

# Generating Benchmarks for Multiple Sequence Alignments and Phylogenetic Reconstructions

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## Abstract

We present a new probabilistic model of evolution of RNA-, DNA-, or protein-like sequences and a tool *rose* that implements this model. By insertion, deletion and substitution of characters, a family of sequences is created from a common ancestor. During this artificial evolutionary process, the “true” history is logged and the “correct” multiple sequence alignment is created simultaneously. We also allow for varying rates of mutation within the sequences making it possible to establish so-called sequence motifs. The results are suitable for the evaluation of methods in multiple sequence alignment computation and the prediction of phylogenetic relationships.

## Introduction

It is useful for many reasons to have a family of sequences with well-known evolutionary history: For instance, in the study of evolutionary processes, for the evaluation of methods which compute multiple sequence alignments and/or reconstruct phylogenetic trees, and for other tools in computational molecular biology. Unfortunately, nature does not provide “benchmark” problems suited for all these applications.

It is generally not possible to retrieve all sequences of a given sequence family and there is no way to learn the exact phylogeny of the sequences involved. Therefore it is common practice to artificially create sequences with the aim to obtain sequences similar to “real world” biological data. Of course, a family of unrelated random sequences does not provide us with the data that we need since species are part of a hierarchically structured phylogeny.

The simulation of evolutionary processes on the molecular sequence level has a long tradition. Starting with the model of Jukes and Cantor (Jukes & Cantor 1969), several generalizations and alterations have been presented, e.g. (Kimura 1980; Felsenstein 1981;

Hasegawa, Kishino, & Yano 1985; Schöniger & von Haeseler 1995). But in none of the methods known to the authors, the length of the sequences is altered by insertion and/or deletion (*indels*) of subsequences.

We have added indels and “sequence motifs” (patterns in a family of related sequences (Wu & Brutlag 1995)) to the so-called HKY-model (Hasegawa, Kishino, & Yano 1985) to create more realistic sequence families. The data created by our tool *rose*<sup>1</sup> (random-model of sequence evolution) has been extensively tested with our *Divide and Conquer Alignment* (Tönges *et al.* 1996; Stoye 1996) and *GeneFisher* (Giegerich, Meyer, & Schleiermacher 1996; Meyer & Schleiermacher 1996) software packages.

We simulate an evolutionary process by iterated mutation of a “common ancestor sequence” following the edges of a given “mutation guide tree”. This way, the topology of the tree induces the relationships of the sequences. The mutations are performed by insertion, deletion, and substitution of single characters or whole subsequences of the ancestor sequence. In addition to knowing the exact evolutionary *distance* of the sequences, our approach provides us with their whole evolutionary *history*. Therefore, in contrast to biological applications, it is easily possible to verify predictions about phylogenetic relationships drawn from the sequences simply by comparing the predicted phylogeny to the tree that was used in the creation process.

## The Model

Given

- an alphabet  $\mathcal{A}$  of size  $l$ , e.g. the DNA-alphabet  $\{A, C, G, T\}$  or the 20 character amino acid alphabet,
- initial character frequencies  $f_1, \dots, f_l$  satisfying 
$$\sum_{i=1}^l f_i = 1,$$

<sup>1</sup>The software is available on our Bioinformatics Web-Server *BiBiServ* under the following address:  
<http://bibiserv.TechFak.Uni-Bielefeld.DE/rose/>

- a mutation matrix  $M$  of size  $l \times l$  representing pairwise mutation frequencies,
- and a rooted, edge labeled mutation guide tree  $T$ ;

we generate

- a family of  $k$  sequences  $S = \langle \mathbf{s}_1, \dots, \mathbf{s}_k \rangle$  with average length  $n$  and average pairwise evolutionary distance  $d_{av}$ ,
- a multiple sequence alignment  $A$  of the sequences  $\mathbf{s}_1, \dots, \mathbf{s}_k$  that is optimal with respect to the creation process, i.e. it reflects the “true” evolutionary history of  $\mathbf{s}_1, \dots, \mathbf{s}_k$ ,
- and a *relatedness tree*  $T'$  representing the phylogenetic relationship of the created sequences.

The underlying idea of our method is the following: First of all, a common ancestor sequence of length  $n$  is created and attached to the root of the tree  $T$ . Then, the nodes connected with the root are filled with “child” sequences created from the root sequence by mutations where the length of the edge stands for the evolutionary distance between the sequences. This process is continued until the leaves of  $T$  are reached and thus all nodes of the tree are filled with a sequence. Finally, from the sequences created this way, the required number of  $k$  sequences is selected in a random manner. These sequences form the family  $S = \langle \mathbf{s}_1, \dots, \mathbf{s}_k \rangle$ .

The multiple sequence alignment  $A$  is created by replacing deleted subsequences with gap characters in the selected sequence and inserting gap characters into all but the selected sequence for insertions. In this way we generate an alignment that reflects the “true” evolutionary history of the sequences.

$T'$  is the smallest subtree of  $T$  which contains all the nodes corresponding to the selected sequences (and possibly some additional inner nodes which can be seen as extinct ancestors).

### The Root Sequence

Each of the  $n$  positions in the root sequence is filled by a random process that takes the character frequencies  $f_1, \dots, f_l$  for the given alphabet into account. For amino acid sequences, we implemented as default values the *normalized frequencies* of the amino acids given in (Dayhoff, Schwartz, & Orcutt 1979) and for nucleotides we use the frequencies given in (Agarwal & States 1996).

Alternatively, we also allow a pre-given root sequence.

### Creation of Child Sequences

Starting from a given sequence  $\mathbf{s}_{old}$ , we create a new sequence  $\mathbf{s}_{new}$ . The following steps are used to create a new “descendant” sequence:

1. We apply the mutation function *mutate* for the given alphabet to every position  $i$  in  $\mathbf{s}_{old}$ :

$$\mathbf{s}_{new}[i] = \text{mutate}(\mathbf{s}_{old}[i], b)$$

where  $b$  is the branch length for leading to the new node. The mutation matrix is selected with respect to  $b$ , please see below.

2. We delete one or more subsequences from  $\mathbf{s}_{new}$ ; an arbitrary deletion probability  $p_{del}$  and arbitrary deletion length function  $l_{del}$  can be specified.
3. We insert one or more gaps of arbitrary length at arbitrary positions in  $\mathbf{s}_{new}$ ; again, an insertion probability  $p_{ins}$  and insertion length function  $l_{ins}$  can be specified. The characters inserted maintain the initial character distribution.

Note that the average sequence length remains  $n$  only if  $p_{ins} = p_{del}$  and  $l_{ins} = l_{del}$ .

In case the mutation matrix  $M$  is the probability matrix of one *accepted amino acid substitution per hundred sites* (1 PAM) given in (Dayhoff, Schwartz, & Orcutt 1979) – which is our default for proteins – we denote this new unit of measure for the distance of a child sequence from its ancestor including insertions and deletions by 1 PAM\* where the parameters for insertions and deletions have to be specified additionally.

Evolutionary rates of more than 1 PAM\* are obtained by applying the creation procedure repeatedly. As Schöniger and v. Haeseler (Schöniger & von Haeseler 1995) have shown, the use of a custom matrix (such as PAM 10) helps to save time when the number of substitutions exceeds an upper bound. At each step along an edge of the guide tree, depending on the mutation rate, the decision is made either to use one of the precomputed PAM\* matrices or to compute a new custom matrix in order to save time.

### Sequence Motifs

Up to this point, we have assumed a constant rate of mutation over the whole length of the sequences. However, this is not very realistic: The mutation rate of genomic sequences found in nature is not constant for all positions in the genome. Mutations in regions with strong functional and/or structural importance are less often observed.

Therefore, we implemented a feature that allows the use of different rates of mutation for different regions

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(a) FSAEALVSPGKGDDEQVPNKDKCVYHGHKDGKRMNVKTPPTGPLVVGWHQ
    YEGANEVGATCEESSYCVVKEQAIQVKESQECTDFARHEVKSFRGVPGLTEVIPVPL
    YGAAHPVGDPIKLGSLFLNHYESKGHTAAMCLLGMKTELIEPIEVQA
    SGVTEPVPNPVVPATGIKLDKYTREENCLGMCLMGMGPPMVTIGEVGI

(b) FSAEALVSP-----GKGDDEQVPNKDKCVYHGHKDGKRMNVKTPPTGPLVVGWHQ
    YEGANEVGATCEESSYCVVKEQAIQVKESQECTDFARHEVKSFRGVPGLTEV-IPVPL
    YGAAHPVGDPIKLGSLFLNH---YESKGHTAAMCLLGMKTELIEP-IEVQA
    SGVTEPVPNP-----VPATGIKLDK---YTREENCLGMCLMGMGPPMVTI-GEVGI

(c) FSAEALVSP-----GKGDDEQVPNKDKCVYHGHKDGKRMNVKTPPTGPLVVGWHQ
    YEGANEVGATCEESSYCVVKEQAIQVKESQECTDFARHEVKSFRGVPGLTE-VIPVPL
    YGAAHPVGDPIKLGSLFL---NHYESKGHTAAMCLLGMKTELIE-PIEVQA
    SGVTEPVPNP-----VPATGIKLDK---DKYTREENCLGMCLMGMGPPMVT-IGEVGI

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Figure 1: (a) Sample family of random sequences obtained with the procedure described in the text for  $n = 50$  and  $k = 4$ ; (b) “true” alignment of these sequences; (c) an optimal alignment according to PAM 250 substitution matrix and gap function  $g(l) = 8 + 12l$ .

of the sequence. We use a vector to specify the degree of conservation at every position in the sequence. This way it is possible to establish well-conserved motifs in the whole sequence family created.

### The Mutation Guide Tree

In the general case, the user can specify any rooted, edge-labeled mutation guide tree. Given a mutation guide tree, the average sequence distance  $d_{av}$  (in units of PAM\*) can be computed, i.e. the expected length of a shortest path between two randomly chosen nodes in the tree. If no tree is entered by the user, we compute a pruned binary mutation guide tree for a user defined average sequence distance.

### Example: A Protein Sequence Family

The following example shows some of the features of our approach. In Figure 1 (a), a sample family with  $k = 4$  sequences of average length  $n = 50$  is shown. This family is created with the default settings of *rose*: A uniform binary mutation guide tree of depth  $m = 9$  and uniform edge length  $R = 18$  PAM\*. The probability for insertions and deletions is set to  $p_{ins} = p_{del} = 0.3\%$  and the insertion and deletion length functions are exponentially decreasing with a maximal length value of 10. These parameters can be shown to yield a family of amino acid sequences with average sequence distance of  $d_{av} = 250$  PAM\* (Stoye 1997).

The alignment given in Figure 1 (b) is the “true” alignment corresponding to the creation process of the sequences. Figure 1 (c) shows the “optimal” alignment according to the PAM 250 substitution matrix (Dayhoff, Schwartz, & Orcutt 1979) (in distance form with values between 0 and 24) and gap

function  $g(l) = 8 + 12l$  computed with the program MSA (Lipman, Altschul, & Kececioğlu 1989; Gupta, Kececioğlu, & Schäffer 1995). While the overall optimal alignment is correct, the exact location of the gaps does not coincide in all cases. The scores for both alignments show these differences as well: The “true” alignment has an alignment score of 5184, while the “optimal” alignment a has a score 5166. This shows that – as is well-known – an optimal alignment is not necessarily the correct one.

Figure 2 shows the corresponding relatedness tree  $T'$  of these sequences.

### Conclusion

The data sets created by *rose* are the first artificially created sequence families that contain both *indels* and *motifs*. The evaluation of multiple sequence alignment tools and phylogenetic reconstruction tools is now possible with the benchmarks created.

However, we still have used some approximations: While we do not assume that the characters of the sequences evolve independently and with the same rate in the whole family, we have not yet included a feature that simulates different rates of evolutionary pressure in different branches of the tree, enabling different lineages to evolve independently within our tree. This has been observed by a number of biologists several times (Greer 1981; 1990; Schulz *et al.* 1986; Benner, Cohen, & Gonnet 1994). While we are planning to include this feature in a future release of *rose* and extend the scope of our model even further, it is important to note that simulations can only aid the evaluation of algorithms. The use of *real* sequences is indispensable for the evaluation of software in Bioinformatics.

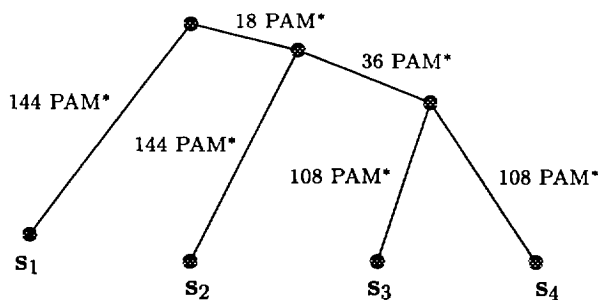


Figure 2: Relatedness tree for the sequences shown in Figure 1.

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