CS 466 – Introduction to Bioinformatics Lecture 16

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These notes are based on Ref. [1].

We are given a binary matrix $B \in \{0,1\}^{n \times m}$ with n taxa (think of them as species) and m characters. We say that taxon $f \in [n]$ possesses character $c \in [n]$ if $b_{f,c} = 1$. We consider the large maximum parsimony phylogeny problem. That is, given matrix B, infer a phylogenetic tree T where each taxon in B uniquely corresponds to a leaf of T and the internal vertices of T are labeled by n binary characters with minimum parsimony score (i.e. the number of state changes is minimum). In general, this problem is NP-hard but we consider a version of this problem with two additional constraints. First, the root vertex of T must have state 0 for each character. Second, each character changes state from 0 to 1 only once on the tree and never reverts back from 1 to 0. The latter constraint is known as the infinite sites assumption in populations genetics, and also known as the two-state perfect phylogeny model. Let's formalize this.

Definition 1. A rooted tree T with n leaves is a two-state perfect phylogeny for a given binary matrix $B \in \{0,1\}^{n \times m}$ provided:

- (i) each taxon (row of B) labels only one leaf,
- *(ii)* each character labels only one edge, and
- (iii) only the characters possessed by a taxon (leaf of T) are present on the unique path to the root.

Importantly, there need not exist a two-state perfect phylogeny for a given matrix B. This gives rise to the following problem.

Problem 1 (Two-state Perfect Phylogeny). Does a given binary matrix $B \in \{0, 1\}^{n \times m}$ have a two-state perfect phylogeny T? If so, construct T.

The way that this problem is posed should remind you of the large additive distance phylogeny problem, where solutions where edge-labeled trees that generated a given distance matrix D. The question that we asked there was to decide if D is additive. In our first attempt, we had only a constructive definition of solutions (relying on the existence of a tree T that generates D). Only later did we identify the four-point condition as a *complete characterization* of the solution set.

Similarly, here, we only have a constructive definition of binary matrices that are twostate perfect phylogeny matrices. This is not good enough! We want to identify a condition Φ that is both necessary and sufficient for a binary matrix B to be generated by a two-state perfect phylogeny tree T. That is, should T be a two-state perfect phylogeny for B then B must satisfy Φ (necessary); and should B satisfy Φ then there must exist a two-state perfect phylogeny T for B (sufficient). If Φ is necessary and sufficient then we say that Φ is a *complete characterization* of the set of two-state perfect phylogeny matrices. Can we find such a condition?

The answer is yes. There exists a condition Φ that can be computed in O(nm) time. It will be helpful to sort the columns of B by the number of ones they contain, in descending order (largest first). Ties are broken arbitrarily. Let \overline{B} denote the sorted binary matrix. We make the following observation, which follows directly from Definition 1.

Observation 1. Let $\overline{B} \in \{0,1\}^{n \times m}$ be obtained from $B \in \{0,1\}^{n \times m}$ by sorting columns of B in descending order by the number of ones they contain. Matrix B has a two-state perfect phylogeny if and only if matrix \overline{B} has a two-state perfect phylogeny.

Moreover, identical columns can be trivially removed.

Observation 2. Let $B' \in \{0, 1\}^{n' \times m}$ be obtained from $B \in \{0, 1\}^{n \times m}$ not containing repeated columns present in B (clearly, $n' \leq n$). Matrix B has a two-state perfect phylogeny if and only if matrix B' has a two-state perfect phylogeny.

We have the following definition.

Definition 2. Binary matrix $B \in \{0,1\}^{n \times m}$ is conflict free if no pair of columns c and d contain the three binary pairs (0,1), (1,0) and (1,1).

Clearly, using a naive algorithm we can check in $O(n^3m^2)$ if a matrix B is conflict free. We have the following lemma.

Lemma 1 (Shared-prefix property). Let d be the rightmost column in \overline{B} possessed by two taxa f and g. Then, if no pair of columns conflicts then f and g must be identical from column 1 to column d.

Proof. By the premise, we have $\bar{b}_{f,d} = \bar{b}_{g,d} = 1$. Consider a column c < d possessed by f. That is, $\bar{b}_{f,c} = 1$. Since the columns are distinct and are sorted by the number of ones they contain, we have that c and d must also contain the binary pair (1,0), say in taxon h. We thus have the following situation:

taxon	c	d
f	1	1
g	?	1
h	1	0

By the premise, we have that $\bar{b}_{g,c}$ cannot be equal to 0, as that would introduce a conflict. Hence, taxon g must also possess character c, i.e. $\bar{b}_{g,c} = 1$. Now the choice of considering taxon f first was arbitrary, and the same argument can be given when g possesses a column c < d. In other words, when either f or g possess a character c then the other taxon must possess c as well. Hence, taxa f and g are identical from columns 1 to d.

We have the following theorem.

Theorem 1. Matrix B has a two-state perfect phylogeny tree if and only if B is conflict free.

Proof. (\Rightarrow) We start with the forward direction. Let *T* be a two-state perfect phylogeny tree for *B*. Consider two characters *c* and *d*. Let e_c (e_d) be the edge where *c* (*d*) was introduced. By Definition 1, taxa that possess *c* (or *d*) must be present as leaves below the edge e_c (or e_d). We distinguish four cases.

1. $e_c = e_d$.

There cannot be a taxon with state (1,0) or (0,1) for the considered characters (c,d), as taxa that possess either c or d are in the same subtree below $e_c = e_d$.

2. The edge e_c is on the unique path from the root to e_d .

There cannot be a taxon with state (c, d) = (0, 1), as character c was introduced prior to character to d in T.

3. The edge e_d is on the unique path from the root to e_c .

There cannot be a taxon with state (c, d) = (1, 0), as character d was introduced prior to character to c in T.

4. The two unique paths from the root to e_c and e_d are edge disjoint.

There cannot be a taxon with state (c, d) = (1, 1), as T does not contain a path from the root containing both character c and d.

Hence columns c and d are conflict free. Since we chose c and d arbitrarily, matrix B itself is conflict free (recall that conflict-free definition considers all pairs of column).

 (\Leftarrow) We use Observation 1 and 2 and consider without loss of generality a sorted matrix B obtain from B that does not contain any repeated columns. Observe that in any two-state perfect phylogeny T for B it must hold that the characters label the edges of the unique path from the root to a taxon f are exactly the characters that taxon f possesses. Moreover,

the characters that taxon f possesses will appear in the same order in which they occur in \hat{B} . To see why, suppose that f possesses characters c < d. Per the previous statement, characters c and d label edges on the unique path from the root to f. Now \hat{B} contains more 1s for character c than for character d. Thus, the edge e_c must occur prior to e_d . Hence, the characters that a taxon f possesses will appear in the same order in which they occur in \hat{B} . This must hold for any two-state perfect phylogeny T for \hat{B} . Thus, all that remains to show is that the n paths for each taxon can assembled into a single tree if \hat{B} is conflict free.

We show constructively how to construct a two-state perfect phylogeny T for a conflictfree matrix \hat{B} . The algorithm will construct T one row at a time. Initially, we create a root vertex. Next, we consider taxon 1. We construct a path T_1 composed of labeled edges for each character possessed by this first taxon maintaining the order imposed by \hat{B} . We extend the path with an unlabeled edge leading to a new vertex that will correspond to the first taxon. Clearly, T_1 is a two-state perfect phylogeny tree for taxon 1.

Let T_f be the partial tree constructed from taxa 1 to f, and assume inductively that T_f is a two-state perfect phylogeny tree for the first f taxa in \hat{B} . We now describe how to construct T_{f+1} . We traverse the edges in T_f starting from the root walking down the tree as long as the traversed edges contain characters that are possessed by f in the same order as in \hat{B} . Let v be the last vertex visited on this traversed path, and let c denote the last matched character. As T_f is a two-state perfect phylogeny, this path is unique. We then create a new path extending from v and containing all characters d > c that are possessed by f + 1 that have not been matched. This new path maintains the order of the columns in \hat{B} . Finally, we extend the new path by one edge, leading to a new leaf that corresponds to f + 1. We claim that T_{f+1} is a two-state perfect phylogeny tree for the first f + 1 taxa in \hat{B} .

First, observe that each path path to a leaf $h \leq f + 1$ in T_{f+1} contains exactly the characters that taxon h possesses. Moreover, no character on the path to v is anywhere else in T_{f+1} , as T_f is a two-state perfect phylogeny. Thus, we only need to show that none of the characters that are in the new path from v to f + 1 are in T_f . Let d be the rightmost character in \overline{B} that taxon f + 1 possesses and that is also possessed by a taxon in T_f . Let e_d denote the edge in T_f labeled by d. By definition, any taxon (leaf) h that is below e_d possesses d. We can apply the shared-prefixed property as \hat{B} is sorted and conflict-free. Thus, by the shared-prefix property, rows h and f + 1 are identical from column 1 to d. As such, the walk from the root to v is also a walk from the root to h. Moreover, by the choice of d, taxa h and f + 1 do not possess any other common character e > d. Thus, none of the characters that are in the new path from v to f + 1 are in T_f . Hence, T_{f+1} is a two-state perfect phylogeny from the first f + 1 taxa of \hat{B} .

When all taxa have been processed the resulting tree is thus a two-state perfect phylogeny for \hat{B} , and in turn for B upon re-introducing characters that correspond to repeated columns (introducing multiple labels per edge).

References

[1] Dan Gusfield. ReCombinatorics: The Algorithmics of Ancestral Recombination Graphs and Explicit Phylogenetic Networks. The MIT Press, 2014.