case of constant rate evolution for 32 sequences and the tree in C represents a varying rate case with 32 sequences. Multiple copies of these trees, connected at the roots, generated increasingly larger model trees consisting of 32 to 4,096 sequences (see also ref. 10). The third set of model trees was generated by using an agglomerative algorithm. In this algorithm we combined a given set of sequences and randomly selected pairs or groups of sequences for making different model trees in a stepwise manner. This process was continued until the required number of sequences was clustered. Model trees in Fig. 2D and E are two examples of such randomly generated trees. For model trees in Fig. 2B and D, the exterior branch lengths (0.2 substitutions per site) were 20 times longer than the interior branch lengths (0.01 substitutions per site). For model trees in Fig. 2C and E, the exterior branches were either long (0.2) or short (0.01). For the type of model trees in Fig. 2E, the exterior branches were assigned a long or short length randomly. The number of nucleotides per sequence (n) used in the second and third data sets was 1,000.

For the second and third sets of model trees, nucleotide substitution was simulated by using the TN model with two sets of biologically realistic values of substitution parameters: (i) $k_1 = k_2 = 4$ and $g_A = g_T = g_C = g_G = 0.25$ to simulate nuclear gene evolution and (ii) $k_1 = k_2 = 20$, $g_A = g_T = 0.40$, and $g_C = g_G = 0.10$ to simulate animal mitochondrial DNA gene evolution.

Using the standard simulation procedure (e.g., refs. 1 and 10), we generated simulated sequence data for each model tree with a set of nucleotide substitution parameters and sequence length (n). A NJ tree was then constructed and compared with the true tree. The accuracy of the NJ tree was measured by the percentage of phylogenetic clades correctly inferred (P_C). This was obtained by $P_C = 100 [1 - d_T/(2m - 6)]$, where d_T is the

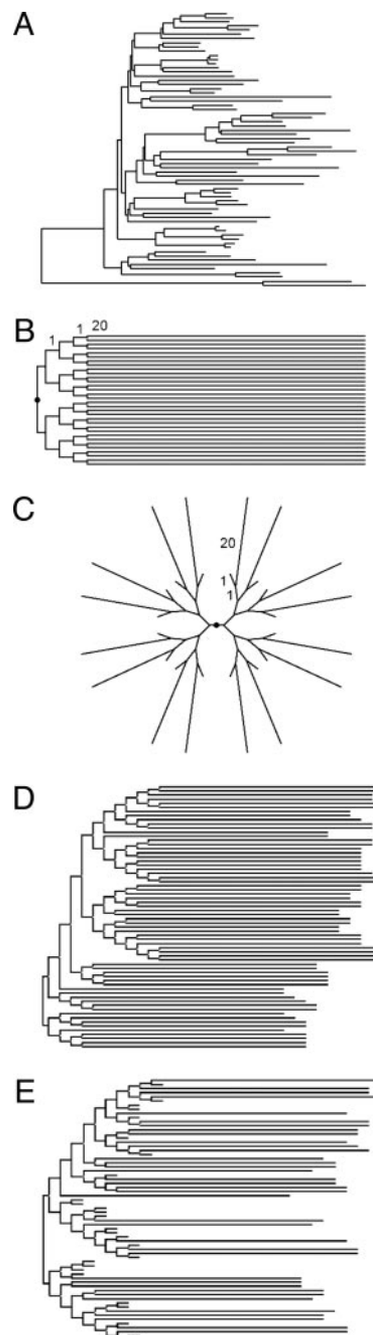


Fig. 2. Some of the model trees used in computer simulations for examining the accuracy of NJ trees. (A) A 66-sequence model tree based on the eutherian phylogeny (see ref. 24). Branches are drawn with relative lengths. (B and C) Trees with constant (model tree B) and variable (model tree C) evolutionary rates that were used to generate increasingly larger trees by connecting their copies at the roots (marked by filled circle) (see also ref. 10). (D and E) Two examples of randomly generated model trees, with equal (tree D) and unequal (tree E) exterior branch lengths. For all model trees (except in A), the expected interior branch lengths were assumed to be the same (see text for details).

topological distance between the inferred and model trees and m is the number of sequences used (1, 26, 27). P_C is 100% when all clades are correctly inferred and is 0% when none of the clades is correctly identified. For each model tree (model tree in Fig. 2A, B, or C) for a given number of sequences, 100 replicates of simulated sequence data were generated for the same topol-

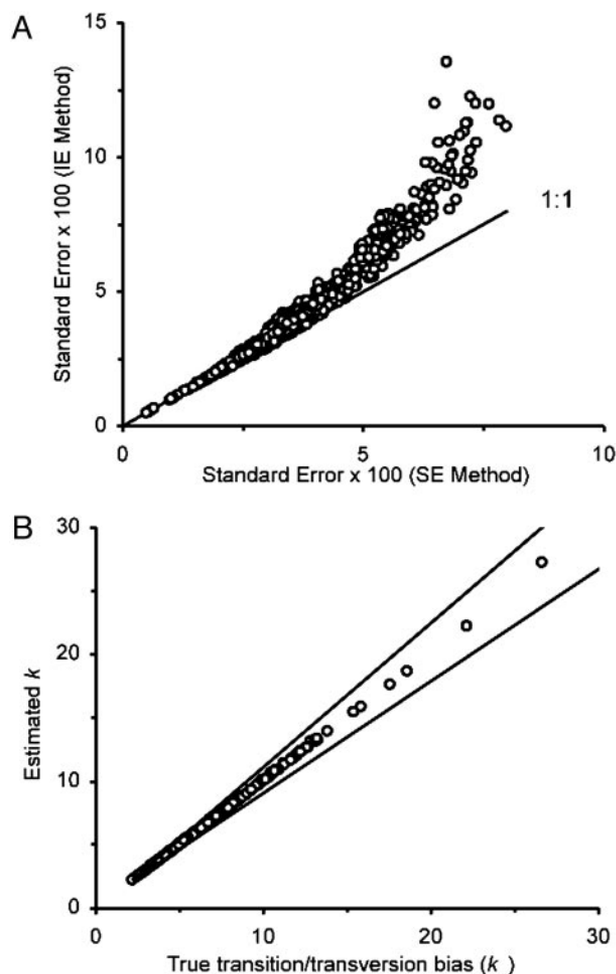


Fig. 3. Standard errors and transition/transversion rate ratios obtained by the SE method. (A) Relationships between the standard errors of evolutionary distances obtained by the IE and SE methods for a data set of 66 computer-generated sequences according to the model tree in Fig. 2A. The TN model was used for generating sequence data with the transition/transversion rate ratios k_1 and $k_2 = 4.4$, G + C content (θ) = 0.61, and sequence length $n = 1,066$ nucleotides. There are $(66 \times 65)/2 = 2,145$ data points in this figure. (B) Relationships between the estimated and true values of k ($k = [k_1 + k_2]/2$) for 448 different patterns of nucleotide substitutions (see text). For each pattern of nucleotide substitution, the average value (circles) and the 95% confidence limits (lines) of the estimate of k were obtained from 100 data sets generated by computer simulation with the model tree in Fig. 2A.

ogy, whereas in the case of randomly generated trees (trees in Fig. 2D and E), a new model tree was generated in each replicate simulation.

To make the P_C values comparable among model trees of different sizes (except Fig. 2A), the expected interior branch lengths were assumed to be the same (0.01 substitutions per site) for all topologies. This approach is different from that used in most previous studies, in which the maximum depths of trees or the maximum evolutionary distances were assumed to be the same (e.g., refs. 11 and 28). The latter approach makes the interior branch lengths shorter in large phylogenies than in small phylogenies and, therefore, the comparison of P_C values among trees of different sizes becomes improper.

Results

Performance of the SE Method. Using the model tree in Fig. 2A, we first compared the standard errors of distance estimates obtained by the IE and SE methods. Fig. 3A shows that the standard

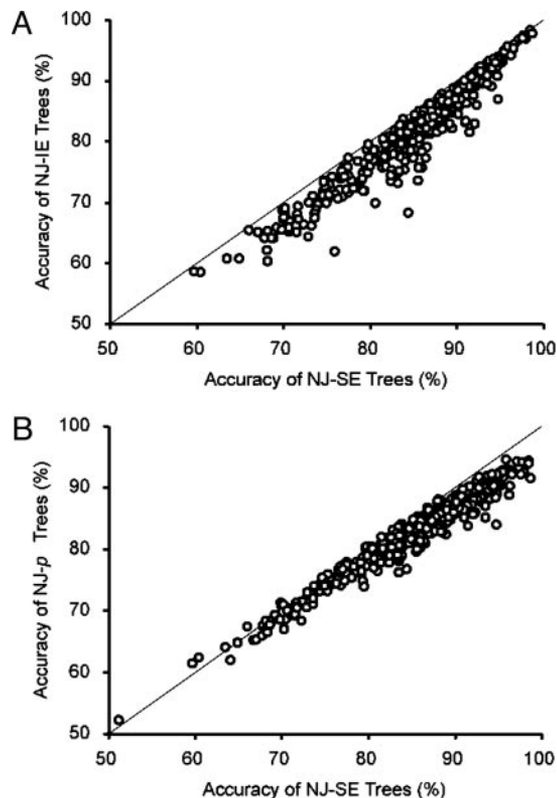


Fig. 4. Accuracy of NJ-SE trees. (A) Accuracies (P_C) of NJ trees when the SE and IE methods with the TN model were used for the 66-sequence model tree in Fig. 2A (see text). (B) Comparison of P_C values for NJ-SE trees with those for NJ trees obtained with p -distance (NJ- p). For each simulation condition, average P_C values from 100 replicate simulations are plotted.

errors for the SE method are always smaller than those for the IE method and that the extent of reduction of the standard errors for the SE method increases as the standard errors for the IE method increases. In this case we used the TN model with $k_1 = k_2 = 4.4$, $\theta = 0.61$, and $n = 1,066$. Essentially the same results were obtained for each of the 448 simulation conditions examined (data not shown). Fig. 3B shows the accuracy of the estimates of k obtained by the SE method. In each of the 448 simulation conditions, the mean estimate of k was close to the true value, and the 95% confident interval was narrow. These results indicate that the SE method is effective in obtaining reliable estimates of k without knowing the topology of the tree.

Fig. 4A shows the P_C values of NJ trees obtained by the SE and IE methods for each of the 448 cases. Each data point in this figure represents the average P_C from 100 replicate simulations for each case. The P_C values of NJ trees obtained by the SE method (NJ-SE) are always higher than those of NJ trees obtained by the IE method (NJ-IE). On average, 19% of all erroneously identified phylogenetic clades of NJ-IE trees were corrected in NJ-SE trees. Large improvements were observed when evolutionary distances were large or when the base composition or the transition/transversion biases were high. In these cases $>50\%$ of erroneous clades of NJ-IE trees were corrected in NJ-SE trees. Essentially the same results were obtained for other model trees in Fig. 2.

Previously we reported that the accuracy of NJ trees is often higher when p distances (proportions of different nucleotide sites) are used than when IE distance estimates for the correct substitution models are used (1, 11, 29, 30). This occurred mainly because p distances have smaller standard errors than distances estimated by the IE method. The same results were observed in

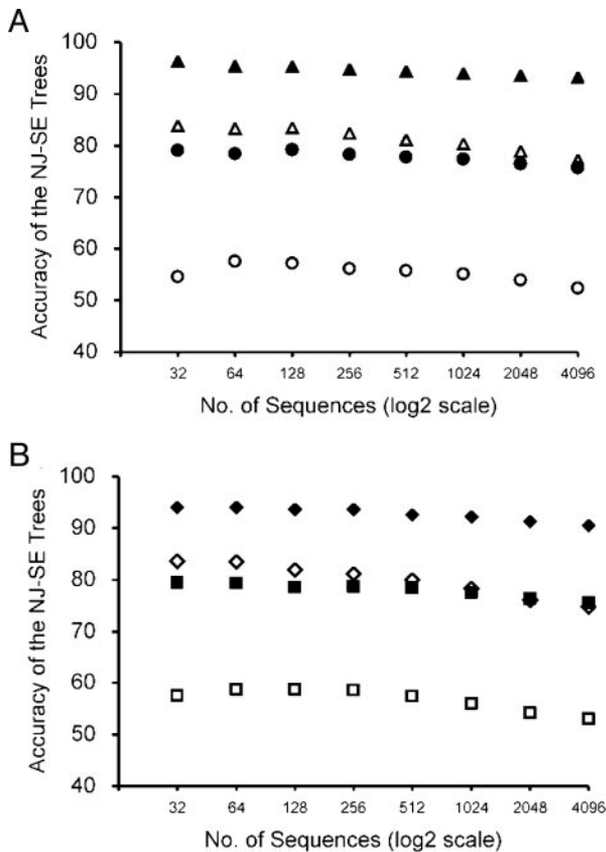


Fig. 5. Accuracies (P_C) of NJ-SE trees with increasing numbers of sequences when the TN model was used. (A) Type B and type C model trees in Fig. 2 were used. For nuclear gene evolution, $k_1 = k_2 = 4$, $g_A = g_T = g_C = g_G = 0.25$, and $n = 1,000$ were used for both type B trees (filled circles) and type C trees (filled triangles). For mitochondrial gene evolution, $k_1 = k_2 = 20$, $g_A = g_T = 0.40$, $g_C = g_G = 0.10$, and $n = 1,000$ were used for both type B trees (open circles) and type C trees (open triangles). (B) Type D and type E model trees in Fig. 2 were used. For nuclear gene evolution, $k_1 = k_2 = 4$, $g_A = g_T = g_C = g_G = 0.25$, and $n = 1,000$ were used for both type D trees (filled squares) and type E trees (filled diamonds). For mitochondrial gene evolution, $k_1 = k_2 = 20$, $g_A = g_T = 0.40$, $g_C = g_G = 0.10$, and $n = 1,000$ were used for both type D trees (open squares) and type E trees (open diamonds).

the present simulation; P_C was higher in NJ- p trees than in NJ-IE trees for 60.4% of the simulation conditions when the TN model was used. However, NJ-SE trees had a higher P_C value than NJ- p trees in 92.9% of the cases (Fig. 4B). Therefore, the SE method considerably improved the accuracy of NJ trees when more realistic models were used. The same results were obtained for other tree topologies given in Fig. 2. For this reason, we consider only NJ-SE trees in the following comparison of P_C values among trees of different sizes.

Accuracy of NJ Trees with Increasing Number of Sequences. One might expect P_C to decline significantly as the number of sequences used (m) increases, because the number of possible topologies rapidly increases with increasing m . To study this problem, we examined the P_C value for different numbers of sequences (32, 64, 128, 256, 512, 1,024, 2,048, and 4,096 sequences) using the model trees in Fig. 2. When type B model trees (constant-rate) with nuclear gene evolution were considered, P_C was 79% for $m = 32$ (Fig. 5A, filled circles). When m was increased to 256, P_C declined only by 0.8%. Even for $m = 4,096$ (128-times increase), P_C was lower than that for $m = 32$ only by 3.4%. These results indicate that the accuracy of NJ-SE trees does not decline significantly even when m increased by 4,064 sequences. When the evolution of mitochondrial

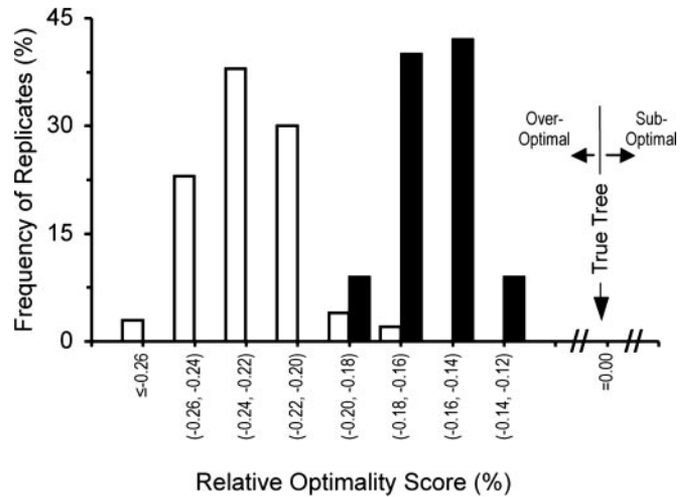


Fig. 6. Distribution of relative optimality scores (R) of NJ-SE trees (filled bars) and NJ-IE trees (open bars) when a type B model tree (Fig. 2B) with 4,096 sequences is used. R is defined as $(S_{NJ} - S_T)/S_T$, where S_{NJ} and S_T are the sum of all branch lengths for a NJ tree and the true tree, respectively (32). Therefore, when a NJ tree is the same as the true tree, R is 0. The sum (S_{NJ}) of branch lengths of a NJ tree is often smaller than that of the true tree when the number of sequences is large, and R is usually negative. However, in general, the R values for NJ-SE trees are closer to 0 (R value for the true tree) than those for NJ-IE trees are. In this simulation the TN model with $k_1 = k_2 = 4$, $g_A = g_T = g_C = g_G = 0.25$, and $n = 1,000$ was used.

DNA was considered, P_C was much lower than that for nuclear genes (55% for $m = 32$) (Fig. 5A, open circles). Yet, the decline of P_C with increasing m was quite small (Fig. 5A, open circles). The P_C value for $m = 4,096$ was smaller than that for $m = 32$ only by 2.2%. A small extent of decline of P_C with increasing m was observed even when type C model trees (varying evolutionary rates) were used (Fig. 5A, open and filled triangles) or when randomly generated model trees (types D and E) were used (Fig. 5B, squares and diamonds). The decrease in P_C was <10% in every case in Fig. 5. These results indicate that P_C does not decline very much when m increases from 32 to 4,096 and, therefore, NJ can be used efficiently even when hundreds or thousands of sequences are used.

Similar results were obtained for NJ-SE trees when the substitution pattern and G + C content vary with evolutionary lineage or when the sequence length is reduced to a half (results not shown). Therefore, although the absolute values of P_C depend on the substitution pattern or sequence length, the relationship between P_C and m is nearly the same for all cases. In other words, P_C generally declines with increasing m , but the extent of the decline is remarkably small.

Discussion

We have seen that the SE of evolutionary distances reduces the variances of distance estimates considerably and consequently increases the accuracy (P_C) of NJ trees. We have also seen that the P_C of NJ trees does not decrease significantly when the number of sequences (m) increases from 32 to 4,096. However, this latter property is not necessarily due to the use of SE distances, because a similar property was observed even with IE distances when m was ≤ 100 (10). Some authors have reported a substantial decrease of P_C when m increased from 50 to 100 (e.g., ref. 28). In the latter study the lengths of interior branches were considerably smaller for large trees than for small trees, and this difference contributed to the substantial decrease of P_C . Therefore, it is important to maintain the same interior branch lengths for a comparison of P_C for large and small trees, as mentioned earlier.

In reality, however, our assumption that all interior branches have the same length irrespective of the tree size is unrealistic. When the number of sequences used is very large, it is quite likely that some interior branches are relatively long and others are very short. If the expected length of an interior branch is very short, no substitutions may occur in the branch when the number of nucleotides examined is small. In this case it would be difficult to resolve the clades associated with the short branches. Therefore, our results should be interpreted only as a guideline. As long as the interior branch is sufficiently long and a sufficient number of nucleotides per sequence is used, the NJ method is capable of producing reasonably accurate trees.

The relatively high P_C values obtained for large trees in this study are partly due to the use of SE distances, which have smaller standard errors than IE distances. This SE approach is effectively the same as the use of a larger number of nucleotides in the IE approach. For example, when model tree *B* was used, the average standard error of SE distances for the TN model with $k_1 = k_2 = 20$, $g_A = g_T = 0.4$, $g_C = g_G = 0.1$, and $n = 1,000$ was similar to that of IE distances with $n = 1,650$. Furthermore, another merit of using the SE approach instead of the IE approach exists. That is, when the number of sequences (m) is as large as 1,000, estimates of IE distances are not always obtainable, because the arguments of logarithms in the analytical formulas may become negative by chance (31). The proportion of such unestimable cases increases as m increases in the IE method (Fig. 1C). By contrast, we have encountered no such cases in our simulation when the SE method was used. Of course, unestimable cases might happen even with the SE method, if the extent of sequence divergence is very high, but the probability of occurrence of such events is much smaller in the SE approach than in the IE approach.

The higher accuracy of NJ-SE trees also appears to be related to the fact that the sum of branch lengths (S) for NJ-SE trees is on average closer to that of the true tree than the S value for NJ-IE trees (Fig. 6). It is already known that the S value for NJ-IE trees is on average considerably smaller than that of the true tree when the number of sequences relative to the sequence length is small (8, 11, 12, 32). Our simulation results indicate that the S values for NJ-SE trees are closer to that of the true tree although, in general, they are still smaller than that of the true tree (Fig. 6). These results also indicate that although the NJ method examines a minuscule portion of the entire tree space, a further search for trees with smaller S values does not necessarily lead to a tree closer to the true tree. This is because the optimization principle does not work well in phylogenetic inference when m is large relative to n , whether the minimum evolution, maximum parsimony, or maximum likelihood method is used (1, 32).

We therefore conclude that the prospects are bright for using the NJ method for generating initial evolutionary hypotheses for very large species and multigene family trees. These large phylogenies often span long evolutionary times in which the pattern of nucleotide substitution is expected to be complex (1, 4). In these cases, the use of the SE approach with sophisticated models of DNA substitution is expected to improve the accuracy of phylogenetic inference.

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