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RH: Missing data in Total Evidence matrices

Effect of missing data in matrices containing living and fossil taxa on topological accuracy

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Abstract.— Living species represent less than 1% of all species that have ever lived. Ignoring fossil taxa may lead to misinterpretation of macroevolutionary patterns and processes such as trends in species richness, biogeographical history or paleoecology. This fact has led to an increasing consensus among scientists that both fossil and living taxa must be included in macroevolutionary studies. One approach, the total evidence approach, uses molecular data from living taxa and morphological data from both living and fossil taxa to infer phylogenies with both fossil and living taxa at the tips. Although the total evidence approach seems very promising, it requires a lot of data and is therefore likely to suffer from missing data issues which may affect its ability to infer correct phylogenies.

In this study we assess the effect of missing data on tree topologies inferred from total evidence matrices. Using simulations we investigate three major factors that directly affect the completeness of the morphological part of the matrix: (1) the proportion of living taxa with no morphological data, (2) the amount of missing data in the fossil record and (3) the overall number of morphological characters in the matrix. We find that, in a Bayesian framework, difficulties in recovering a stable topology are mainly driven by the missing data in the molecular part of the matrix (for which fossil taxa have no data). In a Maximum Likelihood framework, however, topology is not directly affected by missing data per se, but by the number of morphological characters shared among the taxa. Therefore, the two main drivers of incorrect topologies are the overall number of morphological characters and the number of living species with no morphological data.

Our results suggest that, in order to use total evidence approaches, one should reduce the missing data in the morphological part of the matrix for living species and use a Maximum Likelihood framework to fix the topology prior to the overall Bayesian phylogenetic inference process. We apply our method to a comprehensive data set of both living and fossil primates. We find that using this integrative method modifies previous estimates of rates of body mass evolution within primates.

(Keywords: missing data, total evidence, bla)

Living species represent less than 1% of all species that have ever lived (Novacek and Wheeler 1992; Raup 1993). However, the majority of macroevolutionary studies focus solely on living species (Cooper and Purvis 2009; Meredith et al. 2011; Healy et al. 2014). Ignoring fossil taxa may lead to misinterpretation of macroevolutionary patterns and processes such as timing of diversification events (e.g. Pyron 2011), relationships among lineages (e.g. Manos et al. 2007) or niche occupancy (e.g. Pearman et al. 2008). These factors have led to increasing consensus among scientists that fossil taxa must be included in macroevolutionary studies (Jackson and Erwin 2006; Quental and Marshall 2010; Dietl and Flessa 2011; Slater and Harmon 2013; Fritz et al. 2013). However, to do this we need to be able to place living and fossil taxa into the same phylogenies which still remains complex despite an increasing number of significant efforts (Pyron 2011; Ronquist et al. 2012a; Schrago et al. 2013).

Three main approaches have been used for combining fossil and living taxa data in phylogenies. These approaches differ mainly in whether they treat fossil taxa as tips or as nodes in the phylogeny and which part of the available data of the fossil is used (i.e. the fossil occurrence age only or both the age and the morphology). Classical cladistic methods uses matrices containing morphological data from both fossil and living taxa and allows to treat each taxon as a tip in the phylogeny; the relation between the taxa can be infered using optimal criteria such as the maximum parsimony criterion (Simpson 1945). This approach is commonly used by paleontologists but it ignores the additional molecular data available from living species. Neontologists, on the other hand, more commonly use probabilistic methods (e.g. Maximum Likelihood or Bayesian) based on matrices containing only molecular data from living species to infer phylogenies. Because fossil taxa do not usually have available DNA, fossils are used as nodes rather than tips in these analyses and only their occurrence age can be used to time calibrate phylogenies (Zuckerkandl and Pauling 1965). There have been great improvements in the theory and

application of these two approaches (e.g. Bapst 2013; Stadler and Yang 2013; Heath et al. 2013) as well as much debate about the "best" approach to use (e.g. Spencer and Wilberg 2013) but none of them uses all the available data.

A final approach, known as total evidence approach use matrices containing molecular data from living taxa and morphological data from both living and fossil taxa. It allows to use probabilistic methods, to treat every taxon as a tip on the phylogeny and to use the occurrence age of the fossil to time calibrate the phylogeny (Eernisse and Kluge 1993). Here we focus on this total evidence approach because they have been recently successfully developed and applied to empirical data (Pyron 2011; Ronquist et al. 2012a; Schrago et al. 2013). Although the total evidence approach seems very promising, there is one big drawback in using this approach: it requires a lot of data. In particular it requires morphological data form both living and fossil taxa, both of which are scarce. Therefore total evidence approaches are likely to suffer from having lots of missing data which may affect their ability to infer correct phylogenies.

The effect of missing data on phylogenetic inferences has been widely studied (Wiens 2003, 2006; Wiens and Moen 2008; Lemmon et al. 2009; Roure and Philippe 2011; Sansom and Wills 2013). Missing molecular data has been seen by some authors as an issue because it can decrease the phylogenetic signal in some parts of the tree, especially when using large matrices (Lemmon et al. 2009). However other authors do not see missing molecular data as a major issue because the phylogenetic signal is more likely to increase by having at least a "modest" number of highly covered genes (i.e. approximatively half of the genes - Roure and Philippe (2011)), a higher number of taxa (especially slowly evolving taxa or taxa close to the outgroup) and by choosing more adequate models of sequence evolution rather than by reducing the amount of missing data (Wiens 2006; Wiens and Moen 2008; Roure and Philippe 2011). Similarly, depending on the study, missing morphological data might be seen as either a major or minor issue for accurately inferring

phylogenies depending on the study (Wiens 2003; Sansom and Wills 2013). Because soft-tissues characters are rarely preserved in the fossil record, missing data is mainly found in soft tissues characters, and is therefore not randomly distributed, which can lead to biased placement of fossil taxa in phylogenies (Sansom and Wills 2013). However, the phylogenetic signal is not related to the level of missing data per se but to the number of informative characters per taxa, therefore missing data is less an issue than the number of shared informative characters (Wiens 2003). Although missing data as been shown to be no major problem separately in molecular and morphological matrices (Wiens 2003, 2006; Wiens and Moen 2008; Roure and Philippe 2011) it may become an issue when combined in a total evidence type matrix and no attempt has been made to study the impact of this issue on phylogenetic inference until now (especially because a major part of the matrix is missing - the molecular data from the fossil taxa).

Here we assess the effect of missing data on tree topology inferred from total evidence matrices. The molecular part of a total evidence matrix contains no fossil taxa so will act like a classical molecular matrix (Ronquist et al. 2012a). The effect of missing data on such matrices is well known, therefore, we only focus on the missing data issue in the morphological part of the matrix. Using simulations we investigate three major factors that directly affect the completeness of the morphological part of the matrix: (1) the proportion of living taxa with no morphological data, (2) the amount of missing data in the fossil taxa (i.e. the preservation quality of the fossil record) and (3) the overall number of morphological characters for both living and fossil taxa in the matrix. We assess how changing the values of these three parameters affects the topology of total evidence approach phylogenies.

METHODS

To explore the effect of missing data on total evidence trees topologies we used the following

protocol (note that we explain each step in detail below this general outline -Fig. 1):

- Generating the matrix We built a randomly generated birth death tree (hereafter called the "best" tree, Table 1) to infer a matrix containing both molecular and morphological data for living and fossil taxa (hereafter called the "complete" matrix, Table 1).
- 2. Removing data We removed data from the "complete" matrix to simulate the effects of missing data by modifying three parameters (1) the proportion of missing living taxa (M_L) , (2) the proportion of missing data in the fossil taxa (M_F) and (3) the proportion of missing morphological characters (M_C) (the resulting matrix is called hereafter the "missing-data" matrix, Table 1).
- 3. Building phylogenies We inferred Bayesian phylogenetic trees from the "complete" matrix and from the "missing-data" matrices resulting in respectively a tree generate from a matrix containing no missing data in the morphological part (hereafter called the "best" tree, Table 1) and trees inferred from matrices with missing data in their morphological part (hereafter called the "missing-data" trees, Table 1).
- 4. Comparing topologies We then compared the "best" tree to the "missing-data" trees to assess the influence of each parameter $(M_L, M_F, M_C \text{ and their interactions})$ on the topologies of the phylogenies we estimated.

To measure the effect of missing data distribution, we repeated steps 1 to 4 with the exact same fixed parameters 50 times. A list of all the terms used in this paper is available in table 1.

Generating the matrix

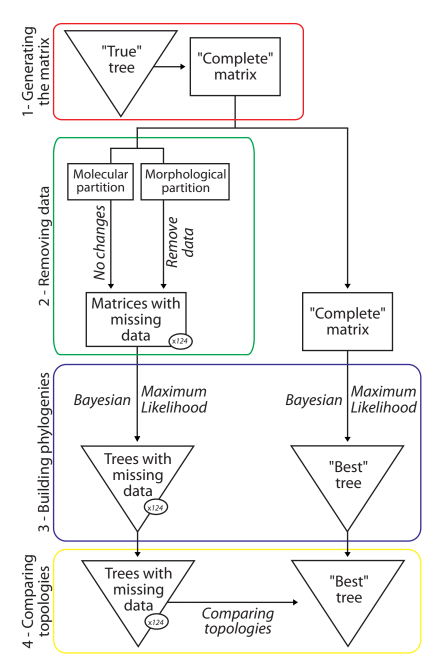


Figure 1: Protocol outline. (1) We generated a random tree (hereafter called the "true" tree) to infer a matrix with no missing data (hereafter, the "complete" matrix). (2) We removed data fron the morphological partition of the "Complete" matrix resulting in 124 matrices with missing data. (3) We build Bayesian phylogenetic trees from each matrix. (4) We then compared the trees with missing data (from the matrices with missing data) to the tree with no missing data (hereafter, the "best" tree infered from the "complete" matrix). We repeated step 1 to 4 50 times.

First we randomly generated a "true" tree of 50 taxa in R v3.0.2 (R Core Team 2014) using the package diversitree v0.9-6 (FitzJohn 2012). We generated the tree using a Birth Death process by sampling the values of the speciation events (λ) and extinction events (μ) from a uniform distribution but maintaining $\lambda > \mu$ (Paradis 2011). We implemented a rejection sampling algorithm to select only random trees with 25 living and 25 fossil taxa. We then added a species to the resulting Birth-Death tree as the outgroup of the tree. The mean branch length of the tree was used to separate the outgroup from the rest of the taxa and the branch length leading to the outgroup was set as the sum of the mean branch length and the longest root-to-tip length of the tree.

Next, we created a molecular and a morphological matrix from the "true" tree. The molecular matrix was infered from the "true" tree using the package phyclust v0.1-14 (Chen 2011). The matrix was made of 1000 characters sites for 51 taxa and generated using the seqgen algorithm (Rambaut and Grassly 1997). We used the HKY model (Hasegawa et al. 1985) with a random base frequencies and with the transition/transversion rate of 2 (Douady et al. 2003) as parameters for generating the matrix. The substitution rates were distributed following a gamma distribution with an alpha (α) shape of 0.5 (Yang 1996). We chose a low value of α to lower the number of sites with high substitution rates, thus avoiding too much homoplasy and a decrease in phylogenetic signal. These parameters were selected to generate data with no special assumption about how the characters evolved as well as to reduce the computational time required if these parameters were estimated rather than defined (65 CPU years).

We inferred the morphological matrix using the ape package v3.0-8 (Paradis et al. 2004) to generate a matrix of 100 character sites for 51 taxa. We assigned the number of character states (either 2 or 3) for each morphological character by sampling with a probability of 0.85 for two states characters and 0.15 for three state characters. These probabilities were selected using the overall distribution of characters states extracted from

100 published empirical morphological matrices (See supplementaries). We then ran an independent discrete character simulation for each character using the "true" tree branch length and topology with the randomly selected number of states (2 or 3) and assuming an equal rate of change (i.e. evolutionary rate) from one character state to an other (Pagel 1994). This method allows us to have only two parameters per character i.e. the number of states and the evolutionary rate. For each character, the evolutionary rate was sampled from a gamma distribution with $\alpha = 0.5$. All the molecular information for fossil taxa was replaced by missing data ("?"). Finally, we combined the morphological and molecular matrices obtained from the "true" tree. Hereafter we call this the "complete" matrix, i.e. the matrix with no missing data except for the molecular data of the fossil taxa.

Removing data

Once we obtained the "complete" matrix we modified it to get a set of matrices with missing data We randomly replaced data with "?" in the morphological part of the matrices according to the following parameters (Fig. 2):

- 1. The proportion of living taxa with no morphological data (M_L) : 0%, 10%, 25%, 50% or 75%. This parameter illustrates the number of living taxa that are present in the molecular part of the matrix but not in the morphological one. Because of the increasing facility to sequence DNA for living species, the number of living species with molecular data is highly superior the the number of species with molecular and morphological data.
- 2. The proportion of missing morphological data across all fossil taxa (M_F) : 0%, 10%, 25%, 50% or 75%. This parameter illustrates the quality of the fossil record.
- 3. The proportion of missing morphological characters across all taxa (living and fossil -

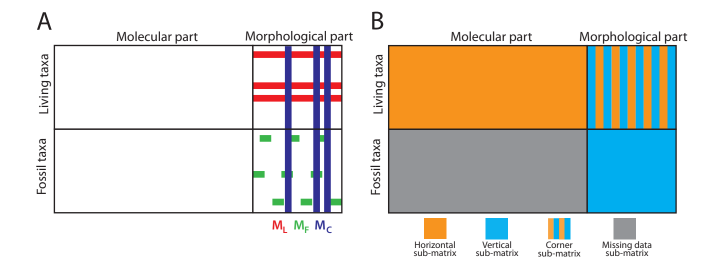


Figure 2: Different parts of the matrix. A: Missing data parameters: Missing living - The proportion of living taxa with no morphological data (M_L) ; Missing fossil - The proportion of missing morphological data across all fossil taxa (M_F) ; Missing character - The proportion of missing morphological characters across all taxa (living and fossil) (M_C) . B: Different parts of the matrix: The "horizontal" sub-matrix (orange) contains molecular and morphological data for living taxa; The "vertical" sub-matrix (blue) contains morphological data for living and fossil taxa; The "corner" sub-matrix (orange and blue striped) contains morphological data for living taxa; The "missing-data" sub-matrix (grey) is the molecular part of the fossil taxa and contains no data.

 M_C): 0%, 10%, 25%, 50% or 75%. This parameter illustrates the number of available morphological characters for both living and fossil taxa.

In practice, each parameter represent a way of removing data in the morphological part of the matrix: M_L removes a proportion of rows from the living taxa; M_F removes a proportion of cells from the fossil taxa; and M_C removes a proportion of columns across both living and fossil taxa. Note that M_L is different to M_F not only because of the region of the matrix affected: for M_L , all the morphological data of a proportion of the living taxa is removed (i.e. removing rows), as for M_F , a proportion of data is removed across the whole of the morphological matrix for fossil taxa (i.e. removing cells).

We tested all parameters combinations resulting in 125 (5³) matrices. Because some parameter combinations introduce a lot of missing (e.g. M_L =75%, M_F =75% and M_C =75%), some matrices contained fossil taxa without any data at all. When this occurred we repeated the random deletion of characters until every species had at least 5% data.

Building phylogenies

From the resulting matrices we generated two types of trees, the "best" tree that is inferred from the "complete" matrix and the "missing-data" trees inferred from the 125 matrices with various amounts of missing data. The "true" tree was used to generate the "complete" matrix and reflects the "true" evolutionary history in our simulations. The "best" tree, on the other hand, is the best tree we can build using the state-of-the-art phylogenetic methods. In real world situations, the "true" tree is never available to us because we cannot know the true evolutionnary history of a clade (except in very rare circumstances, e.g. Rozen et al. (2005)). Therefore, here we focus on comparing the trees inferred from the matrices with missing data to the "best" tree, rather than the "true" tree, as the "best" tree is generally what biologists have to work with.

Maximum Likelihood.— The "best" tree and the "missing-data" trees were infered using RAxML v8.0.20 (Stamatakis 2014). For the molecular data, we used the GTR + Γ_4 model (Tavaré (1986); default GTRGAMMA in RAxML v8.0.20; Stamatakis (2014)) as a generalisation of the HKY + Γ_4 model (Hasegawa et al. 1985) for the molecular data. The GTR model can be seen as a generalisation of the HKY model (the 2 parameters from the HKY model are implicitly included in the 6 from GTR model - Stamatakis et al. (2008)). For the morphological data, we used the implemented Markov k state model (Lewis 2001) wich is a generalisation of the JC69 model (Jukes and Cantor 1969) with $k \geq 2$ assuming

an equal state frequency and a unique overall substitution rate (μ) following a gamma distribution of the rate variation with four distinct categories (M $k + \Gamma_4$; -K MK option in RAxML v8.0.20; Stamatakis (2014)) We used the fast bootstrap algorithm and performed 1000 bootstraps per tree inference to assess the topological support. The bootstrap algorithm used in RAxML is the Lazy Subtree Rearrangement (LSR) which consists in pruning one subtree from the tree and subsequently reinserting it to all neighboring branches (Stamatakis et al. 2008). Subtree Pruning and Reinserting methods (SPR) have been demonstrated as being better than others (e.g. Nearest Neighbouring Interchange - NNI) in recovering good bootstrap values (Salamin et al. 2003).

Bayesian.— The "best" tree and the "missing-data" trees were infered using MrBayes v3.2.2 (Ronquist et al. 2012b). We partitioned the data treating the molecular part as a non-codon DNA partition and the morphological part as a multi-state morphological partition. The molecular evolutionary history was infered using the HKY model with a transition/transversion ratio (Douady et al. 2003) of two and a gamma distribution for the rate variation with four distinct categories (HKY + Γ_4). For the morphological data, we used the Markov k state model (Lewis 2001), with equal state frequency and a unique overall substitution rate (μ) with four distinct rates categories (M $k + \Gamma_4$). We chose these models to be consistent with the parameters used to generate the "complete" matrix.

Each Bayesian tree was estimated using two runs of four chains each for a maximum of 50×1^6 generations. We used the average standard deviation of split frequencies (ASDS) as a proxy to estimate the convergence of the chains and implemented a stop rule when the ASDS went below 0.01 (Ronquist et al. 2012b). The effective sample size (ESS) was also checked on a random subsample of runs in each simulation to check that ESS >> 200 (Drummond et al. 2006). For each run, we removed 25% of the iterations as burnin. We used the following priors for each tree (see supplementaries):

- 1. the "true" trees topology as a starting tree (with a starting value for each branch length of 1),
- 2. an exponential prior on the shape of the gamma distribution of α =0.5 for both partitions
- 3. and a transition/transversion ratio prior of 2 sampled from a strong beta distribution $(\beta(80,40))$.

We used these priors to speed up the Bayesian process. These priors biased the way the Bayesian process calculated the branch length by giving non-random starting points and boundaries for the parameters estimation process, however, we are focusing on the effect of missing data on the topology and not on the branch length. Even using these priors, it took 65 CPU years to build 50 sets of 125 Bayesian trees (8 core nodes 2.30GHz clock speed).

Comparing topologies

We compared the topology of the "missing-data" trees infered from the matrices with missing data to the "best" tree to measure the effect of the three parameters M_L , M_F and M_C . Note that we only investigate differences in topology and not in branch length because the aim of this study is to look at the effect of missing data on the topology of trees infered from matrices with living and fossil taxa and molecular and morphological characters. To compare the topology of the resulting trees, we used two metrics to assess number of conserved taxa and clades position using respectively the Triple (Dobson 1975) and the Robison-Fould (Robinson and Foulds 1981) distance. We normalised the two metrics using the Normalised Tree Similarity index (Bogdanowicz et al. 2012) to generalise our results for any n number of taxa. The two metrics and the index are detailed below.

Triple distance $(T_{x,y})$ (Dobson 1975).— This metric measures the number of different subtrees of three clades between two given trees. Each triplet can be written as $I_{ijk} = (ijk)$. Where I_{ijk} is equal to 0 if the two triplets (ijk) are the same in the two trees otherwise I_{ijk} is equal to 1. For any rooted binary tree there are only three possible combinations per triplets: ((j,k),i);, ((i,k),j); and ((i,j),k); (Johnson and Soltis 1998). If the trees used are not fully binary, a fourth triplet combination is possible: (i,j,k);. One can calculate S_n , the triplet distance between two trees as:

$$S_n = \sum_{ijk} I_{ijk} \tag{1}$$

Where:

$$\sum_{ijk} = \binom{n}{4} = \frac{n!}{4!(n-4)!} \tag{2}$$

And where n is the number of taxa in both trees (modified from Critchlow et al. (1996)). If $S_n=0$, the trees are the same (i.e. no taxa as been displaced). When $S_n=\binom{n}{4}$, the trees are the most different possible (i.e. every taxa as been displaced). This metric therefore illustrates the amount of displaxed taxa but is less sensitive to the placement of individual taxa and to taxa of highly uncertain placement (e.g. fossil taxa) than the Robinson-Foulds distance (Critchlow et al. 1996; Johnson and Soltis 1998; Wiens 2003) and can therefore be used as a proxy to estimate the robustness of the tree to flying taxa (see supplementaries).

Robinson-Fould distance (RF) (Robinson and Foulds 1981).— This metric measures the number of shared clades among two trees and therefore illustrates the number of exactly conserved groups among the trees. The Robison-Fould distance (also called path difference) between two trees reflects the distance between the distributions of the tips among clades

in the two trees (Robinson and Foulds 1981) and can be expressed as following:

$$RF_{x,y} = N_x + N_y - 2C_{x,y}$$
 (3)

Where $C_{x,y}$ is the number of clades in common in the two trees. The minimal value of C is 1 if the two trees have the same n taxa; the maximal value in C=n-2. This metric is more sensitive to taxa displacement than the triple metric (if one taxa gets out of a clade, then the clades are no longer considered as similar - Critchlow et al. (1996); Johnson and Soltis (1998); Wiens (2003)) and therefore a low value will show a good clade conservation between two trees and a high value will show a bad recovery of common clades (see supplementaries).

Normalised Tree Similarity NTS (Bogdanowicz et al. 2012).— For any tree with n taxa compared using a tree distance metric m, NTS_m represents the similarity score between the two trees given the expected distance between two random Yule trees with n taxa. Let $\bar{d}_{m,n}(rand)$ be the average distance between two random Yule trees with n taxa and $d_{m,n}(x,y)$ the distance between the two trees x and y containing each n taxa, then:

$$NTS_{m,n}(x,y) = \frac{\bar{d}_{m,n}(rand) - d_{m,n}(x,y)}{\bar{d}_{m,n}(rand)}$$

$$\tag{4}$$

NTS ranges from 1 to $-\infty$. For any m,n when NTS=1, the trees are the same; when NTS=0 the trees are not more different than expected by chance; when NTS<0, the trees are more different than expected by chance (see supplementaries).

We compared the "missing-data" tree to the "True" and the "Best" tree for each chain. For the Maximum Likelihood trees we performed pairwise comparisons between the

"True" and "Best" tree and the "missing-data" tree (where the "missing-data" tree is one of the 125 trees resulting from the 125 super matrices including various amount of missing data) for both the Robinson-Fould and the Triple metric. We calculated the difference between the trees using the metrics described above by using the TreeCmp java script (Bogdanowicz et al. 2012). For each metric, we normalized the value using the Normalised Tree Similarity scaled with the mean value of 1000 pairwise random tree comparison for the same metric and the same number of taxa n=51 (see supplementaries). We ran the comparison for every "missing-data" tree being one of the 125 combination of parameters values M_L , M_F and M_C in each chain resulting in 50 comparisons for every "missing-data" trees. We calculated the mode and the 50% and 95% confidence intervals from the resulting distribution using the hdrcde R package (v3.1 with contributions from Jochen Einbeck and Wand (2013)).

For the Bayesian trees, we had to transform the trees' posterior distribution to a single value in order to be able to compare it to the results of the Maximum Likelihood trees comparisons. Traditionally people use the 50% consensus rule in order to summarise a Bayesian tree posterior distribution (e.g. Ronquist et al. 2012b). However, when the Bayesian inference fails to recover a stable tree topology, the resulting consensus tree may contain many polytomies up to the maximum (resulting in a star tree, i.e. no resolution at all). Because one cannot compare trees with no resolution we didn't use the consensus tree but instead we used the mode of 1000 random pairwise comparisons within the Bayesian trees posterior distribution. In other words, we summarized the difference between two Bayesian tree posterior distributions using the mode of the distribution of 1000 random trees drawn from the first Bayesian trees posterior distribution to 1000 random trees drawn from the second Bayesian trees posterior distribution. We used the mode to summarize the distributions because it reflects more accurately what a consensus tree represents (the topology the most represented in the posterior distribution). A matrix with a good

phylogenetic signal will lead to a posterior trees distribution containing many identical topologies. Using a normalised tree similarity metric to compare the trees within such a posterior distribution will lead to a high value of tree similarity, reflecting reality.

Alternatively, a matrix with a poor phylogenetic signal will lead to a posterior distribution containing many different topologies. Using a normalised tree similarity metric to compare the trees within such a posterior distribution will lead to a low value of tree similarity, reflecting reality. We compared the pairs of Bayesian tree posterior distributions in the same way as for the Maximum Likelihood trees ("True" and "Best" tree compared to the tree "missing-data" tree). This resulted in a distribution of one mode per chain for each "missing-data" trees (i.e. 50 modes) for which we calculated the mode and the 50% and 95% confidence intervals (see supplementaries - Fig. 3).

Effect of missing molecular characters for fossil taxa

To assess the effect of missing molecular characters for fossil taxa in a Bayesian framework, we split the "complete" matrix in sub-matrices containing no missing data at all:

- 1. A first containing both molecular and morphological data for living taxa only (hereafter called the "horizontal" sub-matrix, Table 1);
- 2. A second one containing morphological data for both living and fossil taxa (hereafter called the "vertical" sub-matrix, Table 1);
- 3. A third one containing morphological data for living taxa only (hereafter called the "corner" sub-matrix, Table 1);

We reran the Bayesian tree inference on the different sub-matrices with no missing data in the same way as described above. We then compared the resulting tree posterior distribution to itself (in the same way described above) to assess the ability to recover

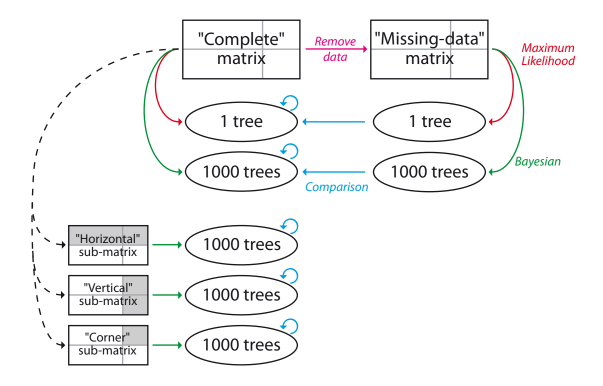


Figure 3: Tree comparison protocol. From each of the 125 parameters combination $(M_L, M_F \text{ and } M_C)$, we inferred a tree in Baysian or Maximum likelihood framework. We then compared this tree with or without fossil/living taxa to the "best/true" tree. In the Bayesian framework, 1000 trees from the posterior distribution where used for each comparison.

topology for each simulation when no missing data was involved in the phylogenetic inference process.

Empirical data

We also compared the results obtained from simulated data by using Ronquist et al. (2012a) empirical data. The matrix contains 67 living species plus one outgroup and 45 fossil species of Hymenopteras with 5097 molecular characters and 354 morphological characters. From the 68 living species used in the matrix, only 66 had molecular data, we therefore treated these 66 taxa as "living" taxa and all the other 47 as "fossil" taxa. We treated the matrix in the exact same way as described in step 2 and 3 resulting in 125 matrices with various amount of missing data and the same number of Maximum Likelihood and Bayesian trees. We used the same settings as for the simulated data in the Maximum Likelihood framework. For the Bayesian inferences however, we didnt used any priors except that we provided a starting tree with the topology of the 68 living species (topology with the highest posterior probability from non-clock analysis - Ronquist et al. (2012a)). Contrary to Ronquist et al. (2012a) analysis, we didnt performed any clock analysis since we were only interested in the topology of the inferred tree and not the branch length.

RESULTS

The ability of recovering the best topology in maximum likelihood method is function of the amount of data missing. However, in Bayesian, topology is badly (if not at all) recovered independently of the amount of data missing.

DISCUSSION

What about the "best" trees with fairly bad bootstrap support values in ML? This is probably due to the low number of characters used in the simulation. The number of morphological characters (10²) is in the same order of magnitudes of empirical studies (cite) but decreases unrealistically with the M_C parameter. The number of molecular characters (10³) is at least two orders of magnitude lower than in empirical studies that can reach 10⁶ characters (cite).

Two different matrices from the "True" tree gives two different phylogenetic interpretations (i.e. seqGen and ape performs badly at generating a data matrix from a tree)? That seems to not really be a problem since we are interested in recovering topology with missing data, not assessing if phylogenetic matrices generation software performs correctly. Also, the "noise" that might be created by this potential "bad" phylogenetic matrices generation software is probably diminished by the number of replicates (i.e. 50 chains).

Different phylogenetic signal between the morphological and the molecular matrix? That is not really a problem since the phenotypic evolution is not necessary correlated to the genotypic one. Also it is important to note that both information are not exactly the same. The main difference between molecular and morphological characters is that, because of historical and practical reasons, a morphological character matrix do no contains autapomorphies and invariant characters which are usually more common than synampomorphies in molecular character matrices (cite).

What about homoplasy in the morphological matrix? Here, we made the assumption that theoretically, morphological characters are randomly distributed on an organism however it seems clear that empirical morphological data does not act randomly If one states that morphological characters accumulate through time in the same way as the majority of the molecular characters (neutral theory, cite) then, homoplasic characters are expected to appear randomly through time. Therefore, homoplasy is expected to be

more important (by chance) in bigger morphological matrices (Davalos et al. 2014).

CONCLUSION

A Bayesian approach fails to recover accurate topology in a total evidence approach framework, whatever the amount of missing data. We think this failure is due to the intrinsic structure of the Total Evidence matrices that includes a vast amount of not randomly distributed missing data (i.e. the molecular part of the matrix for the fossil taxa). This missing data is equal to the number of fossil taxa × the number of molecular characters leading to a vast amount of likely trees to be sampled in the Bayesian mcmc tree building. However, one can use a Maximum Likelihood approach to obtain the likeliest topology and to use this one as a fixed.

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SUPPLEMENTARIES

Morphological characters states

In order to obtain a realistic probabilistic value for of k characters states for each simulated morphological character, we downloaded 100 random morphological characters (with more than 100 characters each) from TreeBASE database (http://treebase.org/) published between 1985 and 2013 and covering 19 taxomomic classess (Chordata, Arthropoda, Annelida, Angiosperm, Gymnosperm and Pteridophyta). We selected a total of 22563 characters ranging from 2 to 10 states. We calculated the proportion of characters with 2, 3, 4, 5, 6, 7, 8, 9 or 10 states. We then sampeled 22563 k values between 2 and 10 with the

same proportion of characters from the empirical data. We then used a simple t-test to check if our simulation was equal to the empirical data. In this study, we only simulated characters with 2 or 3 states because of the high proportion of ordered characters ecountered on characters with more than 3 states and the difficulties of simulate biologicaly sensible ordered characters.

Tree Building Software settings

Maximum Likelihood - RAxML v8.0.20 (Stamatakis 2014).—

Model:

Molecular data:

 $GTR + \Gamma_4$ (-m GTRGAMMA)

Morphological data:

 $\mathrm{Mk} + \Gamma_4 \; (\text{-K MK})$

Support:

Rapid Boostrap algorithm (LSR), 1000 replicates

Bayesian - MrBayes v3.0.2 (Ronquist et al. 2012b).—

Priors:

Molecular data:

rates distribution shape $(\alpha) = 0.5$

Transition/Transversion ratio = $2 (\beta(80,40))$

Morphological data:

rates distribution shape $(\alpha) = 0.5$

Models:

Molecular data: HKY + Γ_4

Morphological data: $Mk + \Gamma_4$

Frequencies of character states number from 100 matrices

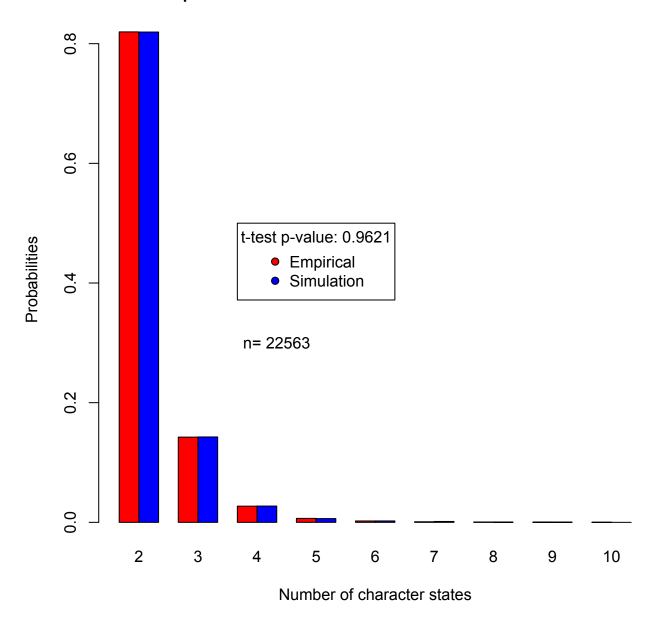


Figure 4: Character states distribution in empirical matrices. Characters states number distribution extracted from 100 random morphological matrices downloaded from RreeBase.

MCMC:

2 runs

4 chains per run

generations; 50×1^6

sample frequency = 1050×1^3

ASDS diagnosis frequency = 50×1^3

ASDS < 0.01

ESS >> 200

Burnin = 25%

Triplets metric details $(T_{x,y})$

Each triplet can be written as $I_{ijk}=(ijk)$). Where I_{ijk} is equal to 0 if the two triplets (ijk) are the same in the two trees otherwise I_{ijk} is equal to 1. For any rooted tree there are only four possible combinations per triplets: ((j,k),i); ((i,k),j); and ((i,j),k); and (i,j,k); (Johnson and Soltis 1998). One can calculate S_n , the triplet distance between two trees as:

$$S_n = \sum_{ijk} I_{ijk} \tag{5}$$

Where:

$$\sum_{ijk} = \binom{n}{4} = \frac{n!}{4!(n-4)!} \tag{6}$$

And where n is the number of taxa in both trees (modified from Critchlow et al. (1996)). When all triplets across the two trees are the same, S_n is equal to 0 and when all the triplets are different S_n is equal to $\binom{n}{4}$. Because the possible number of triplets per clade is a finite number, the probability of two random trees with the same n taxa to have the same triplet is:

$$P(I_{ijk} = 0) = \frac{1}{4} \tag{7}$$

Therefore one can calculate the probability of two random trees having the same triplets:

$$P(S_n = 0) = \sum_{ijk} P_{I_{ijk} = 0}$$
(8)

$$P(S_n = 0) = \frac{n!}{4(3!(n-3)!)}$$
(9)

And in the same way:

$$P(S_n = 1) = \frac{3n!}{4(3!(n-3)!)}$$
(10)

RF metric details

The RF distance (or path difference) between two trees reflects the distance between the distributions of the tips among clades in the two trees (Robinson and Foulds 1981) and can be expressed as following:

$$RF_{x,y} = N_x + N_y - 2C_{x,y} (11)$$

Where $C_{x,y}$ is the number of clades in common in the two trees. The minimal value of C is equal to 1 if the two trees have the same n taxa; the maximal value in C=n-2. For a fully unresolved tree (star tree) N=1 and for a fully resolved tree (binary tree) N=n-2. The minimal and maximal topological distance for n taxa is:

$$RF_{min} = 1 + 1 - 2C_{x,y} (12)$$

And:

$$RF_{max} = 2(n-2) - 2 (13)$$

One can then rescale RF.scaled by using the maximal and minimal value for any n taxa:

$$RF.scaled_{x,y} = \frac{RF_{x,y} - RF_{max}}{RF_{max}}$$
(14)

This metric is more sensitive to taxa displacement than the Triplet distance (Critchlow et al. 1996; Johnson and Soltis 1998; Wiens 2003) and therefore a low value will show a good clade conservation between two trees and a high value will show a bad recovery of common clades.

Codes

All codes are available at: https:

 $//github.com/TGuillerme/Total_Evidence_Method-Missing_data/tree/master/Functions$

Table 1: Glossary

Term	Definition
living taxa	taxa with both molecular and morphological data available
fossil taxa	taxa with only morphological data available
"complete" matrix	matrix with no missing data except for the molecular data for the fossil taxa
"missing-data" matrix	matrix with various amount of missing data
M_L	missing living taxa in the morphological part of the matrix
M_F	missing data for the fossil taxa in morphological part of the matrix
M_C	missing morphological characters for both living and fossil taxa
"true" tree	tree used to simulate the matrix
"best" tree	tree inferred from the "complete" matrix
"missing-data" tree	tree inferred the "missing-data" matrices
"horizontal" sub-matrix	sub-matrix with morphological and molecular characters for living taxa
"vertical" sub-matrix	sub-matrix with morphological data for both living and fossil taxa
"corner" sub-matrix	sub-matrix with morphological data for living taxa
"missing-data" sub-matrix	molecular part of the matrix for the fossil taxa (no data)