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## Exercise Sheet 5 zur Vorlesung Computational Biology (Part 2), WS 12/13

Hand In: Until Monday, 07.01.2013, 10:00 am, email to wild@cs... or in lecture.

## Exercise 11

3 Points

Design an algorithm that computes a *perfect phylogenetic tree*—or shows that none exists—for n taxa and m binary characters<sup>1</sup> in time  $\mathcal{O}(m \cdot n)$ . Your solution should comprise

- a clear description of the algorithm (pseudocode or prose is sufficient),
- a proof of its correctness
- and an analysis showing that the algorithm satisfies the given time bound.

You can built upon the algorithm sketched in lecture, but make sure to argue precisely how your algorithm stays in  $\mathcal{O}(m \cdot n)$ .

## Exercise 12

Let  $\mathcal{T} = \{T_1, \ldots, T_m\}$  be a set of trees obtained by repeated execution of steps 2-4 of the *quartet puzzling* algorithm (script page 182). We *root* all trees in  $\mathcal{T}$  by the following procedure: First, we pick an arbitrary, but fixed taxon a. Then in each  $T_i$ , we insert a new *artificial* node in the middle of the (unique) egde to the leaf corresponding to a and declare this new node the root of this tree.

Now, we label the resulting trees bottom-up: A leaf for taxon  $a_i$  is assigned label  $\{a_i\}$  and each inner node's label is the union of the labels of its children. Note that these labels fully describe the topology of a tree, so it suffices to consider the set of labels for each tree  $T_i$ .

From these *m* label sets, we now pick all labels that occur in  $> \frac{m}{2}$  label sets. Show that the resulting collection of labels always encodes a (not necessarily binary) tree *T* whose leaves exactly correspond to the taxa.

This means, T is a valid phylogenetic tree for the given taxa. It is called the *majority* consensus tree for  $\mathcal{T}$ .

2 Points

<sup>&</sup>lt;sup>1</sup>deutsch: Merkmale