Assessing Confidence in Root Placement on Phylogenies:
An Empirical Study Using Non-Reversible Models

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ABSTRACT

Using time-reversible Markov models is a very common practice in phylogenetic analysis, because although we expect many of their assumptions to be violated by empirical data, they provide high computational efficiency. However, these models lack the ability to infer the root placement of the estimated phylogeny. In order to compensate for the inability of these models to root the tree, many researchers use external information such as using outgroup taxa or additional assumptions such as molecular-clocks. In this study, we investigate the utility of non-reversible models to root empirical phylogenies and introduce a new bootstrap measure, the rootstrap, which provides information on the statistical support for any given root position.

Availability and implementation: A python script for calculating rootstrap support values is available at https://github.com/suhanaser/Rootstrap.

[phylogenetic inference, root estimation, bootstrap, non-reversible models]
The most widely used method for rooting trees in phylogenetics is the outgroup method. Although the use of an outgroup to root an unrooted phylogeny usually outperforms other rooting methods (Huelsenbeck, et al. 2002), the main challenge with this method is to find an appropriate outgroup (Watrous and Wheeler 1981; Maddison, et al. 1984; Smith 1994; Swofford, et al. 1996; Lyons-Weiler, et al. 1998; Milinkovitch and Lyons-Weiler 1998). Outgroups that are too distantly-related to the ingroup may have substantially different molecular evolution than the ingroup, which can compromise accuracy. And outgroups that are too closely related to the ingroup may not be valid outgroups at all.

It is possible to infer the root of a tree without an outgroup using molecular clocks (Huelsenbeck, et al. 2002; Drummond, et al. 2006). A strict molecular-clock assumes that the substitution rate is constant along all lineages, a problematic assumption especially when the ingroup taxa are distantly related such that their rates of molecular evolution may vary. Relaxed molecular-clocks are more robust to deviations from the clock-like behaviour (Drummond, et al. 2006), although previous studies have shown that they can perform poorly in estimating the root of a phylogeny when those deviations are considerable (Tria, et al. 2017).

Other rooting methods rely on the distribution of branch lengths, including Midpoint Rooting (MPR) (Farris 1972), Minimal Ancestor Deviation (MAD) (Tria, et al. 2017), and Minimum Variance Rooting (MVR) (Mai, et al. 2017). Such methods also assume a clock-like behaviour; however, they are less dependent on this assumption as the unrooted tree is estimated without it. Similar to inferring a root directly from molecular-clock methods, the accuracy of those rooting methods decreases with higher deviations from the molecular-clock assumption (Mai, et al. 2017).
Other less common rooting methods that can be used in the absence of outgroup are:

rooting by gene duplication (Dayhoff and Schwartz 1980; Gogarten, et al. 1989; Iwabe, et al. 1989), indel-based rooting (Rivera and Lake 1992; Baldauf and Palmer 1993; Lake, et al. 2007), rooting species tree from the distribution of unrooted gene trees (Allman, et al. 2011; Yu, et al. 2011), and probabilistic co-estimation of gene trees and species tree (Boussau, et al. 2013).

All the methods mentioned above, apart from the molecular-clock, infer the root position independently of the ML tree inference. The only existing approach to include root placement in the ML inference is the application of non-reversible models. Using non-reversible substitution models relaxes the fundamental assumption of time-reversibility that exists in the most widely used models in phylogenetic inference (Jukes and Cantor 1969; Kimura 1980; Hasegawa, et al. 1985; Tavaré 1986; Dayhoff 1987; Jones, et al. 1992; Tamura and Nei 1993; Whelan and Goldman 2001; Le and Gascuel 2008). This in itself is a potentially useful improvement in the fit between models of sequence evolution and empirical data. In addition, since non-reversible models naturally incorporate a notion of time, the position of the root on the tree is a parameter that is estimated as part of the ML tree inference. Since the incorporation of non-reversible models in efficient ML tree inference software is relatively new (Minh, et al. 2020), we still understand relatively little about the ability of non-reversible models to infer the root of a phylogenetic tree, although a recent simulation study has shown some encouraging results (Bettisworth and Stamatakis 2020).

Regardless of the rooting method and the underlying assumptions, it is crucial that we are able to estimate the statistical confidence we have in any particular placement of the root on a phylogeny. A number of previous studies have sensibly use ratio likelihood tests such as the Shimodaira-Hasegawa (SH) test (Shimodaira and Hasegawa 1999) and the Approximately Unbiased (AU) test (Shimodaira 2002) to compare a small set of potential root placements,
rejecting some alternative root placements in favour of the ML root placement e.g. (Nardi, et al. 2003; Steenkamp, et al. 2006; Jansen, et al. 2007; Moore, et al. 2007; Williams, et al. 2010; Kocot, et al. 2011; Zhou, et al. 2011; Whelan, et al. 2015; Zhang, et al. 2018), these tests are still somewhat limited in that they do not provide the level of support the data have for a certain root position.

There is strong empirical evidence that molecular evolutionary processes are rarely reversible (Squartini and Arndt 2008; Naser-Khdour, et al. 2019), but few studies have explored the accuracy of non-reversible substitution models to root phylogenetic trees (Huelsenbeck, et al. 2002; Yap and Speed 2005; Williams, et al. 2015; Cherlin, et al. 2018; Bettisworth and Stamatakis 2020). Most studies that have looked at this question in the past have focused on either simulated datasets (Huelsenbeck, et al. 2002; Jayaswal, et al. 2011; Cherlin, et al. 2018; Bettisworth and Stamatakis 2020) or relatively small empirical datasets (Yang and Roberts 1995; Yap and Speed 2005; Jayaswal, et al. 2011; Heaps, et al. 2014; Williams, et al. 2015; Cherlin, et al. 2018). In both cases, the addressed substitution models were nucleotide models, and to our knowledge, no study has yet investigated the potential of amino acid substitution models in inferring the root placement of phylogenetic trees.

In this paper, we focus on evaluating the utility of non-reversible amino acid and nucleotide substitution models to root the trees, and we introduce a new metric, the *rootstrap support value*, which estimates the extent to which the data support every possible branch as the placement of a root in a phylogenetic tree. Unlike previous studies that used Bayesian methods with non-reversible substitution models to infer rooted ML trees (Heaps, et al. 2014; Cherlin, et al. 2018), we will conduct our study in a Maximum likelihood framework using IQ-TREE (Minh, et al. 2020). A clear advantage of the Maximum likelihood over the Bayesian analysis is that there is no need for a prior on the parameter distributions, which sometimes can affect tree inference (Huelsenbeck, et al. 2002; Cherlin, et al. 2018). Even though estimating
the non-reversible model’s parameters by maximizing the likelihood function seems more computationally intensive than calculating posterior probabilities (Huelsenbeck, et al. 2002), the IQ-TREE algorithm sufficiently fast to allow us to estimate root placements, with rootstrap support for very large datasets.

MATERIAL AND METHODS

The “Rootstrap” Support, and measurements of error in root placement

To compute rootstrap supports, we conduct a bootstrap analysis, i.e., resampling alignment sites with replacement, to obtain a number of bootstrap trees. We define the rootstrap support for each branch in the ML tree, as the proportion of bootstrap trees that have the root on that branch. Since the root can be on any branch in a rooted tree, the rootstrap support values are computed for all the branches including external branches. The sum of the rootstrap support values along the tree are always smaller than or equal to one. A sum that is smaller than one can occur when one or more bootstrap replicates are rooted on a branch that does not occur in the ML tree (Fig. 1).

FIGURE 1. Illustration of the rootstrap concept. (a) The bootstrap replicates trees. (b) The ML tree with the rootstrap support values for each branch. Note that the sum of the rootstrap support values is less than 100% due to 100 bootstrap replicates trees (green) that have their root at a branch that does not exist in the ML tree.
If the true position of the root is known (e.g. in simulation studies) or assumed (e.g. in the empirical cases we present below), we can calculate additional measurements of the error of the root placement. We introduce two such measurements here: root branchlength error distance (rBED) and root split error distance (rSED). Since the non-reversible model infers the exact position of the root on a branch, we define the root branchlength error distance (rBED) as the range between the minimum and maximum distance between the inferred root position and the “true root” branch. If the true root is on the same branch as the ML tree root, then rBED will be between 0 and the distance between the ML tree root and the farthest point on that branch (Fig. 2). Since rBED is based on branch lengths only, it ignores the absolute number of splits between the ML tree root and the true root; and therefore, the rBED for the true root being on the same ML root branch can be bigger than the rBED for the true root being on a different branch (e.g. Fig. 2). In order to account for the number of splits (nodes) between the ML tree root and the true root, we define root split error distance (rSED) as the number of splits between the ML root branch and the branch that is believed to contain the true root (Fig. 2).

**Figure 2.** An example to illustrate the root error distance. (a) the ML rooted tree, (b) the root branch-length error distance (rBED) if the true root is believed to be on the same ML
root branch (rSED = 0), (c) the rBED if the true root is believed to be on the branch between D and the clade of C + B (rSED = 3).

The rootstrap, rBED, and rSED assess different aspects of the root placement. While the rootstrap offers an indication of the support that the data have for a certain branch to be the root branch, rBED and rSED provide an estimation to the accuracy of the method in estimating the exact root position if the root position is known or assumed in advance. In other words, the rootstrap value is a measure for the adequacy of the data to validate a root placement given the model, while rBED, and rSED are measures of the accuracy of the non-reversible model to find the root placement given the data.

Empirical Datasets

Because non-reversible amino acid models require the estimation of a large number of parameters, and because we suspected that the information in any such analysis on the placement of the root branch of a tree might be rather limited, we searched for empirical datasets that met a number of stringent criteria:

(1) Existence of both DNA and amino acid multiple sequences alignments (MSA) for the same loci.

(2) Genome-scale MSAs to ensure that the MSAs have as much information as possible with which to estimate the non-reversible models’ free parameters and the root position. Since we do not know the number of sites required to correctly infer the rooted ML tree, we define 100,000 sites as the minimum number of required sites. This also allows us to subsample the dataset to explore the ability of smaller datasets to infer root positions.

(3) Highly-curated alignments: since the quality of the inferred phylogeny is highly dependent on the quality of the MSA (Philippe, et al. 2011), we focussed on datasets that were highly-curated for misalignment, contamination, and paralogy.
(4) Existence of several clades for which there is a very strong consensus regarding their root placement. Since we are interested in evaluating the performance of non-reversible models to infer root placements in an empirical rather than a simulation context, we need to identify monophyletic sub-clades for which we can be almost certain about their root position. This enables us to divide the dataset into non-overlapping sub-clades for which we are willing to assume we know the root positions. Furthermore, we define the minimum number of taxa in each sub-dataset as five.

We initially identified a number of genome-scale datasets that contained large numbers of nucleotide and amino acid MSAs. In many cases, it was difficult to determine whether these alignments had been rigorously curated, and even more challenging to find datasets for which the root position of a number of subclades could be assumed with confidence. The only dataset that met all of our criteria was a dataset of placental mammals with 78 ingroup taxa and 3,050,199 amino acids (Wu, et al. 2019). This dataset was originally published as an MSA (Liu, et al. 2017) based on very high-quality sequences from Ensembl, NCBI, and GenBank databases. After receiving detailed critiques for potential alignment errors (Gatesy and Springer 2017), the dataset was further processed to remove potential sources of bias and error, and an updated version of the dataset was recently published (Wu, et al. 2018). The fact that this alignment comes from one of the most well-studied clades on the planet, has been independently curated and critiqued by multiple groups of researchers and includes truly genome-scale data, makes it ideally suited for our study.

Selecting Clades with a Well-Defined Root

Since our main objective in this study is to evaluate the effectiveness of non-reversible models and the rootstrap value in estimating and measuring the support for a given root placement on empirical datasets, we must identify a collection of sub-clades of the larger
mammal dataset for which it is reasonable to assume a root position. We acknowledge, of course, that outside a simulation framework it is not possible to be certain of the position of the root position of a clade. Nevertheless, it is possible to identify clades for which the position of the root is well supported and non-controversial, thus minimising the chances that the assumption of particular root position is incorrect. To achieve this, we analysed the root position of each order and superorder in the dataset, and defined “well-defined clades” that fulfilled all of the following criteria:

1. It contains at least five taxa. This ensures that the probability of obtaining a random ML rooted tree to be at most 0.95%. For clades with four taxa, there are 15 different rooted topologies, and therefore a 6.7% probability to get any of these topologies by chance. On the other hand, for clades with at least five taxa, there are at least 105 different rooted topologies and maximum probability of 0.95% to randomly get one of them as the ML tree.

2. The bootstrap support for the deepest two levels of branches leading to that clade in the phylogenetic tree calculated from the whole dataset is 100%: since the bootstrap value indicates the support the data have for a certain branch, we require 100% support for the deepest two levels of branches leading to a certain clade in the whole tree (Appendix Fig. A.1). This requirement ensures that there is strong support in the dataset for the root position of the clade when the entire dataset is rooted with an outgroup.

3. The gene concordance factor (gCF) and the site concordance factor (sCF) for the deepest two levels of branches leading to the clade are significantly greater than 33%. The site Concordance Factor (sCF) is calculated by comparing the support of each site in the alignment for the different arrangements of quartet around a certain branch. In other words, an sCF of 33% means equal support for any of the possible arrangements.
Therefore, we require that the sCF of the deepest two levels of branches leading to that clade to be significantly greater than 33%. The gene Concordance Factor (gCF) of a branch is calculated as the proportion of gene trees contain that branch. Although there is no threshold regarding the required proportion of genes concordant with a certain branch, for convenience, we define branches with gCF significantly greater than 33% as branches that are concordant with enough genes in the alignment (Minh, et al. 2020). To test whether the sCF and the gCF are significantly greater than 33%, we use a simple binomial test with a success probability of 0.33.

(4) At least 95% of the studies that have been published in the last decade support this clade: we searched google scholar for all published papers since 2009 that determine the root of the addressed clade. We then checked if at least 95% of those papers agree that the root position of the clade matches that in the ML tree we estimate from the whole dataset (see supplementary material).

Estimating unrooted Phylogenies

For the whole nucleotide and amino-acid datasets with ingroup and outgroup taxa, we inferred the unrooted phylogeny using IQ-TREE (Nguyen, et al. 2015) with the best-fit fully partitioned model (Chernomor, et al. 2016) and edge-linked substitution rates (Duchene, et al. 2020). We then determined the best-fit reversible model for each partition using ModelFinder (Kalyaanamoorthy, et al. 2017). See the algorithm for finding well-defined clades in Appendix Algorithm A.1.

Estimating Rooted phylogenies

For each well-defined clade, we first removed all other taxa from the tree and then sought to infer the root of the well-defined clade using non-reversible models without outgroups. Using the best partitioning scheme from the reversible analysis, we inferred the
rooted tree for each well-defined clade with the non-reversible models for amino acid (NR-AA) and nucleotide (NR-DNA) sequences (Minh, et al. 2020). This approach fits a 12-parameter non-reversible model for DNA sequences, and a 380-parameter non-reversible model for amino acids. Details of the command lines used are provided in the supplementary material section “Algorithm A.2”. Each analysis returns a rooted tree. We performed 1000 non-parametric bootstraps of every analysis to measure the rootstrap support.

To assess the performance of the rootstrap and the ability of non-reversible models to estimate the root of the trees on smaller datasets, we also repeated every analysis on subsamples of the complete dataset. For each well-defined clade, we performed analysis on the complete dataset (100%) as well as datasets with 10%, 1% and 0.1% of randomly-selected loci from the original alignment.

The confidence set of root branches using the Approximately Unbiased test

In addition to the rootstrap support, we calculate the confidence set of all the branches that may contain the root of the ML tree using the Approximately Unbiased (AU) test (Shimodaira 2002). To do this, we re-root the ML tree with all possible placements of the root (one placement for each branch) and calculate the likelihood of each tree. Using the AU test, we then ask which root placements can be rejected in favour of the ML root, using an alpha value of 5%. We define the root branches confidence set as the set of root branches that are not rejected in favour of the ML root placement.

Reducing systematic bias by removing third codon positions and loci that fail the MaxSym test

As it is common in many phylogenetic analyses to remove third codon positions from the alignment (Swofford, et al. 1996), we wanted to assess the effect of removing third codon positions on the root inference and the rootstrap values in nucleotide datasets. For that
purpose, we remove all the third codon positions from the nucleotide alignments and re-ran
the analysis using the NR-DNA model.

Moreover, although the NR-AA and NR-DNA models relax the reversibility assumption,
they still assume stationarity and homogeneity. To reduce the systematic bias produced by
violating these assumptions, we used the MaxSym test (Naser-Khdour, et al. 2019) to remove
loci that violate those assumptions in the nucleotide and amino acid datasets, and then re-ran
all analyses as above.

Applying the methods to two clades whose root position is uncertain

In addition to the well-defined clades, we used the methods we propose here to infer
the root of two clades of mammals whose root position is controversial; Chiroptera and the
Cetartiodactyla.

There is a controversy around the root of the Chiroptera (bats) in literature. The two
most popular hypotheses are: 1) the Microchiroptera-Megachiroptera hypothesis; where the
root is placed between the Megachiroptera, which contains the family Pteropodidae, and the
Microchiroptera, which contains all the remaining Chiroptera families. This hypothesis is
well supported in the literature (Agnarsson, et al. 2011; Meredith, et al. 2011). However,
more recent studies seem to provide less support for this hypothesis; 2) the
Yinpterochiroptera-Yangochiroptera hypothesis, in which the Yangochiroptera clade includes
most of Microchiroptera and the Yinpterochiroptera clade includes the rest of
Microchiroptera and all of Megachiroptera. There is growing support for this hypothesis in
the literature (Meganathan, et al. 2012; Tsagkogeorga, et al. 2013; Ren, et al. 2018; Reyes-
Amaya and Flores 2019).

Similar to Chiroptera, the root of Cetartiodactyla remains contentious in the literature.
The three main hypotheses regarding the root of Cetartiodactyla are: 1) Tylopo...
group for all other cetartiodactylans; 2) Suina as the sister group for all other
cetartiodactylans; 3) the monophyletic clade containing Tylopoda and Suina as the sister
group for all other cetartiodactylans.

To ascertain whether certain sites or loci had very strong effects on the placement of
the root we follow the approach of Shen et. al. (Shen, et al. 2017) and calculate the difference
in site-wise log-likelihood scores (ΔSLS) and gene-wise log-likelihood scores (ΔGLS) between
the supported root positions for each clade.

RESULTS

Inference of the mammal tree and selection of well-defined clades

The trees inferred from the whole datasets with the nucleotide-reversible model and
the amino-acid-reversible model (Appendix Fig. A.2, Appendix Fig. A.3, Appendix Table
A.2) are consistent with the published tree (Liu, et al. 2017). Five clades met all the criteria of
well-defined clades, namely, Afrotheria, Bovidae, Carnivora, Myomorpha, and Primates in
both amino acid and nucleotide datasets (see Appendix Table A.1 and Appendix Table A.2).

High accuracy of the AA non-reversible model in inferring the root

Using NR-AA, we inferred the correct root with very high rootstrap support for all
five well-defined clades (Appendix Table A.3). Moreover, for all the five clades, the true root
was the only root placement in the confidence set of the AU test.

Our results show that using only 10% of the sites in the amino acid alignments gave
very high rootstrap support values (> 98%) for most of these clades (Fig. 3). Moreover, in
some datasets, 1% of the sites were enough to give a very high rootstrap support value. Yet,
using only 0.1% decreased the rootstrap support value noticeably in all datasets (Appendix
Table A.3). These values are shown for each dataset in Figure 3, where the X-axis is plotted in terms of parsimony-informative sites to allow for a more direct comparison between datasets. Although the rootstrap support for the true root improves as the number of parsimony-informative sites increase, in some datasets (e.g. Afrotheria nucleotide dataset) this is not the case (Fig. 3).

**FIGURE 3.** The rootstrap support value for each clade as a function of the number of parsimony-informative sites.

**Poor performance of the DNA non-reversible model in inferring the root**

We correctly inferred the root for four out of the five nucleotide datasets with the NR-DNA model. However, the rootstrap support was generally lower than in the amino-acid datasets (Fig. 3, Appendix Tables A.3 and A.4). In addition, our results show that removing the third codon positions does not improve the rootstrap support value. In contrast, in some datasets removing third codon positions decreased the rootstrap support value and increased the rSED (Table 1).

**TABLE 1.** Rootstrap support and rSED values in whole nucleotide datasets and nucleotide datasets without third codon positions.

| Clades  | All loci | Without 3rd | Without 3rd |
|---------|----------|-------------|-------------|
|         | rootstrap | rSED | rootstrap | rSED |
| Afrotheria | 0.0% | 2 | 0.0% | 2 |
| Primates | 90.1% | 0 | 90.1% | 0 |
Removing loci that violate the stationarity and homogeneity assumptions improves the rootstrap support

As expected, our results show that removing loci that fail the MaxSym test improves the root placement inference and the rootstrap support values when the rootstrap support value was less than 100% and/or the root placement was inferred incorrectly, as the case in some nucleotide datasets (TABLE 2).

**TABLE 2.** Rootstrap support values in whole datasets and datasets with loci that passed the MaxSym test only.

| Clade       | Amino Acid | Nucleotide |
|-------------|------------|------------|
|             | all loci   | MaxSym     | all loci   | MaxSym     |
| Afrotheria  | 100.0%     | 100.0%     | 0.0%       | 8.4%       |
| Primates    | 100.0%     | 100.0%     | 99.7%      | 99.9%      |
| Myomorpha   | 100.0%     | 100.0%     | 73.2%      | 88.3%      |
| Carnivora   | 100.0%     | 100.0%     | 100.0%     | 100.0%     |
| Bovidae     | 100.0%     | 100.0%     | 100.0%     | 100.0%     |

Microchiroptera-Megachiroptra or Yinpterochiroptera-Yangochiroptera?

Using the whole amino acid dataset, our results show 65.5% rootstrap support for the Yinpterochiroptera-Yangochiroptera hypothesis and 23.2% for the Microchiroptera-Megachiroptra hypothesis. The remaining 11.3% of the rootstrap support goes to supporting the branch leading to Rhinolophoidea as root branch of the bats (Fig. 4). Removing amino acid loci that fail the MaxSym test (110 loci) gives similar results, with 65.9% rootstrap support for the Yinptero-Yango hypothesis and 25.6% rootstrap support for the Micro-Mega hypothesis. In both cases, the AU test could not reject any of the three root positions that received non-zero rootstrap support (Appendix Table A.5).
**Figure 4.** The ML rooted tree as inferred from the whole Chiroptera amino acid dataset. Bold branches are branches in the AU confidence set. Blue values under each branch are the rootstrap support values.

Using the NR-DNA model gives 100% rootstrap support for the Yinptero-Yango hypothesis, and we can confidently reject the Micro-Mega hypothesis in favour of the Yinptero-Yango hypothesis using the AU test (Appendix Fig. A.4). Yet, removing nucleotide loci that fail the MaxSym test (~25% of the loci) decreases the support for the Yinptero-Yango hypothesis to 90.1%, although we can still confidently reject the Micro-Mega hypothesis using the AU test (Appendix Table A.5).

Interestingly, when we randomly subsample 10%, 1%, and 0.1% of the loci in the nucleotide dataset, we consistently get the Yinptero-Yango hypothesis as the ML tree and the solely rooted topology in the AU confidence set (Appendix Table A.5). Moreover, the rootstrap support value for the Yinptero-Yango hypothesis increases and the rootstrap support value for the Micro-Mega hypothesis decreases as more parsimony-informative sites are added to the alignment, for both nucleotide and amino acid datasets (Fig. 5, Appendix Table A.5).
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![Graph showing rootstrap support as a function of the number of parsimony-informative sites for DNA and amino acid datasets.](image)

**Figure 5.** Rootstrap support value as a function of the number of parsimony-informative characters in the Chiroptera nucleotide and amino acid datasets.

The ΔGLS and ΔSLS values (Shen, et al. 2017) reveal that approximately half of the nucleotide and amino acid loci prefer the Yinptero-Yango hypothesis while the other half prefers Micro-Mega hypothesis. Furthermore, slightly less than half of the nucleotide sites prefer the Yinptero-Yango hypothesis. However, more than two-thirds of the amino acid sites prefer the Yinptero-Yango hypothesis (Appendix Fig. A.5). The distributions of ΔGLS and ΔSLS (Appendix Fig. A.6) show that a small proportion of the amino acid loci (~1%) have very strong support for the Micro-Mega hypothesis, and removing those loci from the alignment increased the rootstrap support for the Yinptero-Yango hypothesis to 76.6%.

Nonetheless, both root placements are still in the confidence set of the AU test (Appendix Table A.5) with the amino acid dataset. On the other hand, removing nucleotide loci with the highest absolute ΔGLS value still gives the Yinptero-Yango hypothesis as the ML tree and the sole topology in the AU confidence set. We conclude that while the nucleotide data show a clear preference to the Yinptero-Yango hypothesis, the amino acid data do not allow us to distinguish between the two leading hypotheses for the placement of the root of the Chiroptera based on rooting with non-reversible models.
The ambiguous root of Cetartiodactyla

The ML tree inferred with the whole amino acid dataset places the clade containing Tylopoda (represented by its only extant family; Camelidae) and Suina as the most basal cetartiodactylan clade with 71.8% rootstrap support (Fig. 6). Yet, The AU test did not reject Tylopoda as the most basal clade. On the other hand, the ML tree inferred with the whole nucleotide dataset places Tylopoda as the most basal clade with 71.0% rootstrap support, and we can confidently reject the Tylopoda + Suina hypothesis using the AU test (Appendix Fig. A.7).

Removing the amino acid loci that failed the MaxSym test (~1%) still places Tylopoda + Suina as the basal-most clade, yet, it decreases the rootstrap support for the Tylopoda + Suina hypothesis to 63.3% and increases the rootstrap support for the Tylopoda hypothesis to 28.5%. However, we still cannot reject either of the hypotheses using the AU test (Appendix Table A.6).

The ML rooted tree of as inferred from the whole Cetartiodactyla amino acid dataset. Bold branches are branches in the AU confidence set. Blue values under each branch are the rootstrap support values.
Removing the nucleotide loci that failed the MaxSym test (~1%) still places Tylopoda as the basal-most clade and the only rooted topology in the AU confidence set. However, it decreases the rootstrap support for the Tylopoda hypothesis to 68.7% and increases the rootstrap support for the Tylopoda + Suina hypothesis to 20.1% (Appendix Table A.6). The results from the subsample datasets are mixed (Fig. 7). Analyses on smaller datasets show no clear pattern in the placement of the root (Appendix Table A.6), leading us to conclude only that the analyses of the whole dataset is likely to provide the most accurate result, but that it is plausible that adding more data may lead to different conclusions in the future.

![Rootstrap Support](image)

**Figure 7.** Rootstrap support value as a function of the number of parsimony-informative characters in the Cetartiodactyla nucleotide and amino acid datasets.

ΔGLS analyses reveal that approximately, half of the amino acid and nucleotide loci favour the Tylopoda+Suina hypothesis, while the other half of loci favour the Tylopoda hypothesis (Appendix Figs. A.8-9). On the other hand, two-thirds of the amino acid sites and more than 80% of the nucleotide sites favour the Tylopoda+Suina hypothesis. Removing 1% of the amino acid loci with the highest absolute ΔGLS values still places Tylopoda + Suina as the most basal clade. However, the rootstrap support of the Tylopoda + Suina decreased to 63.2% and the rootstrap support for the Tylopoda hypothesis remains approximately the same.
(~14.5%), while the rootstrap support for the Suina hypothesis increases from 13.7% to 22.4%. Yet, both the Tylopoda + Suina hypothesis and the Tylopoda hypothesis are in the confidence set of the AU test, while the Suina hypothesis is rejected by the AU test (Appendix Table A.6). Removing 1% of the nucleotide loci with the highest absolute ΔGLS values gives the Tylopoda+Suina as the most basal clade of Cetartiodactyla with 39.7% rootstrap support. However, the solely rooted topology in the AU confidence set is the topology in which the root is placed on the branch leading to Suina (Appendix Table A.6). We conclude that neither the nucleotide nor the amino acid data are adequate to infer the root placement of Cetartiodactyla with non-reversible models.

**DISCUSSION**

In this paper, we introduced a new measure of support for the placement of the root in a phylogenetic tree, the rootstrap support value, and applied it to empirical amino acid and nucleotide datasets inferred using non-reversible models implemented in IQ-TREE (Minh, et al. 2020). The rootstrap is a useful measure because it can be used to assess the statistical support for the placement of the root in any rooted tree, regardless of the rooting method. In a Maximum Likelihood setting, interpretation of the rootstrap support is similar to the interpretation of the classic nonparametric bootstrap. In a Bayesian setting, the same procedure could be used to calculate the posterior probability of the root placement given a posterior distribution of trees. It is noteworthy that the rootstrap support value is not a measure of the accuracy of the root placement and therefore should not be interpreted as such. However, it provides information about the robustness of the root inference with regard to resampling the data. This interpretation is consistent with the interpretation of the
ROOTSTRAP SUPPORT

nonparametric bootstrap (Holmes 2003) but with regard to the root placement instead of the whole tree topology.

In addition to the rootstrap support value, we introduced another two metrics; the root branch-length error distance (rBED), and the root split error distance rSED. Similar to the rootstrap metric, these additional metrics can be used in with any approach that generates rooted phylogenetic trees. We note that both metrics require the true position of the root to be known (or assumed) and that the rBED requires the rooting method to be able to accurately place the root in a specific position of the root branch.

In this study, we used these and other methods to assess the utility of non-reversible models to root phylogenetic trees in a Maximum Likelihood framework. We focussed on applying these methods to a large and very well curated phylogenomic dataset of mammals, as the mammal phylogeny provides perhaps the best opportunity to find clades for which the root position is known with some confidence. As expected, our results show an exponential increase in the rootstrap support for the true root as we add more information to the MSA. Our results suggest that non-reversible amino-acid models are more useful for inferring root positions than non-reversible DNA models, which is consistent with results from previous simulations using the NR-DNA model (Bettisworth and Stamatakis 2020). One explanation for this difference between the NR-DNA and the NR-AA models is the bigger character-state space of the NR-AA models. These models have 400 parameters (380 rate parameters and 20 amino acid frequencies) whereas NR-DNA models have only 16 parameters (12 rate parameters and 4 nucleotide frequencies). This could allow the NR-AA model to capture the evolutionary process better than the NR-DNA model, potentially providing more information on the root position of the phylogeny. This hypothesis requires some further exploration though, and we note that the actual character-space of amino acids is much smaller than
accommodated in NR-DNA models due to functional constraints on protein structure (Dayhoff, et al. 1978).

Another explanation for the difference in performance between the NR-AA and NR-DNA models is that higher compositional heterogeneity in nucleotide datasets may bias tree inference. In principle, this bias can be alleviated by removing loci that violate the stationarity and homogeneity assumptions (Naser-Khdour, et al. 2019). Our results suggest that this may be the case for the datasets we analysed: we show that removing loci that violate the stationarity and homogeneity assumptions improves the accuracy and statistical support for the placement of the root. This is not surprising since the robustness of the rootstrap, similar to the bootstrap, relies on the consistency of the inference method, so removing systematic bias should improve its performance.

We used the non-reversible approach to rooting trees along with the rootstrap support to assess the evidence for different root placements in the Chiroptera and Cetartiodactyla. Using the amino acid datasets we found that in both cases, although there tended to be higher rootstrap support for one hypothesis, neither of the current hypotheses for either dataset could be rejected. These results emphasize the importance of the rootstrap support value as a measure of the robustness of the root estimate given the data. In both the Chiroptera and Cetartiodactyla datasets the root placement varied among subsamples of the dataset, and the rootstrap support reflects this uncertainty. In both cases, the amino acid data is inadequate to distinguish between certain root placements. On the other hand, in both the Chiroptera and Cetartiodactyla, the nucleotide datasets appear to show stronger support for a single root placement. This difference between the amino acid and the nucleotide datasets results may be due to greater phylogenetic signal contained in the nucleotide characters.
Our results demonstrate that both non-reversible models can be surprisingly useful for inferring the root placement of phylogenies in the absence of additional information (such as outgroups) or assumptions (such as molecular clocks). Indeed, we show that root placements appear to be accurate even with fairly datasets as small as 50 well-curated loci between fairly closely-related taxa such as orders of mammals. We hope that the combination of non-reversible models and rootstrap support will add another tool to the phylogeneticist’s arsenal when it comes to inferring rooted phylogenies.

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