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An analysis of maximum parsimony algorithms to predict parasitism in Eukaryota

using a large multifurcated phylogenetic synthesis tree

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Lydia Buntrock E-Mail: info@irallia.de

Supervisor:

Prof. Dr. Bernhard Y. Renard

&

Prof. Dr. rer. nat. Emanuel Heitlinger

Abstract

Parasitism can be defined as an interaction between species in which one of the interaction partners, the parasite, lives in or on the other, the host. The parasite draws food from its host and harms it in the process. According to estimates, above 40% of all eukaryotes are parasites. Nevertheless, it is computationally difficult to obtain information whether a particular taxon is a parasite making it difficult to query large sets of taxa.

Here we test in how far it is possible to use the Open Tree of Life (OTL), a synthesis of phylogenetic trees on a backbone taxonomy (resulting in unresolved nodes), to expand available information via phylogenetic trait prediction. We use the Global Biotic Interactions (GloBI) database to categorise 25,992 and 34,879 species as parasites and free-living, respectively and predict states for over \sim 2.3 million (97.34%) leaf nodes without state information.

We estimate the accuracy of our maximum parsimony based predictions using cross-validation and simulation at 60-80% overall, while strongly varying between clades. The cross-validation results in an accuracy of 98.17% which is explained by the fact that the data are not uniformly distributed. We describe this variation across taxa as associated with available state and topology information. We compare our results with several smaller scale studies, which used manual expert curation and conclude that computationally inferred state changes largely agree in number and placement with those. In clades in which available state information is biased (mostly towards parasites, e.g. in Nematodes) phylogenetic prediction is bound to provide results contradicting conventional wisdom.

This represents, to our knowledge, the first comprehensive computational reconstruction of the emergence of parasitism in eukaryotes. We argue that such an approach is necessary to allow further incorporation of parasitism as an important trait in species interaction databases and in individual studies on eukaryotes e.g. in the microbiome.

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1 Introduction

This thesis is about the analysis of ancestral state reconstruction algorithms for non-binary trees, applied to the currently largest phylogenetic synthesis tree of Open Tree of Life (OTL) [1] to predict parasitism in Eukaryota.

For about 50 years, people have been working on ancestral state reconstruction, the inference of evolution that leads to the given data.

One of the first papers was written by Camin and Sokal, who in 1965 were working on algorithms for discrete-state data [2]. Different methods have been developed and the question is which method is the most suitable for the problem at hand: the ancestral state reconstruction for a huge non-binary tree with two discrete states.

Royer-Carenzi et al. distinguish two major classes of ancestral state reconstruction methods: The first is maximum parsimony: explain the current state with the least number of state changes between the child and its ancestor.

The other class they present describes modeling the character evolution as a stochastic process and using the likelihoods to compute the possible ancestral character states. This is generally done with a continuous time Markov model [3].

TODO: Pasqualin et al. unterscheiden noch eine weitere Methode: stochastic mapping...

One of the major disadvantages of parsimony methods is that, unlike likelihood approaches, they can not take divergence times (branch length) into account. Since the OTL does not include development times of species, this can not be considered here.

Another problem pointed out by Royer-Carenzi is that parsimony approaches are either based on predefined parameters (generalized parsimony) or on strong and often controversial assumptions, like irreversibility of transitions for dollo parsimony. This problem is unimportant to the present task because in the analysis of the entire Eukaryota tree only generalized

models are meaningful.

The comparison of methods gives us the maximum parsimony method as the simplest and fastest method that meets all our requirements.

Felsenstein [4] discusses in his book two parsimony algorithms that generalize previous methods (from Camin and Sokal [2], Farris [5] and others): Fitch parsimony [6] and Sankoff parsimony [7]. Therefore, these are the methods used in this work.

For Fitch, the algorithm has been extended from binary to non-binary trees. For the Sankoff algorithm, Louca and Doebeli have presented an implementation for non-binary trees published in an R package named *castor* [8].

For an ancestral reconstruction on the phylogeny of Eukaryota, we use Open Tree of Life (OTL) [1].

Phylogeny describes the evolution of species, while the taxonomy is a classification according to certain criteria in so-called taxa.

OTL is a comprehensive, dynamic and digitally available tree of life constructed from published phylogenetic trees along with a backbone of taxonomic data. Besides this, Hinchliff et al. also offer an OpenTreeOfLife-Taxonomy (OTT) with the help of which we identify the individual nodes.

It follows that the biggest 'phylogenetic tree' is this synthesis of phylogenetic trees filled with a taxonomic trees given by OTL.

For these large phylogenetic synthesis trees, however, ancestral state reconstruction has so far only been done for Bacteria and Archaea for binary traits by Goberna and Verdú [9]. However, this differs from eukaryotes in the sense that complex traits such as parasitism depend on more than one gene.

The present tree structure of OTL is not binary but multifurcated, meaning that each node has multiple (n > 2) children or in other words its degree, number of adjacent nodes, is greater than 3 [4].

Parsimonious in phylogeny refers to favoring the tree that needs the least evolutionary change to explain the observed data. Maximum parsimony methods have been developed for phylogenies, which are usually depicted as binary trees. Therefore, the selected parsimony methods are not directly applicable, as they were specifically applied to much smaller subtrees, where all splits are known.

We will extend and test the existing maximum parsimony algorithms of Fitch [6] and Sankoff [7] for this task and estimate their predictive power.

The Sankoff method is implemented by Louca et al. for the non-binary case and is available as an R package called *Castor* [8].

Note, the original Fitch algorithm has the sole purpose of minimizing the number of transitions and not reconstructing the ancestral nodes. Felsenstein [4] describes a simple extension for the reconstruction. In this work, the algorithm extended to reconstruction is adapted to multifurcated trees, based on the critical reevaluation of this extension by Cunningham et al. [10].

To accomplish this task, next to the phylogenetic tree information about the current species' states is needed.

Most of the largest interaction databases are offline or outdated. For example: IWDB (Interaction Web Database) [11], Webs on the Web [12], Animal Diversity Web [13] and ecoweb [14].

We use the interaction database Global Biotic Interactions (GloBI) [15] because it is including most of the known databases and is still growing actively [15].

The data in GloBI are stored as interactions e.g. species A parasitize species B. From this we conclude that species A is parasitic and species B free-living.

From this data, we could specify \sim 2.3 million leaf nodes 34,879 as free-living and 25,992 as parasitic.

There are many different ways to define parasitism. Since we use GloBI to classify species, we use their definition of parasitism. Again, in GloBi, Ontobee definitions are used [16].

The interaction *has parasite* is defined as: "An interaction relationship between two organisms living together in more or less intimate association in a relationship in which association is disadvantageous or destructive to one of the organisms."¹. This definition includes: ectoand endoparasites, parasitoids, kleptoparasites and pathogenes.

The objectives of this work are the following points: (1) Find a suitable ancestral state reconstruction method. (2) Accomplish reconstructing on the Eucaryotic synthesis tree of OTL.

¹ontobee.org/ontology/RO?iri=http://purl.obolibrary.org/obo/RO_000244; Last checked: 22.03.2018.

The goal of Point 1 is to evaluate the possible methods based on a simulation of our data situation.

The Sankoff algorithm implemented by Louca et al. is the best in our comparisons. Therefore, point 2 consists of reconstructing the ancestral states and predicting the unknown leaf states with the aid of this algorithm. And then perform an evaluation of the results.

2 Aims

The aim of this thesis is the application of maximum parsimony algorithms to non-binary trees and very large data sets to predict unknown leaf node states.

We discuss different ancestral state reconstruction methods and test the most appropriate ones on our data. Hereby, we want to find out how far it is possible to use the Open Tree of Life (OTL) to expandavailable information via phylogenetic trait prediction. As a basis for known states, we use the Global Biotic Interactions (GloBI) database, which cathegorises 34,879 and 25,992 species as free-living and parasites. This data is used to perform an ancestral state reconstruction and predict states for over \sim 2.3 million (97.34%) leaf nodes without state information.

In order to generate a realistic simulation, influencing parameters are investigated. Since the transitions are minimized in an ancestral state reconstruction, this is an important parameter to consider. On the other hand, the completeness of our input data is an influencing value. Therefore, two major types are distinguished:

- i) Biological parameters (a result of the evolutionary process):
 - State distributions and transition probabilities
- ii) Distribution of missing information:
 - Lack of information on topology (→ multifurcations)
 - Lack of information of states of some leaf nodes

3 Methods

In this thesis, a maximum parsimony algorithm is applied to the Eukaryota tree to obtain an ancestral state reconstruction of free-living versus parasite states.

This chapter is divided into the following sections: the description of the data sources (section 3.1) used and the analysis of these data as a preparation for the simulation (section 3.2). A description of the analyzed methods for the ancestral state reconstruction (section 3.3) and then an explanation of how they fit the problem at hand (section 3.4). And at the end, the real data analysis with the Sankoff method (section 3.5).

Figure 3.1 briefly outlines these relationships. A more detailed view of the workflow can be found in the appendix 7.1.

3.1 Data sources

Two types of data are needed for an ancestral state reconstruction: a tree and information about the states.

For the subsequent analyzes on the Eukaryote tree the database of Open Tree of Life (OTL) is used (downloaded on 16.02.2018) [1]. This database gives a synthesis of phylogenetic trees (currently 819 trees) and a taxonomic tree¹. OTL also includes the large phylogenetic database TreeBASE [1].

Furthermore the Open Tree Taxonomy (OTT) from OTL is used because it includes most of the known taxonomies and is synthesised by prefering taxonomies that match with available phylogenetic data. The team from OTL prefer a maximum number of species [1], this results in the synthesis between taxonomy and phylogeny.

¹https://tree.opentreeoflife.org/about/synthesis-release/v9.1; Last checked: 22.03.2018.

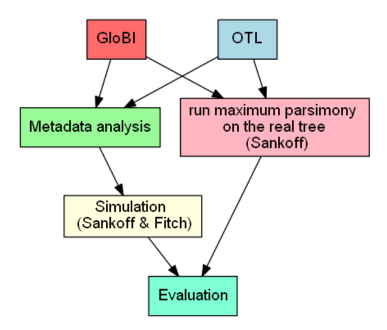


Figure 3.1: The Workflow of the resulting procedure with the following steps:

(1) Retrieve phylogenentic tree data as input for the tree (OTL) and the state information (GloBI). (2) Get metadata of these for a realisitc simulation of the maximum parsimony algorithms (Fitch & Sankoff). (3) Build and run the simulation. (4) Evaluation of parameters for the simulation and the ancestral state reconstruction of the real tree. (5) Evaluate the accuracy of developed algorithms and choose the best. (6) Run the resulting algorithm on the original data. (7) Evaluate and interprete the results.

For the state information the Global Biotic Interactions database (GloBI) is taken [15] (downloaded on 29.01.2018).

This database consists of entries of the form: species A (source) interacts with B (target). A number of interactions have been identified², including those indicating whether the species source or target has become a parasite or a free-living species from the biological perspective. They are the following:

- free-living source: preysOn, eats, flowersVisitedBy, hasPathogen, pollinatedBy, hasParasite, hostOf
- free-living target: preyedUponBy, parasiteOf, visitsFlowersOf, pathogenOf, hasHost
- parasite source: parasiteOf, pathogenOf

²https://github.com/jhpoelen/eol-globi-data/.../InteractType.java; Last checked: 22.03.2018.

• parasite target: hasParasite, hasPathogen

Of these interactions, e.g. species A parasitize species B, the state of the species is determined, here is species A parasitic and species B free-living. In the case that a parasite conquers (parasitizes) another parasite yields conflicting states for the second species. This is solved by preferring parasitic.

For each species known identifyers are stored in GloBI. This includes OTT (the taxonomy of OTL). All species that have stored an OTT indentifyer and have a matching interaction are divided into two lists: parasites and free-livings.

3.2 Metadata analysis

Based on the input data, generalized linear models are compared with poisson respectively binomial regression according to their residuals. In order to compare models of different complexity, the BIC (Bayesian Information Criterion) values are calculated in addition to the residuals.

There are two different information criteria: AIC (Akaike Information Criteria) and BIC.

The advantage of the BIC is that the penalty is dependent on the sample size and is therefore advantageous for large samples.

For all these calculations, the following R functions are used: glm(), anova() and BIC().

For each node, depth, min, max and mean height are noted. The node depth is calculated as the distance (number of edges) to the root node. The node height is calculated in three different ways: min, max and mean height refers to the (mininmal, maximal, mean) distance to a leaf node.

The influence in the modeling of these parameters is tested, additively as well as multiplicatively to the models.

3.3 Ancestral state reconstruction methods

The methods used to reconstruct the ancestral states are Fitch parsimony [6] and Sankoff parsimony [7], which are explained in the following subsections.

3.3.1 Fitch maximum parsimony

Based on the work of Cunningham et al. [10] the Fitch method [6] is implemented and extended by us for the multifurcated tree case. To understand the differences to the multifurcated case, the algorithm for the binary case is briefly explained and referred to the extension.

Input: A rooted, binary tree, with state informations in the leaf nodes. Each node is depicted as a set of states. There are only two states in this thesis, free-living (FL) and parasitic (P). Internal nodes have three sets, which are empty at the beginning, excluding the root node, it has only one. Leaf nodes have their state as a set (e.g. $\{FL\}$ or $\{P\}$, unknown leaf nodes the union of all possible states ($\{FL, P\}$).

The algorithm traverses three times through the tree and fills these sets.

In each step, two sets are considered and their intersection is formed. There are two cases:

- i) The intersection is not empty and corresponds to the new set.
- ii) The intersection is empty. \rightarrow Build the union of these sets as new set.

First traverse from the leaf nodes to the root: each internal node is formed from its child nodes, where the only information lies at the beginning.

Second traverse from the root node to the leafs: each internal node is formed from its parent node and its sibling node.

Last traversion (direction does not matter): build the final state for every node. It is formed from the sets of previous traversals.

(The original Fitch algorithm is designed to minimize transitions without predicting actual states of internal nodes, so it is just the first traversal.)

The extension to the non-binary case is quite obvious, but holds some opportunities. In this case, more than two children may be present for the first traversal, but the incision or union may also be formed over more than two sets. Also in the second traversing, there may be several sibling nodes. However, there are several possibilities here that are all tested and compared in the simulation. Some of these options are already available in the binary case:

- The parent node has (except for the root node) two state sets, because he came through the up-traversing previously. Are both sets used or only the first traversing?
- If there are several siblings, is the cut or union made first of these, or is everything made directly including the parent node.

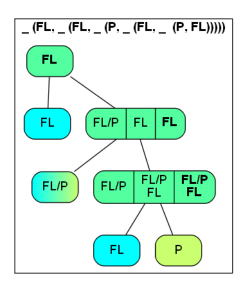


Figure 3.2: Fitch algorithm for binary trees.

The unknown leaf node is discribed with both states. Computed internal nodes (exclusive the root node) consists of three sets, where the last set is the final one (bold).

From the second internal node (seen from the root node) there are several possibilities to create the second and third set.

The first point already has an effect on the binary case. Figure 3.2 shows both possibilities of the three sets.

Cunningham uses only the first state set of the parent node [10].

From these two points four different versions of Fitch are formed:

- i) Fitch 1: First state set of parent node; intersection/union of siblings first.
- ii) Fitch 2: First state set of parent node; intersection/union of siblings together with parent node.
- iii) Fitch 3: Both state sets of parent node; intersection/union of siblings first.

iv) Fitch 4: Both state sets of parent node; intersection/union of siblings together with parent node sets.

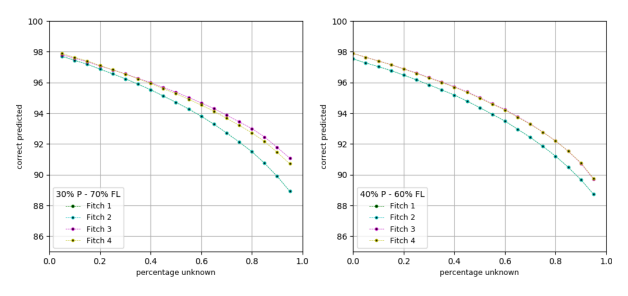


Figure 3.3: Test of Fitch Versions. TODO: beschriften!!

These four versions are tested in the simulation with 100 trees and 10000 leaf nodes and distributions of 70% FL to 30% P and 60% FL to 40% P. Figure 3.3 shows this over all unknown node percentage.

At 95% unknown nodes and 95% of multifurcation of the internal nodes, version 1 is 88.37%, version 2 is 88.37%, version 3 is 88.4%, and version 4 is 88.39% correct. Therefore, only version 3 is used for all further simulations.

Sankoff

Maximum parsimony algorithm from Sankoff implemented in the R package castor [8]. From this the function $hsp_max_parsimony()$ is used with default settings including $transition_costs = "all_equal".$

3.4 Simulation

The simulation compares these different ancestral state reconstruction algorithms with each other.

First different implementations of the Fitch maximum parsimony are compared and then the best of them is compared with the implementation of the Sankoff algorithm of the *Castor* package [8].

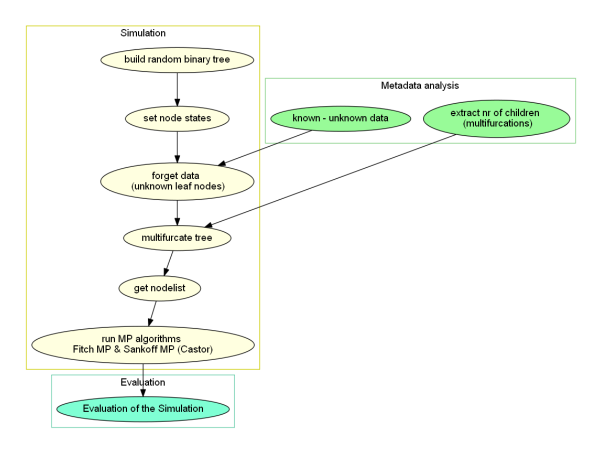


Figure 3.4: A simulation is performed to compare different maximum parsimony algorithms. The course of the simulation with influence of the metadata analysis from the real data can be seen:

- (1) A phylogenetic tree is created randomly. (2) Simulate node states for all nodes. (3) 'Forget' internal states and some leaf node states. (4) 'Lose' phylogeny information.
- (5) Make a nodelist for the algorithm. (6) Run algorithms. (7) Evaluate results. Points 3 and 4 are influenced by metadata of the real-data analysis.

The course of a simulation is shown in Figure 3.4. The individual steps are explained in the following.

A tree is needed to perform a simulation of ancestral state reconstruction. It has to be decided whether to take the real tree or simulate a tree. In this simulation, trees are created randomly, as one can replicate a complete binary phylogentic tree. Thus, there is also the possibility to simulate the multifurcation.

To get a random binary tree, the Phylo package from biopython is used [17]. They offer a randomized() function which returns a BaseTree³.

The next step is to simulate states for all nodes.

Here the influence of the biological parameters as transition probabilities and distributions of states is included.

Since there is no statement about general transition probabilities, these are all set the same:

$$\mathcal{P}(FL \to P) = \mathcal{P}(P \to FL).$$

For the distributions of states different ratios of parasites (P) to free-livings (FL) are simulated with the help of beta distributions and a given threshold:

- 50% P to 50% FL.
- 40% P to 60% FL.
- 30% P to 70% FL and
- 20% P to 80% FL.

Procedure: The root node is defined as ancestor of all subsequent species and in this case, determined to be free-living. Therefore, the beta distribution for free-living is used at the beginning. Now traverse from the root to the leaf nodes, always pulling out of the current

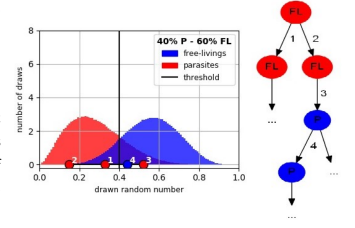


Figure 3.5: Set node states: Distribution of states (left); traversion through the tree (right).

Start with a free-living root node (FL: red)

- (1) + (2) Draw random numbers for its children from the free-living distribution (red), the numbers are under the threshold \rightarrow the nodes are again free-living; go on with the children.
- (3) The number drawn is above the threshold. \rightarrow The node state changes to parasitic (P: blue).
- (4) Now draw random numbers from the parasite distribution (red) until one number lies unter the threshold. Then change back.

³https://github.com/biopython/biopython/blob/master/Bio/Phylo/BaseTree.py

distribution until the randomly drawn number is above the threshold and the new node changes state. Figure 3.5 shows a part of this simulating states and the associated distributions.

After traversing through the tree, each state is saved in a nodelist associated with the node indentifyer which is the OTT from OTL.

Here begins the simulation of the lack of information, such as the lack of information on topology (multifurcation) and of states of some leaf nodes.

In the real tree, there is usually only information about species living today \rightarrow leaf nodes. And beyond only a small percentage of these. All information about the states of the internal node and one leaf node is removed and stored in another column to the node.

Last step for the preparation is the multifurcation of the tree. As previously explained, some divisions in the tree are not known, so the real tree is not binary. This multifurcation is simulated by an equally distributed percentage of forgotten internal nodes.

Different percentages of removed information are simulated.

The last step is the evaluation of the results. This is done with a simple difference calculation of the node states.

In the nodelist, the originally simulated states and the newly calculated states are stored for each node (FL=0, P=1). The sum of the differences of the node states gives the distance of the prediction to the original tree.

3.5 Real data analysis

For the evaluation of real data results, a leave-100-out cross-validation is performed. For this purpose 100 randomly distributed 100 states are left out.

Analogous to the missing leaf node states, generalized linear models with binomial regression are compared according to their residuals, based on the "true" / "false" prediction.

Again, in order to compare models of different complexity, the BIC (Bayesian Information

Criterion) values are calculated in addition to the residuals.

For all these calculations, the following R functions are used: glm(), anova() and BIC().

3.6 Implementation

The complete code is located on GitHub: github.com/Irallia/IZW-HU-Parasites. Most of the code is written in Python, the analyzes and the use of the Castor package in R. Some shell scripts are used to execute whole workflows.

4 Results

This work deals with the ancestral state reconstruction of the entire Eukaryota relatives tree.

For this reconstruction, we first analyzed our data. This is the tree of OTL [1] and the data for the leaf node states (free-living or parasite states) from GloBI [15] (section 4.1), where we have determined that the entire Eukaryota tree is 89.45% multifurcated (missing internal nodes) and that we do not have any state information for 97.34% nodes.

In addition, we tested different models for these two paramters and found out that multifurcation can best be modeled with the taxa 'order' multiplied by the mean height of the knots. Also, the missing states could be modeled with the taxa 'order' and in this case multiplied by the depth of the nodes.

Next we compared different possible methods. We decided to take a closer look at maximum parsimony algorithms because they are fast and best suited to the problem at hand. We tested these on different simulations of the data and compared their predictive power (section 4.2). We compared them with the result that the Sankoff algorithm performs significantly better than Fitch. The multifurcation rate has little impact on Sankoff, but Fitch's prediction significantly breaks down at more than 60% missing internal nodes. However, the amount of missing node state information has a linear impact on both, as does the ratio of free-living to parasitic (the more balanced, the worse is the prediction).

As a result, we used the Sankoff algorithm [7] implemented by Louca et al. [8], which performed best, for the actual reconstruction on the real data.

TODO: ... biological view We evaluated these by performing a leave-100-out cross-validation and predicted approximately 98.17% of the omitted states correct (subsection 4.3.2).

4.1 Availability of internal nodes and state information

As previously presented, we have two types of missing information: unknown states of leaf nodes and multifurcation.

A tree is multifurcated if there are nodes that have more than two children. A binary tree with n leaf nodes has n-1 internal nodes. The present Eukaryota tree of OTL has 2,293,463 leaf nodes and only 241,974 internal nodes, that is:

$$100 - \frac{100}{(2293463 - 1) \times 241974} \approx 89.45\%$$

missing internal nodes. This means that there is a lack of information about the underlying phylogeny. Instead of a binary tree this tree is highly multifurcated.

Subtree of	Unknown States	Multifurcation
Eukaryota	97.34%	89.45%
Metazoa	96.44%	87.93%
Fungi	98.87%	96.97%
Chloroplastida	99.14%	89.46%
Apicomplexa	86.26%	87.16%
Nematoda	89.01%	88.59%
Chordata	88.59%	66.49%
Platyhelminthes	68.73%	80.34%
Insecta	97.11%	90.78%

Table 4.1: Examination of subtrees regarding missing information.

The percentage values show the proportion of missing information of: unknown states (missing state information of leaf nodes) and multifurcation (missing internal nodes). The subtrees are from different taxa: domain (Eukaryota), kingdom (Metazoa, Fungi, Chloroplastida), phylum (Apicomplexa, Nematoda, Chordata, Platyhelminthes) and class (Insecta).

The two by far smallest values are highlighted in green.

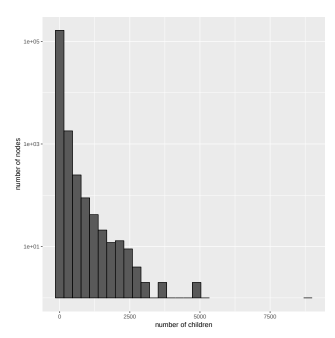
For the missing state information the present Eukaryota tree with 2,293,463 leaf nodes,

34,869 free-livings and 25,962 parasites are found. This gives

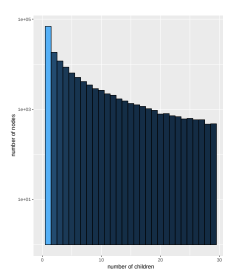
$$100 - \frac{100}{2293463 \times (34860 + 25962)} \approx 97.34\%$$

unknown states of leaf nodes.

We calculated these percentages of missing nodes and also missing state information for some subtrees and plotted them in table 4.1.



(a) Histogram with automatic binwidth.



(b) Histogram with binwidth = 1. Higher multifurcation than 30 has been cut off.

light blue: binary; dark blue: multifurcation

Figure 4.1: Histograms showing the multifurcation of the internal nodes of the synthesis tree. For each node, the number of children (degree -1) is collected. A node is multifurcated if it has more than two children, so we deducted two from each number of children. The two histograms show: number of children -2 on the x-axis with log scale and the number of nodes with this amount on the y-axis.

For a first overview, we collected for each node its number of children (degree -1), and plotted this in two histograms, see figure 4.1.

The multifurcation affects only the internal nodes. We collected the number of children -2 of these nodes (because a node with two children is binary). That means it describes the

number of nodes which we have lost from the real (binary) phylogenetic tree. It can be recognized that we are very far from a binary tree.

4.1.1 Poisson regression of the multifurcation

For a regression analysis of the multifurcation first set up a simple generalized linear model and then output the intercept of it:

Since the multifurcation can also be 0, the intercept describes the expected mean value and thus, if it is greater than 0, it says that we have a multifurcation: $2.62 > 0 \Rightarrow$ there is a multifurcation.

We next compared the intercept of the different kingdoms and found that

```
1.22 (Fungi intercept) > -0.11 (Metazoa intercept) > -0.09 (Chloroplastida intercept)
```

that means the strength of the multifurcation is the same/has the same size sorting TODO: kann das sein?

We investigated several models that could describe multifurcation:

In doing so, we allowed the different influence of the taxa and the heights and depths of a node to be included. From this we got 9 times 4 models of different complexity levels (see first row of table 4.2).

We first compared the models based on their residuals. These give us the error of the model. If the value is small, our data will be well modeled. These values are meaningful only for the same complexity, therefore we then calculated the BIC (Bayesian information criterion) values of the models and used them to evaluate the models.

The deviance table with the residuals can be found in the appendix (table 7.3). Table 4.2. lists the BICs.

Model / Taxa	Kingdom	Phylum	Class	Order
multifurc \sim taxa	8273333	7937828	7842157	7644249
multifurc \sim taxa	8257680	7922207	7826490	7574154
multifurc \sim taxa $+$ depth	8273318	7934322	7839364	7539999
multifurc \sim taxa $+$ max.height	7993515	7749121	7661817	7416211
multifurc \sim taxa $+$ min.height	8251211	7875521	7778327	7516883
multifurc \sim taxa $+$ mean.height	7825417	7644249	7572474	7340741
multifurc \sim taxa * depth	8235932	7836755	7757688	7383808
multifurc \sim taxa * max.height	7963438	7693555	7614820	7335338
multifurc \sim taxa * min.height	8214030	7808940	7690618	7336627
multifurc \sim taxa * mean.height	7768360	7536296	7484953	7206369

Table 4.2: BIC (Bayesian information criterion) values of the multifurcation models. These models are created with the R function glm() and compared with the BIC() function. This results in the listed BIC values.

Within the every complexity class it can be seen that the mean height gives the best additional factor.

Despite higher complexity, the BIC values are getting smaller from model to model, meaning that the finest model available here is also the best one of these. Lower taxa than orders (e.g. family) are computationally too expensive to calculate.

The model $multifurc \sim order * mean.height turns out to be the best of our models, whereby it is possible that e.g. <math>multifurc \sim family * mean.height$ is better.

4.1.2 Binomial regression of the unknown state information

Next to the problem of the multifurcation of the tree is the less of interaction data we have for the species. For the ancestral state reconstruction, we need information about the states (free-living or parasite) in the leaf nodes.

The eukaryotic synthesis tree has 293,463 leaf nodes. The GloBI database has 5,346,414 interactions (at 29.01.2018). Out of this data we got 51,337 distinct free-living species and 47,332 distinct parasite species for the whole tree of life. From the Eukaryota we could

determine 25,992 and 34,879 species as parasites and free-living. With 2,293,463 leaf nodes we still have about 97.34% unknown leaf nodes. In the discussion, section 5.1, we will talk about ways to extend this data.

We also compared different models in terms of their BICs (Table: 4.3). The Residuals are not very meaningful here, since all models have different complexities. For the sake of completeness, the associated deviance tables are located with the residuals in the appendix 7.4.

Model / Taxa	Kingdom	Phylum	Class	Order
multifurc \sim taxa	545799	500004	485121	484681
multifurc \sim taxa $+$ depth	544862	493808	481869	478851
multifurc \sim taxa * depth	544179	489845	481494	478188

Table 4.3: BIC (Bayesian information criterion) values unknown state information models. These models are created with the R function glm() and compared with the BIC() function. This results in the listed BIC values.

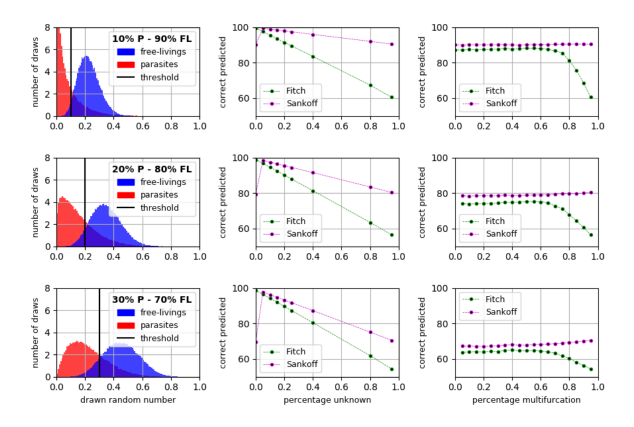
It also follows from this table that the most complex model is the best. In general, the BIC values are smaller than those of the multifurcation models. The modeling here is thus better. Again, the calculation of finer models (e.g. order or family) was too expensive.

These missing data modeling results can be used to better simulate the data.

4.2 Results of simulation / Influence of different parameters

As presented, we compare two methods in our simulation to their prediction accuracy: Fitch and Sankoff.

Figure 4.2 shows the results of examining various parameters.



First, we tested different transition probability distributions including threshold (first column). It can be observed that the more balanced the percentages of free-livings and parasites, the worse the prediction of the algorithms (second and third column).

It can be seen that both algorithms are always over 50% prediction and therefore better than guessing. Moreover, they are usually close to each other, with Sankoff always makes better predictions except for equally distributed states as Fitch.

On the other hand, we examined the influence of missing internal nodes (ridge of multifurcation) and missing leaf node information (unknown leaf nodes).

Both factors have a relatively linear influence on the Sankoff method. Fitch, on the other hand, breaks significantly in his prediction from about 70% unknown leaf nodes or mutifurcation.

Since we have a lot of missing data in most subtrees (both internal nodes and state informations), the Sankoff gives a better prediction and is thus used for real data analy-

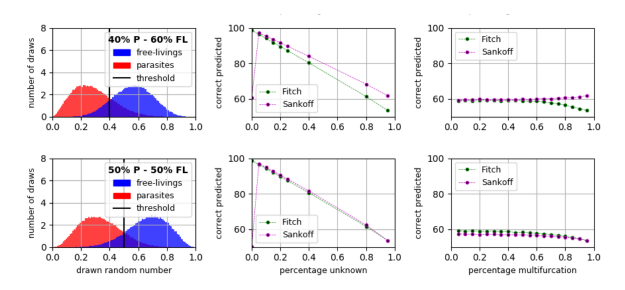


Figure 4.2: Influence of unknown data to prediction.

The first column describes the distributions of free-livings and parasites with a given threshold for the respective simulations to the right.

The middle column investigates the influence of the unknown states, the right the influence of the strength of the multifurcation.

The y-axes indicate the percentage of correctly predicted states (including known states). On the x-axis the percentage of forgotten states or missing internal nodes.

Each point corresponds to the average of one hundred simulations, each with 10,000 leaf nodes.

For the middle column we set the strength of the multifurcations to 0.95% similar to the real data and in the right column the amount of the unknowns to 0.95% also similar to the real data.

4.3 Results of the real data analysis created with Sankoff

This section is about evaluating the prediction of real data using the Sankoff method. It is divided into two subsections. 4.3.1: The analysis of some subtrees using biological background knowledge. 4.3.2: Presentation of the results of the leave-100-out cross-validation.

4.3.1 Biological view

To analyze the results, we have selected some phyla (subtrees) to evaluate our results selectively from the biological point of view: Chordata, Nematoda, Platyhelminthes and Apicomplexa.

In Table 4.4 we compare the given states with the predicted ones.

Several factors such as the distribution of existing input data via parasitics and free-livings, faulty input data from GloBI and reinforcement of errors by multifurcation play an important role, which crystallize out in these examples.

An important factor here is that the credibility of the results. The accuracy of the input data stands and falls with the presence and the correctness of the data of GloBI. Errors of incorrect input data can be amplified by incorrect prediction of unknown species and can be reversed in order to improve the data situation of GloBI.

Since we look at such large trees we can not expect to know all the parasites, so we look at individual positives. This is positive in the sense that the majority have the opposite state.

In contrast to the other phyla examined, the phylogeny in the Chordata is more pronounced (less multifurcation) (see Table 4.1). This results in less variance of errors. What is reflected in the results from the table 4.4. There are 18 parasites as input data and only 8 more are predicted. The Chordata mostly consist of free-living species, so this seems believable.

		original states			final states		
Phylum	# nodes	FL	Р	0 (FL)	0.3	0.5	1 (P)
Chordata	91785	10451	18	91759	0	0	26
		99.83%	0.49%	99.97%			0.03%
Nematoda	30127	21	3289	1604	142	1196	27185
		0.63%	99.37%	5.32%	0.47%	3.97%	90.23%
Platyhelminthes	22683	7	7086	175			22508
		0.1%	99.9%	0.77%	0	0	99.23%
Apicomplexa	1863	1	255	1	0	0	1862
		0.39%	99.61%	0.05%			99.95%

Table 4.4: Some selected phyla (subtrees), only leaf nodes, are examined for the evolution of the amount and percentage of given data to predicted data.

We started with 99.83% species and predict 99.97% species as free-living (including already known nodes).

We mapped the few parasitic species in a rough/simple taxonomy (see Figure 4.3): Known parasitic birds belong to the order Sauria. Here we know from Rothschild here there are breeding parasites, like the cuckoo and clepto-parasites as the skuas [18]. We got 6 input parasites from GloBI and there are no predictions: A woodpecker - *Sphyrapicus varius* and a duck - *Aix sponsa*, a cow bird - *Molothrus ate* known as broodparasite and some others. An example of the amplification of mistakes here are the carp. There is a paper from which GloBI concludes: Grass carp (*Ctenopharyngodon idella*) has Pathogen common carp (*Cyprinus carpio*)¹. Since there is hardly any information about free-living, it follows that all siblings are also predicted to be parasitic.

The Apicomplexa are a parasitic phylum. We found only one input organism: Stemonitis fusca as a free-living species. In GloBI it is listed as being parasitized by Nectria candicans and Nectriopsis sporangiicola². The algorithm has not predicted new free-livings.

Most species of Platyhelminthes (flatworms) are parasites, although there are also free-living, predatory feeding species. These are summarized in the Turbellaria, while the parasites are

¹https://www.globalbioticinteractions.org/?interactionType=hasParasite&targetTaxon=Cyprinus%20carpio

 $^{^2} https://www.globalbiotic interactions.org/? interaction Type = parasite Of \& target Taxon = Stemonitis \% 20 fuscal functions and the property of the prop$

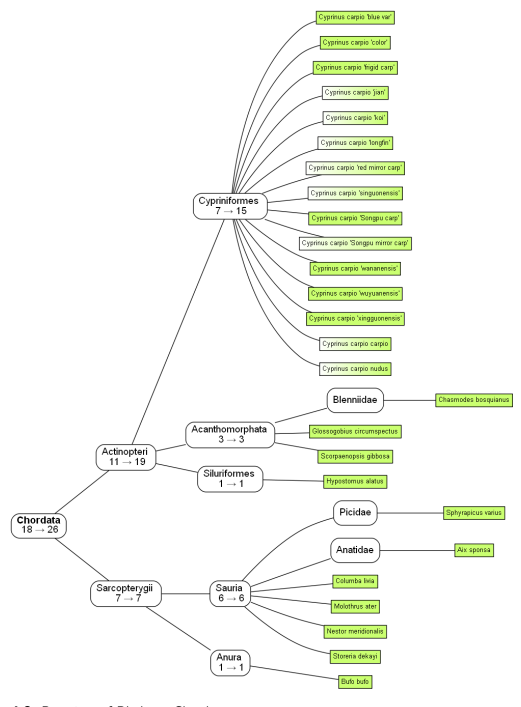


Figure 4.3: Parasites of Phylum: Chordata.

All parasite data of the chordata are mapped into a rough taxonomy (phylum, class, order, family) in order to understand their distribution and affiliation.

The internal nodes are the wanted taxa from OTL (with the addition of # input parasites to $\to \#$ predicted parasites).

The leaf nodes are the input parasites (green) and the predicted parasites (white \rightarrow green).

divided into three other classes [19]. This also corresponds to our observations. There is one class (Rhabditophora) that contains all but one single exception of free-living species of this phylum, which includes the Turbellaria.

It should be noted, however, that this classification is outdated, as it has been proven that the Turbellaria are not monophyletic. But we will not go into that here.

For the Platyhelminthes we have more state information for the leaf nodes compared to the other considered subtrees, see Table 4.1. We start with 0.1% free-livings and predicted 0.77% as free-living species.

With the Nematoda it looks more complicated. Large parts of Nematoda are free-living, but we found only 5.32% of them. Blaxter et al. estimates the order of 25,000 parasites in the Nematoda [20] and speaks of at least seven independently arosed parasitism [21]. In a recent article Blaxter identifies 18 origins [20] in Nematoda.

The problem at this point, however, is: Hallan speaks of the fact that only 23,000 species are described by the Nematoda but is it assumed that one million or more species are in this phylum.³. TODO: Link ist nicht erreichbar! The parasites have been much more studied and thus we start with only 0.63% (table 4.4) free-living species. Against such a shifted data situation, the algorithm is almost powerless to make correct predictions. And yet the percentage has increased to 5.34%.

TODO: Im Folgenden folgen 3 weitere ähnliche tabellen. Einmal eine ähnliche Tabelle inklusive interner Knoten 4.5, zweimal die Übersicht über die Kingdoms Blattknoten 4.6 bzw interne Knoten 4.7. Welche nehmen wir? Rest Appendix oder ganz raus?

Origins and Losses

Weinstein and Kuris have been searching for origins of parasitism in Animalia [22]. They identified 223 parasitic origins: 223 in Metazoa \supset 143 in Arthropoda \supset 87 in Insecta. This has led us to count the origins and losses of parasitism in our investigation as well. We count only one origin / loss in a parent node with different children's nodes.

³J. Hallan, unpublished; https://insects.tamu.edu/research/collection/hallan/

		origina	l states	final states						
Phylum	# nodes	FL	P	0 (FL)	0.4	0.5	0.67	0.75	1 (P)	
Chordata	122546	10451	18	122473	0	0	0	0	73	
Nematoda	33564	21	3289	846	0	1133	0	0	31585	
Platyhelminthes	27142	7	7086	1010	0	175	0	0	25957	
Apicomplexa	2102	1	255	1	0	0	0	0	2101	

Table 4.5: Some selected phyla (subtrees), including internal nodes, are examined for the evolution of the amount and percentage of given data to predicted data.

		origina	l states		final states						
Kingdom	# nodes	FL	Р	0 (FL)	0.25	0.33	0.4	0.5	0.67	0.75	1 (P)
none	75446	45	529	13426	220	24082	0	7792	5302	0	24493
Fungi	31457	577	2983	38520	0	0	0	5723	0	0	266463
Chloroplastida	416478	3519	77	410795	0	0	0	4182	0	0	1501
Metazoa	1491012	30758	22373	1328135	0	0	930	25535	4423	1665	130324

Table 4.6: All kingdoms (subtrees), only leaf nodes, are examined for the evolution of the amount and percentage of given data to predicted data.

		origina	l states		final states						
Kingdom	# nodes	FL	Р	0 (FL)	0.25	0.33	0.4	0.5	0.67	0.75	1 (P)
none	84456	45	529	15035	243	25910	0	8764	6183	0	28140
Fungi	324105	577	2983	39088	0	0	0	5858	0	0	274803
Chloroplastida	460457	3519	77	454211	0	0	0	4688	0	0	1558
Metazoa	1670956	30758	22373	1485749	0	0	1313	29002	5102	1957	147833

Table 4.7: All kingdoms (subtrees), inkluding internal nodes, are examined for the evolution of the amount and percentage of given data to predicted data.

Here we have encountered a problem: The Castor algorithm gives us probabilities for states. That means there are also nodes with state like 0.3 or 0.5. The question is, how to count these state changes. We decided to round the values.

In Table 4.8 we can see, that we found some more origins than Weinstein and on top of that some losses.

Domain	# internal	# leaf	Rootnode	without and	d with rounding
/ Kingdom	node	es	state	# origins	# losses
/ Phylum / Class				(FL -> P)	(P -> FL)
Eukaryota	241974	2293463	1.0	415	363
			Р	462	369
Metazoa	179944	1491012	0.5	294	123
				321	129
Fungi	9534	314571	0.5	80	222
				97	222
Chloroplastida	43486	412434	0.0	40	2
			FL	42	2
Arthropoda	120479	1198981	0.0	260	102
			FL	281	108
Apicomplexa	239	1863	1.0	0	1
			Р	0	1
Nematoda	3437	30127	1.0	0	11
			P	2	11
Chordata	30761	91785	0.0	12	1
			FL	12	1
Platyhelminthes	4459	22683	1.0	0	5
			Р	0	5
Insecta	91256	989572	0.0	234	77
			FL	245	77

Table 4.8: Origins and losses

When more closely regarding the same phyla as in the section before: Chordata, Nematoda, Platyhelminthes and Apicomplexa.

Chordata are full of free-living species and so we see only a few origins of parasitism. The root and mostly all species are predicted as free-living.

In Apicomplexa and the Platyhelminthes also look credible. Our algorithm gives us only one loss of parasitism in Apicomplexa and five in the Platyhelminthes. They are both from the root over mostly all species predicted as parasites.

Nematoda is again full of problems. The root node is predicted as a parasite and so we have more losses of parasitism for the less information of free-living species in this phylum. The rest is parasitic

As we have already mentioned Blaxter et al. found at least seven origins of parasitism [21].

If we assume that the root node of Nematoda is free-living, then some losses would have to turn around and become Origins. So it could be that we end up in a similar size as Blaxter.

4.3.2 Leave-100-out cross-validation

For a further validation of our results, we carried out a leave-100-out cross-validation. In order to achieve about 15% of the 60,871 input node states with a validation, we randomly left out 100 states 100 times. To make smaller amounts up to leave-one-out had too much computational effort.

TODO: warum 15%?

Of these 10,000 nodes, 9,238 are unique. From the unique nodes, we predicted approximately 98.17% correct and thus 1.82% wrong, with duplicate draws always having the same prediction (data not shown).

We have again dealt with the question of how this data is best modeled. The influence of the taxa (kingdom, phylum, class) and the depth of the leaf nodes is re-modeled and the BICs compared (table 4.9). Lower taxa than classes (e.g. order) are computationally too expensive to calculate.

The Residuals are included in the appendix (table 7.5) for the sake of completeness; here too, due to the different complexity, they do not have much meaning for the comparison of the models.

TODO: table for these numbers?

The BIC values this time did not prove that the finest model is the best. Of our calculated models, correctly predicted \sim phylum * depth has the smallest value.

We examined the influence of the omitted data on the prediction. On average, about twice as many leaf nodes are predicted differently. The variance is very high. This means that we have a high degree of dispersion and thus a stochastic situation exists. It also describes the

Model / Taxa	Kingdom	Phylum	Class
correctly predicted \sim taxa	117936	112242	111733
correctly predicted \sim taxa $+$ depth	117776	111304	111273
correctly predicted \sim taxa * depth	117709	111262	113135

Table 4.9: BIC (Bayesian information criterion) values of cross-validation prediction models. These models are created with the R function glm() and compared with the BIC() function. This results in the listed BIC values.

width of the present probability function. The standard deviation is about five times as high as the number of omitted nodes, so the variability is quintupled. Table 4.10 shows these results.

		min	max	mean	variance (σ^2)	σ
	all	0	3587.70	224.96	313650.61	560.05
distance	leaf nodes	0	3021.12	208.69	248103.38	498.10
	internal nodes	0	566.58	16.28	4927.95	70.20
	all states	100	100	100	0	0
lost	FL states	44	66	57.25	19.50	4.42
	P states	34	56	42.75	19.50	4.42

Table 4.10: Statistics about the leave-100-out cross-validation

The distance between original and new states is calculated using the Euclidean metric. This is summed over all states / all leaf node states and all internal node states. The lower half of the table describes the distribution of the 'lost' states between parasites (P) and free-livings (FL).

5 Discussion

TODO: Aus den models kann man auch schließen: highly variable throughout our tree

TODO: noch irgendwo einfügen:

The number of nodes in the tree corresponding to a taxonomic rank is tabulated. For the classical taxonomic ranks phylum, class, order, family, genus xyz is found (appendix 5.2).

This chapter deals with the question of how trustworthy our result of the ancestral state reconstruction of the Eukaryota tree is. And further on how well our simulation can simulate the real problem and thus make statements about the predictive power of the applied Sankoff algorithm.

We have pursued this question in various investigations and yet, of course, further possibilities for improvement remain. Despite these possibilities for improvement, our reconstruction gives a first good assessment of the whole tree.

TODO: Why is the study important?

TODO: How does this study relate to previous studies?

We evaluated our ancestral state reconstruction and the relational state prediction using three methods: a simulation, a selective analysis from a biological perspective and a leave-100-out cross-validation.

Each theme has its own limitations.

There are various expansion options and points of criticism for the simulation, which in particular revolve around the question of how well we can simulate our real data. We dedicate this question to the subsection 5.2.

Above all, the 'biological view' gives a first impression of the credibility of the results. But he can not make any statistical statements. Here the interested researcher may have to analyze the subtrees of relevant to him for himself.

Of course, cross-validation can be expanded. With more computing power, a leave-1-out cross validation could also be performed. However, our result on this point yields a strong result with 98.17% correct prediction that this does not seem necessary.

One last limitation also exists in the data situation itself. We have pointed out in various sections that we have a strong multifurcation and only a few data about leaf node states. In figure 4.2 you can see how the predictive power can improve if we have more data available. We discuss this point in section 5.1.

5.1 Data situation

The used eukaryotic synthesis tree from OTL [1] has 241,974 internal nodes and 293,463 leaf nodes. In addition, we could specify 25,992 parasitic and 34,879 free-living species from GloBI [15].

This gives us a high number of missing internal nodes (high multifurcation) and a low number of node states of the leaf nodes. In table 4.1 we have written down this percentage including some subtrees.

The investigation of the taxonomy revealed that the OTL tree has three kingdoms: Chloroplastida, Metazoa, Fungi, 53 phyla, 195 classes and 924 orders.

Since the analysis of the tree is not part of this work, it should be mentioned here that, according to recent findings, this is not complete and we lack some taxa in every rank. For example, Cavalier-Smith says that one distinguishes between seven and nine kingdoms [23]. In section/page 40 of the appendix is a list of all phyla.

Also, the global database is not infallible. We found out (section \square) that there is some misinformation.

We are able to report some of these directly to Globi.

We found also 57,352 (not distinct!) source species and 809,993 (not distinct!) target species without OTT indentifyers. Since we currently use only OTT indentifyers, we could not use this information.

At this point there is thus the possibility to use more of the existing data, if one performs a matching with the other indentifyers.

5.2 Simulation

TODO: einbauen.... As mentioned above, in an ancestral state reconstruction, transitions are minimized, with transition probabilities playing a role.

This subsection deals with these transition probabilities from free-living (hereinafter / as a formula FL) to parasitic (hereinafter P) and vice versa: $\mathcal{P}(FL \to P)$, $\mathcal{P}(P \to FL)$.

Different parasite types have different transition probabilities. It is very difficult to make a statement about these probabilities. It is generally assumed in this work that there are 40% parasites and 60% free-livings which is based on the estimates by Windsor [24] and $\mathcal{P}(FL \to P) = \mathcal{P}(P \to FL)$, as a result of the diversity of parasites and the lack of general determinations for this. These parameters are debated in subsection 5.2.1 of the discussion.

However, the used castor package [8] offers the possibility to enter different transition probabilities.

Wieviele Übergänge gab es dabei? Hast du die gleich (ähnlich groß) gehalten? Der Anteil von Parasiten und Freilebenden ist ja nicht gleich der Transition probability. Das sind zwei verschiedene Dinge...auch wenn die Transition probability den Anteil ergibt. neben dem Anteil der P/FL ist auch die Anzalhl der Übergänge zu berücksichtigen! 50/50 kann man in einem Baum mit 1000 nodes mit einem einzigen Übergan erreichen, der die eine Häfte des Baumes ändert. 50/50 kann man aber genauso gut mit einem Übergan auf JEDER edge erreichen. Dann würde sich immer ein P mit einem FL knoten abwechselnn....

... für die Diskussion!

TODO:

The aim of the simulation is to test the influence of various unknown or unsafe parameters in order to test the credibility of the prediction.

Some of these influences could not be tested: Firstly, the distribution of parasites and free-livings in the tree produces various parameters that we could not test all (see subsection 5.2.1) and, secondly, we discuss the simulation of a phylogenetic tree on which the evaluation based on certain parts (see subsection 5.2.2).

5.2.1 Transition probabilities

We assume that different parasite types have different transition probabilities. Establishing a common distribution across the eukaryotic tree is therefore difficult.

Based on the estimates of Windsor [24], we have assumed a distribution of 40% parasites to 60% free livings in this work. As a result of the diversity of parasites and the lack of generalizations, we have generally stated that $\mathcal{P}(FL \to P) = \mathcal{P}(P \to FL)$. But it is also reasonable to assume that in general $\mathcal{P}(FL \to P) > \mathcal{P}(P \to FL)$, because a reverse mutation is usually less likely. However, one would have to determine how much this difference is and thus discuss another parameter.

In the simulation, we tested different beta distributions with different thresholds and found that this combination has a considerable influence on the predictive power of the algorithms. At this point it would be possible to test other distributions with equal threshold values or different threshold values with equal distributions.

However, if you choose a strong free-living distribution and balance it with the threshold to achieve a 40:60 parasite free-living distribution, for example, this could be very computationally expensive.

With these further simulations you could find out if the issue of distributions plays a big role. Conversely, one could estimate possible distributions based on the data location in the tree. However, this is likely to be very difficult given the poor data. For this reason, we have decided in this work to accept very general values and not to speculate much.

5.2.2 Simulation of a phylogeny

In our simulation, we start with the simulation of a phylogeny and then depict our data situation.

For this first step we use the *randomized()* function of the phylo package from biopython [17]. The question is how close is such a randomized tree to a phylogeny.

The problems that can arise are that different species develop at different rates. This means that there could be sections in the tree with many branches (for example, subtrees of unicellular species), ie with high depths and opposite sections (of very complex species, for example). It can thus be assumed that the tree is not balanced.

For more precise statements, one would have to take deeper undercurrents, which was not possible in the context of this work.

5.3 Conclusion

From the simulation, we can conclude that we predict correctly about 60% of the nodes in the present data situation (see section 4.2). The leave-100-out cross-validation even showed that we predict the omitted nodes to be 98.17% percent correct (see section 4.3.2).

This allows for the assumption that the data is grouped and not uniformly distributed and thus smaller subtrees are present in which data are to be found in the simulation with smaller multifurcation and smaller value for unknown nodes. This is also confirmed by our biological analysis of the Chordata subtree (see subsection 4.3.1).

However, this means that the ancestral states' data in the direction of root node are probably particularly unbelievable. Which makes the localization of Origins direction root node difficult. The question remains, how much this affects the estimation of the number of origins. Our comparison with the paper from Weinstein and Kuris [22] (see subsection 4.3.1), however, leaves us with optimism.

6 Bibliography

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7 Appendix

7.1 Methods overview

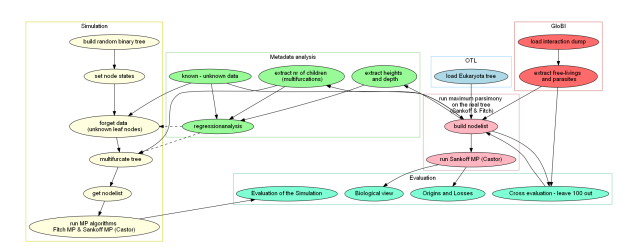


Figure 7.1: Big overview of the whole Workflow

7.2 OTL analysis

7.2.1 List of all phyla

Phyla (53):

Acanthocephala, Amoebozoa, Apicomplexa, Arthropoda, Ascomycota, Bacillariophyta, Basidiomycota, Brachiopoda, Bryozoa, Chaetognatha, Chlorophyta, Chordata, Chromerida,

Chytridiomycota, Ciliophora, Cnidaria, Colponemidia, Ctenophora, Cycliophora, Echinodermata, Entoprocta, Entorrhizomycota, Euglenida, Foraminifera, Gastrotricha, Glomeromycota, Gnathostomulida, Haplosporida, Haptophyta, Hemichordata, Kinorhyncha, Loricifera, Microsporidia, Mollusca, Myzostomida, Nematoda, Nematomorpha, Nemertea, Onychophora, Orthonectida, Phaeophyceae, Picozoa, Placozoa, Platyhelminthes, Porifera, Priapulida, Rhodophyta, Rhombozoa, Rotifera, Streptophyta, Tardigrada, Xanthophyceae Wobei von Streptophyta -> Anthocerotophyta, Marchantiophyta, Bryophyta, Tracheophyta als Phylum im Phylum gefunden und nicht einbezogen wurden und Magnoliophyta als Phylum in Tracheophyta ebenfalls nicht.

Distribution of Taxa

- In the tree we can distinguish 28 different Taxa with the OTL taxonomic tree.
- The most of them are hardly represented. The major taxonomic groups are: ...
- Here you can see some characteristics of the Multifurcation of the tree.

 In a phylogeny, the taxonomic division of the tree is far too coarse, meaning that there



Figure 7.2: Distribution of Nodes in Rank-Cathegories

should be more subtaxa or 'unranked' nodes. But the closer we get to the root, the more the pure taxonomic tree is reflected. If the tree are binary, the taxa would have to double. But the multipliers for some are much bigger and for others much smaller, which you can

```
see in in figure 7.2.
... (see Table 7.2)
extended leaf nodes (real leaf nodes)
```

Distribution of data in the taxa

Mithilfe des taxonomischen Baums von OTL haben wir die Knoten ihren Kingdoms, Phyla und Classes zugeteilt (see Table 7.2).

- 7.3 Missing internal node modeling Residual table
- 7.4 Missing leaf state modeling Residual table
- 7.5 Cross-validation prediction Residual table

Taxa	Number of Nodes	Internal Nodes	Leaf Nodes	Terminal Leaf Nodes
domain	1	1		
kingdom	3	3		
subkingdom	2	2		
infrakingdom	5	5		
superphylum	2	2		
phylum	57	54	3	
subphylum	22	20	2	
infraphylum	2	2		
superclass	8	8		
class	209	175	34	
subclass	120	106	14	
infraclass	14	11	3	
superorder	40	37	3	
order	1014	772	242	
suborder	285	222	63	
infraorder	95	78	17	
parvorder	14	12	2	
superfamily	829	626	203	
family	8449	6143	2306	
subfamily	3090	2250	840	
supertribe	1	0	1	
tribe	1764	1285	479	
subtribe	435	359	77	
genus	164656	105452	59204	
subgenus	1266	869	397	
section	3	2	1	
subsection	1	1	0	
species group	365	288	77	
species subgroup	123	100	23	
species	2247251	7423	2239828	2228993
subspecies	22437	75	22362	22239
variety	1755	7	1748	1726
varietas	5970	19	5951	5909
forma	1181		1181	1181
no rank	954	719	235	7
no rank - terminal	37452		37452	37452
(no entry)	40099	40099		

Table 7.1: TODO: ...

Kingdom (3)	Number of Nodes	Phylum (25)	Number of Nodes	max max height
Metazoa	1 465 207	Arthropoda	1 170 539	54
		Chordata	106 650	74
		Mollusca	80 022	22
		Platyhelminthes	27 141	9
		Nematoda	24 564	23
		Cnidaria	14 878	36
		Porifera	11 737	26
		Echinodermata	10 654	14
		Bryozoa	8 631	11
		Rotifera	3 093	7
		Nemertea	1 793	7
		Tardigrada	1 654	7
		Acanthocephala	1 596	6
		Brachiopoda	1 055	9
		Nematomorpha	633	7
		Chaetognatha	360	7
		Hemichordata	196	5
		Cycliophora	11	5
Fungi	254 871	Ascomycota	157 045	19
		Basidiomycota	92 931	18
		Microsporidia	1 949	6
		Glomeromycota	1 490	6
		Chytridiomycota	1 456	6
Chloroplastida	121 239	Streptophyta	120 731	49
		Chlorophyta	508	6

Table 7.2: TODO: ...

Model / Taxa	Kingdom	Phylum	Class	Order
multifurc \sim taxa	7774454	7435700	7337241	7076068
multifurc \sim taxa $+$ depth	7752303	7431609	7334754	7027578
multifurc \sim taxa $+$ max.height	7730196	7375889	7275856	7005424
multifurc \sim taxa $+$ min.height	7472500	7233486	7144686	6890703
multifurc \sim taxa $+$ mean.height	7304402	7128318	7055313	6815271
multifurc \sim taxa * depth	7714881	7335396	7250759	6843004
multifurc \sim taxa * max.height	7692980	7311241	7187504	6795823
multifurc \sim taxa * min.height	7442387	7177002	7094933	6795099
multifurc \sim taxa * mean.height	7247309	7020258	6965794	6665565

Table 7.3: Residuals of multifurcation models

These models are created with the R function glm() and compared with the anova() function. This results in the listed residuals.

Model / Taxa	Kingdom	Phylum	Class
multifurc \sim taxa	545740	499227	482265
multifurc \sim taxa $+$ depth	544789	493017	478998
multifurc \sim taxa * depth	544062	488366	476382

Table 7.4: Residuals of unknown information models

These models are created with the R function glm() and compared with the anova() function. This results in the listed residuals.

Model / Taxa	Kingdom	Phylum	Class
correctly predicted \sim taxa	117877	111466	108878
correctly predicted \sim taxa $+$ depth	117703	110513	108403
correctly predicted \sim taxa * depth	117592	109827	107994

Table 7.5: Residuals of cross-validation prediction

These models are created with the R function glm() and compared with the anova() function. This results in the listed residuals.