Greedy Selection of Species for Ancestral State Reconstruction on Phylogenies: Elimination Is Better than Insertion

Guoliang Li1, Jian Ma2, Louxin Zhang1*

1 Computational & Mathematical Biology, Genome Institute of Singapore, Singapore, Singapore, 2 Department of Bioengineering, University of Illinois at Urbana-Champaign, Urbana, Illinois, United States of America, 3 Department of Mathematics, National University of Singapore, Singapore, Singapore

Abstract

Accurate reconstruction of ancestral character states on a phylogeny is crucial in many genomics studies. We study how to select species to achieve the best reconstruction of ancestral character states on a phylogeny. We first show that the marginal maximum likelihood has the monotonicity property that more taxa give better reconstruction, but the Fitch method does not have it even on an ultrametric phylogeny. We further validate a greedy approach for species selection using simulation. The validation tests indicate that backward greedy selection outperforms forward greedy selection. In addition, by applying our selection strategy, we obtain a set of the ten most informative species for the reconstruction of the genomic sequence of the so-called boreo-eutherian ancestor of placental mammals. This study has broad relevance in comparative genomics and paleogenomics since limited research resources do not allow researchers to sequence the large number of descendant species required to reconstruct an ancestral sequence.

Introduction

Ancestral sequence reconstruction incorporates DNA or protein sequences from modern organisms into an evolutionary model to estimate the corresponding sequence of an ancestor that no longer exists on the Earth. In 1963, Pauling and Zuckerlandl first discussed how to study an ancient protein by inferring its sequence from the sequences of the corresponding proteins found in extant organisms, and subsequently synthesizing the sequence for functional analysis in laboratory [1]. With many genomic sequences being known and recent advances in DNA synthesis, ancestral sequence reconstruction has become an important approach to the investigation of the origins and evolution of proteins and other molecules [2,3].

Different methods have been proposed to estimate the sequence of an ancestor when the phylogeny that relates the ancestor to the extant taxa whose sequences are used for reconstruction is known [4,5]. Among these methods, the parsimony and maximum likelihood methods are the most widely used. The Fitch (parsimony) method [6] was first used for ancestral sequence reconstruction in 1984 [7]. Since then, it has been used in reconstructing ancestral sequences of digestive ribonucleases [8], chymase proteases [9], and immune RNases [10]. To reconstruct the character state at the root on a phylogeny, the method assigns states to internal nodes of the given phylogeny so as to minimize the total number of state changes on all branches. Here, the states could represent particular traits or morphological features. In ancestral DNA sequence reconstruction, the characters are sequence sites and the states are four nucleotides. The parsimony method is quite accurate and effective for the extant sequences that are closely related to each other [11].

The marginal maximum likelihood (ML) method and its variants were later proposed to infer ancestral states more accurately through an explicit statistical framework [12–16]. Given a phylogeny with branch lengths, a model that specifies change rates on all branches, and a set of observed states at the taxa, the marginal ML method selects a state, as the ancestral state, that has the maximum likelihood, the conditional probability that the observed states would have evolved given the state at the ancestor under consideration. The ML method has been used for reconstructing ancestral sequences of vertebrate rhodopsins [17], steroid hormone receptors [18], elongation factor EF-Tu [19], and the ligand-binding pocket of Family CG protein-coupled receptors [20]. Recently, the Bayesian method has also been proposed and implemented by Huelsenbeck and his coworkers [21].

Different reconstruction methods infer different proxies of an ancestral genome from the same set of extant sequences. But, it is believed that the reconstruction uncertainty involved in deciding which reconstruction method to use is generally less significant than the uncertainty arising from the different evolutionary features of the extant sequences to use for reconstruction [22]. Therefore, we study how taxon selection affects the accuracy of ancestral state reconstruction on a phylogeny in this paper. Relative to the study of phylogeny estimation, less attention has been paid to accuracy analysis for ancestral state reconstruction. There are only several papers on the merits and limitations of
various methods for ancestral sequence reconstruction [22–26], the effect of the given phylogeny topology [27–29], and the reconstruction accuracy in terms of physico-chemical properties of proteins [30].

When the genome of the common ancestor for a group of organisms is reconstructed, one would expect that the accuracy will increase with the number of extant genomic sequences used. For instance, in a recent review, Crisp and Cook [4] recommended that “if ancestral features are to be inferred from a phylogeny, a method that optimizes character states over the whole tree should be used.” However, this is not always true for the Fitch method (see the result section for detailed discussion). In other words, the accuracy of a particular method is not necessarily a monotonic function of the size of taxa used in ancestral state reconstruction. Additionally, biomedical research resource limitations often prevent a researcher from sequencing all extant genomes in an ancestral genome reconstruction project. These facts motivate us to investigate the effect of taxon sampling on inferred ancestral states and how to select taxa for the best reconstruction of an ancestral sequence.

Finally, we remark that there have been a couple of studies on the related problems. Given a set of sequenced organisms and a phylogeny over a set of both sequenced and unsequenced organisms, Sidow and his coworkers propose sequencing the organisms that maximize the additive evolutionary divergence on the phylogeny in [31]. In another work [32], McAuliffe, Jordan and Pachter present an elegant statistical framework for optimal species selection for detecting single-site reservation. Here, we propose an algorithmic approach to taxon selection for ancestral state reconstruction and investigate as an application what species are informative for reconstructing the genomic sequence of the so-called boreoeutherian ancestor.

Results

Monotonicity of Reconstruction Accuracy: Parsimony Method

Intuitively, more taxa should give better estimation of the ancestral sequence at the root of a phylogeny. However, this is not necessarily true for all reconstruction methods. The reason for this fact is probably that reconstruction accuracy is highly sensitive to the topology of the phylogeny under consideration [28,29]. For example, if the root of a phylogeny has a leaf child on a branch that is shorter than the other branch leading to a clade, reconstructing the root state from all taxa in the tree is no better than using the sole child taxon of the root when the Fitch method is used [33].

A natural question to ask is then how often this counterintuitive situation arises in practice. In evolutionary study, branch lengths in a phylogeny may satisfy a molecular clock, so that the path from the root to a leaf has equal length (or evolutionary time). Phylogenies in which branch lengths satisfy a molecular clock are said to be ‘ultrametric’. Even on an ultrametric phylogeny, reconstructing the root state from a subset of taxa can be more accurate than from all taxa. Figure 1 presents such an example, in which the reconstruction from four taxa $a, b, c, e$ is more accurate than from all taxa in the two-state Jukes-Cantor model.

The accuracy of reconstructing the root state in a phylogeny is a continuous function of branch lengths. On the phylogeny in Figure 1, it is still true that the Fitch method reconstructs the root state from taxa $a, b, c, e$ more accurately than from all the taxa when the lengths of branches leading to these four taxa increase by a small amount. Therefore, the accuracy of reconstructing the root state could increase even if some taxa that are close to the root are discarded on a phylogeny for Fitch method [34].

Monotonicity of Reconstruction Accuracy: The Marginal ML Method

Consider two subsets $L_1$ and $L_2$ of taxa such that $L_1 \subset L_2$ on a phylogeny $T$. It is known that, among all reconstruction methods, the marginal ML method has the highest accuracy of reconstructing the root state from all the taxa in the spanning subtree over $L_2$ (see [35] for example). A simple proof of this fact can be found in the Materials and Methods section. Note that reconstructing the ancestral state from the taxa of $L_1$ is just a special reconstruction in the tree spanning over $L_2$, in which the state information carried in the taxa in $A_2$ but not in $A_1$ is ignored. Therefore, when the marginal ML method is used, the reconstruction from the taxa in $L_2$ is at least as accurate as that from the taxa in $L_1$. In other words, the reconstruction accuracy of the marginal ML method is a monotonic function of the size of taxa selected for reconstruction (E. Mossel, personal communication).

Taxon Selection for Ancestral State Reconstruction

The Fitch method is efficient for ancestral state reconstruction. But, as we have shown, its reconstruction accuracy is not a monotonic function of the size of taxon sampling. When it is employed, it is necessary to first identify a subset of taxa to achieve the best reconstruction. Moreover, due to limited research resources, it is usually unlikely to sequence the large number of extant genomes in a comparative genomics project. This motivates us to investigate how to identify a small or medium number of taxa for qualitative reconstruction of an ancestral genome. Formally, we study the following taxon selection problem:

![Figure 1. An ultrametric phylogeny on which the Fitch method does not have monotonic accuracy.](https://example.com/fig1.png)

When the Fitch method is used, the accuracies of reconstructing the root state from taxa $a, b, c, e$ and from all taxa are 0.921926 and 0.915298 in the two-state Jukes-Cantor model, respectively. The Newick format of the tree is ((g:2.1553,d:2.1553):47.8447,(((f:0.8271,a:0.8271):14.1190,(h:5.2352,i:5.2352):9.7109):10.0613,(e:21.7263,(c:4.2160,-b:4.2160):17.5103):3.2811):24.9926), where the branch lengths are scaled 100 times. The conservation probability on a branch of length $t$ is $0.5(1+e^{-2t})$, where $x$ is set to 0.25. 

$\text{doi:10.1371/journal.pone.0008985.g001}$
Given a phylogeny $T$ over a set of $n$ taxa, a Markov evolutionary model of a character in $T$, a reconstruction method $M$, and an integer $k \leq n$, find the $k$ taxa that allow the character state at the root to be reconstructed with the highest accuracy over all $k$-taxon subsets when $M$ is used.

Since the reconstruction accuracy of a method depends on both the topology of the given phylogeny and the evolutionary model of the considered character, the taxon selection problem is unlikely polynomial-time solvable, although its complexity status is not yet known. Here we propose two heuristic strategies for solving this problem which originated in our linear regression study.

The first strategy selects $k$ taxa one by one in terms of incremental accuracy. We call this the forward greedy method. The algorithm first picks a taxon that is the closest to the root, breaking ties arbitrarily. In each of the next $k - 1$ steps, the algorithm selects a taxon that, together with the taxa that have been selected, gives the largest increment in reconstruction accuracy.

The other strategy is called the backward greedy method. It removes taxa one by one by considering the accuracy decrement rather than the accuracy increment. Assume there are $n$ taxa on the input phylogeny. In each of the $n - k$ steps, the method deletes a taxon with the property that the removal of this taxon leads to the least decrement in reconstruction accuracy.

We validated these two methods on 4000 random ultrametric phylogenies over 16 taxa in which branches have different lengths for each of tree heights 0.1, 0.2, 0.5, 1, 2, 5. We selected $k$ taxa by using each method in each random ultrametric phylogeny, calculated the accuracy of reconstructing the root state from the selected taxa and compared it with the accuracy of the reconstruction from all the taxa in the phylogeny. Here the exact accuracy of using the selected $k$ taxa is calculated using the branch substitution rates on the subtree spanned by the taxa. Because the accuracy of reconstructing the root state from $k$ selected taxa varies on different random phylogenies, we used the accuracy difference to measure the performance of the proposed methods on each generated ultrametric phylogeny. Figure 2 shows the average difference between the accuracies of using all the 16 taxa and using the selected $k$ taxa for reconstructing the root state in a random ultrametric phylogeny of height 2 when the Fitch method.

![Forward greedy algorithm](Figure 2)

Figure 2. The box-and-whisker plot of the difference of the accuracies of reconstructing the root state from all 16 taxa and from the selected $k$ taxa on random ultrametric phylogenies of height 2 for the Fitch method. A negative difference indicates that the reconstruction accuracy of using the selected $k$ taxa is higher than that of using all the 16 taxa in the corresponding phylogeny. The (blue) curve represents the average difference. The bottom and top of the box are the 25th and 75th percentile; the bar inside the box indicates the median; and the (red) crosses are outliers.

doi:10.1371/journal.pone.0008985.g002
was used. Note that there are a number of negative outliers for both selection methods in each plot, indicating that the selected subset of taxa leads to a better reconstruction than using all 16 taxa in the corresponding ultrametric phylogeny.

We also ran the selection methods for the marginal ML method. Computing the exact reconstruction accuracy of the marginal ML method by definition takes exponential time. It is still open if it is NP-hard or not. Because of the computational intensity of calculating the accuracy of the ML method, we ran the methods over 324 random ultrametric phylogenies for each tree height given above. Figure 3 shows the average accuracy difference for the marginal ML method in the case of tree height 2.

We repeated our experiments with 4,000 phylogenies over 16 taxa randomly generated by a Yule model using the Fitch method. To investigate how the tree topology affects the reconstruction accuracy, we assign equal length to all branches in each generated phylogeny. We reconstructed the root state of a two-state character on each generated phylogeny in a Jukes-Cantor model. Let \( p \) denote the conservation probability on the branches of a phylogeny. The probability that one state changes into the other is \( 1 - p \) on each branch. For each \( k \leq 16 \) and \( p = 0.75, 0.80, 0.85, 0.90, 0.95, 0.99 \), we selected \( k \) taxa by using each method and calculated the accuracy of reconstructing the root state from the selected taxa in each generated phylogeny. Similar facts were also observed in this case.

We observed the following facts from the above experiments. First, when the branch conservation probabilities \( p \) are smaller than 0.9 on a phylogeny, the methods frequently produced a subset of taxa that have higher reconstruction accuracy than all 16 taxa in the phylogeny. However, this becomes rare as \( p \) exceeds 0.9. In addition, as a function of the size \( k \) of taxon sampling, reconstruction accuracy increases rapidly when \( k \) is less than 10 and becomes stable when \( k \) is medium or large. When \( k \) is small, there are many outliers in the box-and-whisker plots in the figures. This indicates that the accuracy of reconstructing the root state from a subset of \( k \) taxa varies in a wide interval when \( k \) is small.

Secondly, the backward greedy algorithm generally outperforms the forward greedy algorithm. In about 80% of our experiments as

![Figure 3. The box-and-whisker plot of the difference of the accuracies of reconstructing the root state from all 16 taxa and from the selected taxa on random ultrametric phylogenies of height 2 for the marginal ML method. A negative difference indicates that the reconstruction accuracy of using the selected \( k \) taxa is higher than that of using all the 16 taxa in the corresponding phylogeny. The (blue) curve represents the average difference. The bottom and top of the box are the 25th and 75th percentile; the bar inside the box indicates the median; and the (red) crosses are outliers. doi:10.1371/journal.pone.0008985.g003](http://example.com)
shown in Figure 4, the backward greedy method produced a better taxon subset in terms of reconstruction accuracy. But, the backward greedy method is less efficient as it starts the taxon selection from the whole set of taxa in a phylogeny and removes taxa one by one until the required number of taxa remain.

We also analyzed the mean value of the reconstruction accuracy using an arbitrary subset of k taxa in a phylogeny and compared it with that of using the k taxa selected by each method. The reconstruction accuracy using the k taxa selected by each algorithm is significantly higher than using k arbitrary taxa (see Figures S1 and S2).

**Selecting Species for Reconstructing Boreoeutherian Sequence**

We now consider species sampling for reconstructing the genomic sequence of the so-called boreoeutherian ancestor of all placental mammals that lived approximately 100 million years ago. We examined the accuracy using the species selected by the two greedy methods for reconstructing the ancestral genome over the phylogeny over 24 extant species shown in Figure 5 (personal communication from Adam Siepel). Our dataset covers 20,917 base positions in the CFTR region. In the CFTR region, different base positions have, on average, similar relative rate of evolution [31].

We first reconstructed the nucleotides in these 20,917 base positions in the boreoeutherian genome by applying the base-level reconstruction program reported in [25] to the sequences of all the 24 species. Then, we selected a subset of k species (4 ≤ k ≤ 24) using both the backward and forward selection methods. We also inferred the ancestral nucleotides based on the selected species using the same reconstruction program. Since the boreoeutherian genome is unknown, it is impossible to obtain the true reconstruction accuracy of using a selected subset of species. Hence, we estimated the accuracy of reconstructing the boreoeutherian ancestral sequence from all 24 species by taking the simulation approach described in [25]. We first simulated evolutionary process starting from a hypothetical ancestral sequence of 20,000bp. We then ran the reconstruction program reported in Blanchette et al. (2004) to predict the boreoeutherian ancestral sequence from the resulting sequences at all 24 extant species. Finally, we compared the predicted ancestral sequence and the hypothetical ancestral sequence to estimate the reconstruction accuracy. The average accuracy is 99.45% (over 100 simulations). Since the reconstruction from all 24 species are quite accurate, we examined the relative accuracy by comparing the reconstruction from selected species with the reconstruction from all 24 species. The results obtained from the backward selection method are summarized in Table 1 and those from the forward selection method are given in Table 2.

The results show that the two methods output quite different subsets of k species for each k. For instance, when 4 species were selected, the backward greedy method output *marmoset, alpaca, armadillo, and sloth*, whereas the forward greedy method output *gorilla, orangutan, rhesus, and marmoset*. The only common species in these two selected subsets is *marmoset*. When the number of species to be selected is 15, the two subsets output from the backward and forward methods have only the following 7 species in common: *marmoset, mouselemur, treeshrew, mouse, guinea pig, squirrel, and rabbit*. This observation suggests that the effect of a species on ancestral sequence reconstruction should not be examined separately.

**Figure 4. The backward greedy selection vs the forward greedy selection.** Each circle represents a selection instance. If reconstructing the root state from the 12-taxon subset output from the backward selection is more accurate than that from the forward selection on a random 16-taxon phylogeny, the corresponding circle falls in the region above the line \(x = y\); otherwise, it falls in the region below the line. doi:10.1371/journal.pone.0008985.g004
Our test results demonstrate that the backward greedy approach again outperformed the forward greedy approach for the boreoeutherian sequence reconstruction. The results in Table 1 also suggest that using just 20 or more species, one can reconstruct in the base level the boreoeutherian genome with high accuracy once the genomic sequences are aligned.

The species selection made by the backward greedy method is broadly consistent with an early study of Cooper et al. [31]. In investigating the relative divergence of a species with respect to human, mouse and rat, they find that dog is the most divergence informative and cow is more informative than the others. Our backward greedy selection process first eliminated cow, then guinea_pig and then dog, showing that dog is more informative than cow for inferring the boreoeutherian genomic sequence. This consistency further suggests that the backward greedy method is superior to the forward greedy method.

Discussion

Monotonicity of Ancestral State Reconstruction

Reconstructing the ancestral state of a character is far more complicated than we thought. Our first finding is that the accuracy of reconstructing the root state of a character is not a monotonic function of the taxon sampling size even in an ultrametric phylogeny for the Fitch method. In our counterexample, the reconstruction of the root state from a subset of four taxa is more accurate than from all nine taxa in an ultrametric phylogeny (given in Figure 1). This fact is presented under the assumption...
that a character has only two states and evolves in a Jukes-Cantor model. Obviously, our finding also holds under any general evolutionary mode for a multiple-state character. Finally, we remark that the non-monotone property of ancestral state evolution could be a natural choice for ancestral state reconstruction. It has been known for a long time that the parsimony method is computationally fast, but it has limitations. There are two possible reasons for this limitation of the Fitch method. First, the Fitch method ignores character change rates on all branches. Second, the Fitch method is a kind of ‘local’ method in the sense that it estimates the states of an internal species from the states estimated at the children of the species. As such, incorrect estimates made at the internal species close to the taxa can propagate all the way through to the root state.

The Fitch method is computationally fast, but it has limitations. Hence, the weighted parsimony method proposed by Sankoff in [36] could be a natural choice for ancestral state reconstruction. It is not only computationally efficient as the Fitch method, but also takes the branch lengths into account by posing a weight on each branch.

Unlike the Fitch method, the marginal ML method does not have such an undesirable property. However, when the marginal ML method is used, reconstructing the root state from all taxa could have the same accuracy as from a small subset of taxa on a phylogeny. Additionally, the marginal ML method is not as efficient as the Fitch method. Hence, developing a fast method with monotonic reconstruction accuracy is important for future research. One promising approach for improving ancestral sequence reconstruction is to utilize other biological factors. For example, co-evolutionary information of the studied genes or proteins is employed for reconstructing the gene content of the LUCA [37].

Stability of Ancestral State Reconstruction

Our experimental results demonstrate that the accuracy of reconstructing the root state from a subset of taxa varies in a wide range especially when the size of taxon sampling is small for both the Fitch and ML methods. But the situation is much worse for the Fitch method. This is because the Fitch method computes a subset of taxa as an assignment to each internal species without graded ambiguity and hence topology can have big influence on the reconstruction procedure. This further confirms the same observation made in [24], in which different reconstruction accuracies were discussed for the Fitch method.

The accuracy variability of reconstructing the root state suggests that ancestral state reconstruction is sensitive, and the inferred ancestral state could be unreliable when a small number of taxa are used. In the reconstruction of the boreoeutherian ancestral

### Table 1. The species selected by the backward greedy method and their relative reconstruction accuracy in the reconstruction of boreoeutherian sequence.

| Species                        | Sampling size | Percentage identity |
|--------------------------------|---------------|---------------------|
| Marmoset, alpaca, armadillo, sloth | 4             | 96.18               |
| megabat                        | 5             | 97.08               |
| rabbit                         | 6             | 97.95               |
| squirrel                       | 7             | 98.39               |
| treeShrew                      | 8             | 98.77               |
| dog                            | 9             | 98.97               |
| dolphin                        | 10            | 99.18               |
| mouse_pig                      | 11            | 99.31               |
| mouse_rentor                   | 12            | 99.49               |
| horse                          | 13            | 99.62               |
| cow                            | 14            | 99.65               |
| mouse                          | 15            | 99.71               |
| bushbaby                       | 16            | 99.85               |
| tarsier                        | 17            | 99.89               |
| rat                            | 18            | 99.92               |
| kangaroo, rat                  | 19            | 99.96               |
| rhesus                         | 20            | 99.99               |
| gorilla                        | 21            | 100.00              |
| orangutan                      | 22            | 100.00              |
| human                          | 23            | 100.00              |
| chimp                          | 24            | 100.00              |

The species selected for each size include those appearing in the corresponding row or above. The third column is the percentage of base positions at which using the selected taxa and the all 24 taxa give the same nucleotide.

doi:10.1371/journal.pone.0008985.t001

### Table 2. The species selected by the forward greedy method and their relative reconstruction accuracy in the reconstruction of boreoeutherian sequence.

| Species                        | Sampling size | Percentage identity |
|--------------------------------|---------------|---------------------|
| Gorilla, orangutan, rhesus, marmoset | 4             | 90.18               |
| human                          | 5             | 90.19               |
| chimp                          | 6             | 90.19               |
| mouse_rentor                   | 7             | 92.86               |
| bushbaby                       | 8             | 93.68               |
| tarsier                        | 9             | 94.52               |
| rabbit                         | 10            | 95.50               |
| squirrel                       | 11            | 95.72               |
| treeShrew                      | 12            | 97.07               |
| mouse_pig                      | 13            | 97.24               |
| mouse                          | 14            | 97.21               |
| rat                            | 15            | 97.27               |
| kangaroo, rat                  | 16            | 97.32               |
| alpaca                         | 17            | 97.70               |
| megabat                        | 18            | 98.35               |
| dog                            | 19            | 98.46               |
| dolphin                        | 20            | 98.55               |
| horse                          | 21            | 98.61               |
| cow                            | 22            | 98.63               |
| armadillo                      | 23            | 99.67               |
| sloth                          | 24            | 100.00              |

The species selected for each size include those appearing in the corresponding row or above. In the third column, the percentage identity denotes the fraction of base positions at which using the selected taxa and the all 24 taxa give the same nucleotide.

doi:10.1371/journal.pone.0008985.t002
genome, where we used the method reported in [25], the 6 species selected by the forward selection method have a relative accuracy of 90.16%, whereas the 6 species selected by the backward selection method have relative accuracy as high as 97.95%. Therefore, caution should be exercised before drawing conclusions about evolutionary hypotheses about an ancestral sequence when the ancestral sequence was estimated from a small number of taxa.

**Taxa Selection for Ancestral State Reconstruction**

We have studied the taxon selection problem for ancestral state reconstruction under the assumption that the true phylogeny is given. In earlier studies [31] and [32], k taxa with the largest additive divergence are selected to detect single-site conservation. Such a widely-discussed criterion makes good sense [39]. Consider an ultrametric phylogeny in which substitution rate is constant. The k taxa selected based on this criterion induce a k-leaf subtree with the largest total branch length. Since each root-to-leaf path has the same length, the internal branches close to the root are short and hence in such a tree, the lineages are less dependent, giving high reconstruction accuracy. Does a subset of k taxa with the largest additive divergence give the optimal reconstruction accuracy? The answer to this question is negative in general. When the Fitch method is used, k-taxa with the largest additive divergence is not necessarily the best selection in terms of reconstruction accuracy. For instance, in the phylogeny presented in Figure 1, for k=4, taxa a,b,e,d has the largest additive divergence, but their reconstruction accuracy is lower than that of species a,b,e,i. When the marginal ML method is used, our numerical computation indicates that the k species with the largest additive divergence often give nearly-optimal reconstruction accuracy. Over an arbitrary phylogeny, the situation is much more complicated. It is not clear whether the taxa with the largest or smallest additive divergence should be selected. As such, we investigate the taxa selection problem with the algorithmic approach.

We proposed two heuristic methods for taxon selection. These methods have their origin in the linear regression study and can be applied to any reconstruction methods. We tested them for the Fitch and ML methods on random phylogenies generated under a Yule model as well as on random ultrametric trees. The experiment results show that, in most of the cases, the accuracy of reconstructing the root state from the k taxa selected by each method is comparable to the accuracy of the best reconstruction from the same number of taxa; it is also comparable to, if not better than, the accuracy of using all taxa in a phylogeny when the number of selected taxa is medium or large as shown in Figure 6. In summary, the forward selection is straightforward, but the backward selection is more effective. The C++ programs for these two methods are available upon request.

**Reconstructing Boreoeutherian Ancestor Genome**

Our experiments suggest that the reconstruction accuracy increases rapidly when taxon sampling size is small and becomes stable when 10 or more taxa are used. Hence, little is gained when a few taxa are added into or removed from the reconstruction when the number of used taxa is 10 or more. This is consistent with earlier studies [25,31]. A positive aspect of this observation is that ancestral states can be well estimated from about a dozen taxa in a phylogeny. This suggests that reconstructing an ancestral protein sequence or genome that existed millions of years ago is feasible.

Most modern mammalian lineages originated in a burst of speciation events around 80–100 million years ago [38]. This makes the boreoeutherian ancestor an ideal target for ancestral genome reconstruction. Reconstructing the boreoeutherian ancestral sequence is important for decoding the molecular basis for the extraordinary diversity of mammalian forms and capabilities. Blanchette et al. successfully reconstructed a genomic region covering about 1.1 million bases around the CFTR locus from 16 extant sequences with 96.8% accuracy at the nucleotide level including indel events, as estimated by computer simulation [25]. From the data presented in Table 1, we can see that the reconstruction of the boreoeutherian ancestor genome from a subset of 10 or more selected species has nearly-identical accuracy as from all 24 taxa for bases that are not involved in indel events. The 10 species that are most useful for the reconstruction are *marmoset, treeshrew, squirrel, rabbit, alpaca, dolphin, dog, megabast, armadillo*, and *shrew*. Unfortunately, most of these genomes are not completely sequenced in high coverage and resolution yet. With more and more species in the list being sequenced in high quality, the boreoeutherian ancestral genome should be reconstructed with high resolution in the near future.

In this paper, we focus on the taxon selection problem for inferring ancestral states. In reconstructing the boreoeutherian ancestor genomic sequence, we did not consider insertion and deletion events. Inferring the insertion and deletion events on a phylogeny is extremely challenging [40,41]. When the insertion and deletion events are considered, the definition of reconstruction accuracy presented in the method section is no longer valid. How to incorporate insertion and deletion events into the study of taxon selection is another important problem for future research in ancestral sequence reconstruction.

**Materials and Methods**

**Reconstruction Accuracy**

The problem of reconstructing ancestral states is to find the true state of a character in an ancestral species from the states in a set of extant taxa that evolve from that ancestral species. Let T be a phylogeny with root r. We use L(T) to denote the set of the extant taxa on T. For a taxon subset L⊆L(T), we say D is a state assignment for L if it associates a state with each taxon in L. We use S(L) to denote the set of all possible state assignments for L. For a state a and a state assignment D∈S(L), Pr[D|x=a] is used to denote the probability that a at the root evolves into the states specified in D. Such a probability can be calculated recursively given a Jukes-Cantor model (for example, see [42] and [43]).

Consider a reconstruction method M. The accuracy of M for reconstructing the ancestral state of a character at the root r from the states of the taxa of L is defined as the expected probability that M correctly reconstructs the root state, i.e.,

$$A_M(L) = \sum_a p_a \Pr[\hat{x}_r = a | s_r = a],$$

(1)

where \(\hat{x}_r\) denotes the state reconstructed at the root and \(p_a\) is the prior probability of state a at the root.

For a state a and a state assignment D∈S(L), we write

$$\Pr[D | \text{State is at } r \text{ States in } D \text{ are observed in the corresponding taxa}],$$

(2)

as \(Pr[\hat{x}_r = a | D]\) and call it the likelihood of the state a given state assignment D. By the law of total probability,

$$\Pr[\hat{x}_r = a | s_r = a] = \sum_{D \in S(L)} \Pr[\hat{x}_r = a | D] | s_r = a]$$

for any state a. Since the state output by M depends only on D,
In practice, one infers the root state from some state assignment. Since the inference of a state \( a \) by \( \mathcal{M} \) from a state assignment \( D \) is correct only if the states in \( D \) are evolved from \( a \). Combining Eqn (1) and Eqn (3), we obtain:

\[
A_{\mathcal{M}}(L) = \sum_{D \in S(L, a)} p_D \Pr[\tilde{\xi}_r = a | D] \Pr[\tilde{\xi}_r = a],
\]

where \( \Pr[\tilde{\xi}_r = a | D] \) is the probability that \( \mathcal{M} \) outputs \( a \) as the root state on \( D \).

As a parsimony reconstruction method, the Fitch method assigns to each internal node those states that allow for the smallest number of substitutions posed on all branches of a phylogeny over the given taxa [6]. The assignment to each node is computed by considering the assignments previously obtained at the node’s children one by one downward in the phylogeny, starting with the taxa. For each taxon, the observed state forms the state subset. Assume \( A \) is an internal node with children \( B \) and \( C \). The following rule is used to compute the state subset \( S_A \) from the state subsets \( S_B \) and \( S_C \):

\[
S_A = \begin{cases} 
S_B \cup S_C & \text{if } S_B \cap S_C = \emptyset, \\
S_B \cap S_C & \text{if } S_B \cap S_C \neq \emptyset.
\end{cases}
\]

The state subset output by the Fitch method at the root contains all the possible states that are equally parsimonious candidates as the root state. We say that the Fitch method unambiguously reconstructs a state \( a \) at the root \( r \) if the state

\[
\Pr[\tilde{\xi}_r = a | D | \tilde{\xi}_r = a] = \frac{\Pr[\tilde{\xi}_r = a | D]}{\Pr[\tilde{\xi}_r = a]} = \frac{\Pr[D]}{\Pr[\tilde{\xi}_r = a]}.
\]

In each figure, the 1000 tree samples are arranged in ascending order of the accuracy of reconstructing the root state from the best subset of 12 species. (a) and (b) The conservation probability \( p \) on each branch is set to be 85% and the method is the forward and backward greedy method respectively. (c) and (d) \( p = 95\% \) and the method is the forward and backward greedy method respectively.

doi:10.1371/journal.pone.0008985.g006

**Figure 6.** Comparison of the accuracies of reconstructing the root state of a character from the 12 taxa selected by a greedy approach, the best subset of 12 taxa and all 16 taxa, respectively, on a random phylogeny. The graphs are drawn using the accuracy data collected on 1000 random trees generated in a Yule model. In each figure, the 1000 tree samples are arranged in ascending order of the accuracy of reconstructing the root state from the best subset of 12 species. (a) and (b) The conservation probability \( p \) on each branch is set to be 85% and the method is the forward and backward greedy method respectively. (c) and (d) \( p = 95\% \) and the method is the forward and backward greedy method respectively.
subset $S_r$ contains only a and ambiguously reconstructs a state $a$ if $S_r$ contains $a$ and other states. When $S_r$ contains more than one state, we simply pick one of them as the root state. Thus, for a state assignment $D$ of a taxon subset in the given phylogeny, the probability that the Fitch method outputs a state $a$ from $D$ is set to be

$$\Pr[\hat{s}_r = a | D] = \begin{cases} 0 & a \not\in S_r, \\ 1 & a \in S_r \end{cases}$$

(5)

where $|S_r|$ is the number of states in the subset $S_r$ computed from $D$.

Given a phylogeny and a Jukes-Cantor model, the accuracy of reconstructing the root state from a subset of leaf states in a phylogeny can be calculated by using a recurrence system (see [42] or [43]).

To reconstruct the root state more accurately, the marginal maximum likelihood (ML) method selects the state that has the maximum likelihood given $D$, which is defined in Eqn (2), breaking tie by choosing one arbitrarily. For the marginal ML method, we have that

$$\Pr[\hat{s}_r = a | D] = \begin{cases} \frac{1}{k} & \Pr[\hat{s}_r = a | D] \text{ is maximal,} \\ 0 & \text{otherwise} \end{cases}$$

(6)

where $k$ is the number of states that have the same likelihood as $a$. By Eqn (4),

$$A_{M\ell}(L) = \sum_{D \in S(L)} \left( \sum_a p_a \Pr[D|\hat{s}_r = a] \Pr[\hat{s}_r = a | D] \right)$$

$$= \sum_{D \in S(L)} \max_a \{ p_a \Pr[D|\hat{s}_r = a] \}$$

(7)

For any reconstruction method $\mathcal{M}$, by Eqn (4), its accuracy of reconstructing the root state from a taxon subset $L$ in a phylogeny is bounded as

$$A_{\mathcal{M}}(L) = \sum_{D \in S(L)} \sum_a p_a \Pr[\hat{s}_r = a | D] \Pr[D|\hat{s}_r = a]$$

$$\leq \sum_{D \in S(L)} \max_a \{ p_a \Pr[D|\hat{s}_r = a] \} \left( \sum_a \Pr[\hat{s}_r = a | D] \right)$$

$$= \sum_{D \in S(L)} \max_a \{ p_a \Pr[D|\hat{s}_r = a] \}.$$
References

1. Pauling L, Zuckerkandl E (1965) Chemical paleogenetics: molecular restoration studies of extinct forms of life. Acta Chem Scand 17: 89–916.
2. Thornton JW (2004) Resurrecting ancient genes: experimental analysis of extinct molecules. Nature Rev Genetics 5: 366–375.
3. Liberles DA, ed (2007) Ancestral sequence reconstruction. New York: Oxford University Press.
4. Crip MD, Cook LG (2005) Do early branching lineages signify ancestral traits? Trends in Ecol Evol 20: 122–128.
5. Felsenstein J (2004) Inferring Phylogenies. Sunderland: Sinauer Associates, Inc.
6. Finch WM (1971) Toward Defining the Course of Evolution: Minimum Change for a Specific Tree Topology. Syst Zoology 20: 406–416.
7. Baba ML, Goodman M, Berger-Cohn J, Demaille, Matsuda G (1984) The early adaptive evolution of calmodulin. Mol Biol Evol 1: 442–455.
8. Jermann TM, Opitz JG, Stackhouse J, Benner SA (1995) Reconstructing the evolutionary history of the ariodacyl ribonuclease superfamily. Nature 374: 57–59.
9. Chandrasekharan UM, Sanker S, Glynias MJ, Kamik SS, Husain A (1996) Angiotensin II forming activity in a reconstructed ancestral chymase. Science 271: 502–505.
10. Zhang J, Rosenberg HF (2002) Complementary advantageous substitutions in the evolution of an antiviral RNAse of higher primates. Proc Natl Acad Sci USA 99: 5486–5491.
11. Cunningham CW, Onland KE, Oakley TH (1998) Reconstructing ancestral character states: A critical reappraisal. Trends in Ecol and Evol 13: 361–366.
12. Yang ZH, Kumar S, Nei M (1995) A new method of inference of ancestral nucleotide and amino acid sequences. Genetics 141: 1641–1650.
13. Koshi JM, Goldstein RA (1996) Probabilistic Reconstruction of Ancestral Protein Sequences. J Mol Biol 42: 313–320.
14. Papko T, Pe'er I, Shamir R, Graur D (2000) A fast algorithm for joint reconstruction of ancestral amino acid sequences. Mol Biol Evol 17: 890–896.
15. Elias T, Tuller T (2007) Reconstruction of ancestral genomic sequence using likelihood. J Comput Biol 14: 216–237.
16. Pagel M (1999) The maximum likelihood approach to reconstructing ancestral character states. Syst Biol 48: 612–622.
17. Chang BS, Jonsson K, Kazmi MA, Donoghue MJ, Sakmar TP (2002) Recreating a functional ancestral archosaur visual pigment. Mol Biol Evol 19: 1403–1409.
18. Thornton JW, Need E, Crews D (2003) Resurrecting the ancestral steroid receptor: ancient origin of estrogen signalling. Science 301: 1714–1717.
19. Gaucher EA, Thomson JM, Burgan MF, Benner SA (2005) Inferring the palaeoenvironment of ancient bacteria on the basis of resurrected proteins. Nature 432: 203–208.
20. Kuang DH, Yao Y, MacLean D, Wang MH, Hampson DR, et al. (2006) Ancestral reconstruction of the ligand-binding pocket of Family CG protein-coupled receptors. Proc Natl Acad Sci USA 103: 14050–14055.
21. Huchardbeac JP, Rollback JP (2001) Empirical and hierarchical Bayesian estimation of ancestral states. Syst Biol 50: 351–366.
22. Hall BG (2006) Simple and accurate estimation of ancestral protein sequences. Proc Natl Acad Sci USA 103: 5431–5436.
23. Zhang J, Nei M (1997) Accuracies of ancestral amino acid sequences inferred by parsimony, likelihood, and distance methods. J Mol Evol 44(S1): 139–146.
24. Salisbury BA, Kim J (2001) Ancestry state estimation and taxon sampling density. Syst Biol 50: 357–364.
25. Blanchette M, Green ED, Miller W, Haussler D (2004) Reconstructing large regions of an ancestral mammalian genome in silico. Genome Res 14: 2412–2423.
26. Pardi F, Goldman N (2005) Species choice for comparative genomics: Being greedy works. PLOS Genetics 6: e71. Available: http://www.plosgenetics.org/article/info%3Adoi%2F10.1371%2Fjournal.pgen.0010071.
27. Manderson WP (1995) Calculating the probability distributions of ancestral states reconstructed by parsimony on phylogenetic trees. Syst Biol 44: 474–481.
28. Kim J (1996) General inconsistency conditions for maximum parsimony: Effects of branch lengths and increasing numbers of taxa. Syst Biol 45: 563–574.
29. Blanchette M, Kent WJ, Riemer C, Elnitski L, Maitra A, et al. (2004) Aligning multiple genomic sequences with the Threaded Blockset Aligner. Genome Res 14: 707–715.
30. Kent WJ, Sugnet CW, Furey TS, Roskin KM, Pringle TH, et al. (2002) The human genome browser at UCSC. Genome Res 12: 986–1006.