TOWARD DEFINING THE COURSE OF EVOLUTION: MINIMUM CHANGE FOR A SPECIFIC TREE TOPOLOGY

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Abstract

Fitch, W. M. (Dept. of Physiological Chemistry, Unio. of Wisconsin, Madison, Wisconsin, 53706), 1971. Toward defining the course of evolution: minimum change for a specific tree topology. Syst. Zool., 20:406-416.—A method is presented that is asserted to provide all hypothetical ancestral character states that are consistent with describing the descent of the present-day character states in a minimum number of changes of state using a predetermined phylogenetic relationship among the taxa represented. The character states used as examples are the four messenger RNA nucleotides encoding the amino acid sequences of proteins, but the method is general. [Evolution; parsimonious trees.]

It has been a goal of those attempting to deduce phylogenetic relationships from information on biological characteristics to find the ancestral relationship(s) that would permit one to account for the descent of those characteristics in a manner requiring a minimum number of evolutionary steps or changes. The result could be called the most parsimonious evolutionary tree and might be expected to have a high degree of correspondence to the true phylogeny (Camin and Sokal, 1965). It's justification lies in the most efficient use of the information available and does not presuppose that evolution follows a most parsimonious course. There are no known algorithms for finding the most parsimonious tree(s) apart from the brute force method of examining nearly every possible tree. This is impractical for trees involving a dozen or more taxonomic units. Most numerical taxonomic procedures (Sokal and Sneath, 1963; Farris, 1969, 1970; Fitch and Margoliash, 1967) provide dendrograms that would be among the more parsimonious solutions; one just cannot be sure that a more parsimonious tree structure does not exist. Farris (1970) has explicitly considered the parsimony principle as a part of his method which, like the present method, has its roots in the Wagner tree (Wagner, 1961, 1965).

A problem subordinant to finding the most parsimonious tree(s) is finding all possible, most parsimonious assignments of the information to any given particular tree. This is the problem that is treated herein. Its solution provides the most reasonable hypotheses on the ancestral states and therefore is the best estimate of the course of evolution. The biological characteristics to be used are the nucleotides of orthologous genes but the method is applicable to any data for which the underlying assumptions are acceptable. Thus, given a set of descendant nucleotide sequences and a topology presumed to describe their ancestral relationships one can set forth: a), the exact number of nucleotide replacements that are the minimum necessary to account for the descent of those sequences from a common ancestor; b), all possible ancestral

1 Orthologous is used, as previously defined (Fitch, 1970), to denote that particular subset of homologous genes for which there is an exact one-to-one correspondence (ortho = exact) between the ancestral relationships of the genes and the ancestral relationship of the species (or other taxonomic unit) in which those genes are found. It may be distinguished from paralogous which is used to denote homologous genes that arose via a gene duplication and descended side by side (para = in parallel) in any one line of descent. An example of the latter would be the genes for α and β hemoglobin.
sequences at each node (branch point) that are consistent with that minimum; c), those combinations of nodal ancestral sequences that are consistent with that minimum number of replacements; and d), a way of assigning relative weights among the various alternative nucleotide replacements required of the various internodal periods of the several minimal trees.

METHOD

The method assumes that: a), a set of orthologous descendant sequences is available; b), the ancestral (topological) relationships among them are known; and c), any nucleotide may replace any other nucleotide at any position without regard to the nature of prior replacements in that position or elsewhere. The word "may" indicates that all logical options are to be considered; this is not to deny the possibility that selective forces may have precluded some of those options.

The character state of a given nucleotide position is the set of nucleotides that might be present, A, C, G or U or some combination thereof. Ambiguity may arise because one is uncertain which codon is used in the gene. For example, arginine may be encoded by either AGG or CGG (among others) and we can not be sure, knowing only that arginine was encoded, whether the first nucleotide position is A or C. Consequently, the character state is represented by the set AC to indicate the exact range of ambiguity. Ambiguity will also arise from uncertainty about ancestral character states and will be similarly represented.¹

¹Another type of ambiguity concerns missing information which may be missing for one of two reasons, viz. either the character was not examined or the character does not exist. These suggest different treatments. Where the character was not examined, it may be best to represent the state as completely ambiguous (e.g., ACCU) or, in set theoretic terminology, as the universal set. The result is that the character has no influence whatsoever on the procedures described or on the results obtained. Where the character does not exist, it may be best to represent the state by the empty (null) set. The result is that a change of state will be detected (viz, the deletion of the character) but there is no other effect on the procedures or results. One must exercise some discretion, however, since, if five contiguous codons were eliminated in a single genetic event by unequal crossing over and all 15 (3 x 5) nucleotide positions assigned to empty set state, the final result would show 15 changes of state where only one deletion occurred.
Fig. 1.—Reconstruction of Ancestral Nucleotide Forms. On the left side, Figure 1a shows the preliminary reconstruction beginning with an assumed topology and an assignment of nucleotides to the present taxonomic units at the bottom. Each ancestral set is composed of those nucleotides common to its immediate descendants, if any, otherwise it is composed of all of them. Arabic numerals indicate the set number and their order of construction. Figure 1b on the right shows the result when the preliminary phase has been converted to the final phase by the method described in the text. Nucleotides in parentheses have been added.

Reconstruction of possible ancestral nucleotides—preliminary phase

The preliminary reconstruction has been previously described (Fitch, 1970). That reconstruction proceeds from the descendent character sets by formulating an ancestral character set for an immediate ancestor and working backward, one successive node at a time, until finally the nodal character set for the most distant ancestor has been formed. The formulation of each nodal set follows the following simple rule: The preliminary nodal set shall be comprised of all characters (nucleotides) common to both immediately descendent sets; if none are common to both, then the preliminary nodal set will be comprised of all characters found in either. In mathematical terms, the nodal set is the intersection of its immediately descendent sets if the intersection exists (i.e., is not empty) otherwise it is the union of those sets. An example is shown in Figure 1a (on the left). Because there are so many different nucleotides in only 6 taxonomic units, many of the descendants have no nucleotides in common, with the result that the first (lowest) three preliminary nodal sets are formed by unions. At the penultimate node [4], the intersection CU ∩ ACU = U and so U is the preliminary nodal set. The ultimate node is once again a union. The example is chosen for its completeness in representing possible problems rather than for its representativeness. With real amino acid sequences such as the cytochromes c, the vast majority of the nodal sets are simpler in that they are formed by intersections of identical elements rather than by unions as in this case. It should be noted that for every occasion that a union is required to form the preliminary nodal set, a mutation (nucleotide replacement) must have been fixed in this nucleotide position at some point during
Fig. 2.—Deficiencies of Preliminary Phase Reconstruction. In each case, the trees on the left (2a top, 2c middle and 2e bottom) show the results of a preliminary phase reconstruction of ancestral nucleotides. In each instance, there is an interpretation, shown on the right, which is not inherent in the preliminary reconstruction. See text for explanation. Arrowheads are shown on segments to indicate the occurrence of nucleotide replacements.

the evolution of this position. Thus counting the number of unions gives one the minimum number of fixations (or changes of state) required to account for descendent nucleotides (characters) from a common ancestor, given the phylogeny assumed at the outset.

Reconstruction of possible ancestral nucleotides—final phase

To understand why there is a final phase following the preliminary phase, it is necessary to understand in what respects that preliminary phase may be deficient. These are shown in Figure 2.

In Figure 2a (upper left) is shown a preliminary phase reconstruction of a position from three taxonomic units. The ambiguous AC shown at the lower ancestral node represents an initial inability to decide whether the ancestral form was an A that was replaced by a C in the line of descent to the middle taxonomic unit or was a C that was replaced by an A in the leftmost line of descent. The only certainty is that a replacement is required. The third taxonomic unit, possessing an A, requires the ultimate ancestor to possess the A unless we postulate additional replacements beyond the minimum required to account for
the data. But this additional information about the upper ancestral node also makes it clear that the first node can not then be a C. The only formulation that will permit the descendent positions to be accounted for in a single replacement requires that replacement to be from A to C in the descent from the first node as shown in Figure 2b (upper right). The elimination of the C from the first node is determined by what may be called the *rule of diminished ambiguity*. Its precise formulation is encompassed in steps I and II of the algorithm, to be presented further on, that contains the complete set of rules for the final phase of reconstructing the nodal sets.

In Figure 2c (middle left) is shown another preliminary phase reconstruction which accounts, using two replacements, for the descent of the characters of the three taxonomic units given. Figure 3d (middle right), however, shows an equally adequate solution which is not encompassed by the possible alternatives available in Figure 3c. Clearly C is a valid alternative for the first node. This case is encompassed by the *rule of expanded ambiguity* which is precisely described in steps III and IV of the forthcoming algorithm.

In Figure 2e (lower left) is shown a third preliminary phase reconstruction that accounts for four descendants using two replacements. In Figure 2f (lower right) is an equally valid solution. Indeed, the C at the lowest node in the preliminary reconstruction is a valid alternative to the A if and only if a C is allowed at the penultimate node above. It is characteristic of this type of case that two nodes, separated by a single node, both contain a nucleotide not present in the intervening node because of the intersection process. Hence, this is called the *rule of encompassing ambiguity* which is formulated as step V of the forthcoming algorithm.

In the preliminary phase, the nodes in Figure 1 were formulated in the order of increasing ancestral remoteness (1→5, with the order for formulating nodes 1 and 2 being arbitrary). In the final phase, the order for correcting the nodal sets must be reversed (5→1).

The preliminary set for the ultimate node is made the final set for that node. We then go to the penultimate node (4 in this case) and proceed according to the following six step algorithm.

I. If the preliminary nodal set contains all of the nucleotides present in the final nodal set of its immediate ancestor, go to II, otherwise go to III.

II. Eliminate all nucleotides from the preliminary nodal set that are not present in the final nodal set of its immediate ancestor and go to VI.

III. If the preliminary nodal set was formed by a union of its descendent sets, go to IV, otherwise go to V.

IV. Add to the preliminary nodal set any nucleotides in the final set of its immediate ancestor that are not present in the preliminary nodal set and go to VI.

V. Add to the preliminary nodal set any nucleotides not already present provided that they are present in both the final set of the current ancestor and in at least one of the two immediately descendent preliminary sets and go to VI.

VI. The preliminary nodal set being examined is now final. Descend one node as long as any preliminary nodal sets remain and return to I above.

Figure I illustrates the operation of the algorithm. The left hand side (Figure 1a) depicts the preliminary nodal sets. The ultimate ancestral nodal set 5 (AU) is considered the final set and we turn our attention to preliminary nodal set 4. This nodal set does not contain an A and therefore, according to step I, we proceed to step III. Nodal set 4 was not formed by a union and therefore we are directed by step III to go to step V. Following the directions of step V we discover that A is present in both nodal sets 3 and 5 (the rule of encompassing ambiguity) and must therefore be added to nodal set 4. Mathematically, \((1 \cap 5) \cup (3 \cap 5)\) = AU.
Since A is not present in nodal set 4, it must be added. The A is shown in parentheses to denote the fact that it was added in the final phase. We now proceed to step VI which tells us we are finished with node 4 and may proceed to node 3 and step I.

Preliminary nodal set 3, AGU, contains all of the nucleotides in its immediate ancestor and so we are directed by step I to go to step II. Step two tells us to eliminate all nucleotides in nodal set 3 not present in nodal set 4 (the rule of diminished ambiguity). Hence G is removed from nodal set 3 and we proceed to step VI which tells us nodal set 3 is now final and we may proceed to node 2 and step I.

Preliminary nodal set 2, GU, does not contain all of the nucleotides in its immediate ancestor and so we are directed by step I to go to step III. Because nodal set 2 was formed by a union, step III directs us to step IV. According to step IV, we must add any nucleotides in final nodal set 3 not present in preliminary nodal set 2 (rule of expanded ambiguity). Hence A is added and we proceed to step VI which tells us nodal set 2 is now final and we may proceed to node 1 and step I.

Preliminary nodal set 1, CU, like nodal set 2 before it, will pass through steps I, III, IV and VI, acquiring an A by the rule of expanded ambiguity. Since there are no further preliminary nodal sets remaining, the final phase of reconstruction is completed.

Permitted links between nucleotides in successive nodal sets

The ancestral forms have now been reconstructed such that all nucleotides that
could possibly occupy a given position in an ancestral sequence are shown. Any other nucleotides would require the postulation of additional replacements to account for final descendant forms. Even within the framework of the permitted nucleotides, not all replacements are allowable as one descends from node to node if one is to obtain a most parsimonious tree.

Figure 3 is a redrawing of the ancestral nodes of Figure 1b with links connecting those nucleotides that comprise valid lines of descent in a most parsimonious tree. The arrowheads are on those internodal links that involve a change of nucleotide state in descending from one node to the next. The subscripts give the number of nucleotide replacements required in those cases where one or both immediate descendants are not other ancestral nodes. Since node 4 has only other ancestral nodes for its immediate descendants, its nucleotides have no subscripts. The total number of replacements required must be four since there were unions used to construct nodes 1, 2, 3 and 5 as shown in Figure 1a. In Figure 3, a complete phylogeny will require a network composed of one nucleotide from every node using only the links shown. Such a complete linkage will involve nucleotides with subscripts and possibly links bearing arrowheads. The sum of the arrowheads plus subscripts on any complete valid linkage is four, the minimum number of replacements required to account for the data. There are 11 such linkages. One of them begins with the U in the ultimate ancestor, the other ten begin with the A. Of the ten beginning with A, one involves the replacement of A by U in the descent to node 4. The remaining are the 9 possible combinations of the three alternatives in descending to node 2 from an A in node 3 and the three alternatives in descending to node 1 from an A in node 4.

How then does one discover all possible valid linkages such as those shown in Figure 3? First of all, the nucleotides without parentheses were placed in the nodal set during the preliminary phase of reconstruction while those with parentheses were placed there during the final phase. Such nucleotides will be represented as $N_i$ and $(N_i)$ where the subscript denotes one of the four nucleotides. We shall use an arrow ($\rightarrow$) to denote descent from an ancestor to the next node. Thus $(N_i) \rightarrow N_j$ means that an ancestral nucleotide that was added to a nodal set during the final phase of reconstruction is replaced (since $i \neq j$) by a nucleotide that was originally assigned to the descendant nodal set in the preliminary phase. An example is shown by the $(A) \rightarrow C$ change in Figure 3.

The rules then are as follows: Given that a particular ancestral nucleotide is i,

I. $N_i \rightarrow N_i$ or $(N_i) \rightarrow N_i$ is obligatory if the descendant $N_i$ exists; if the descendant $N_i$ does not exist, then

II. All possible linkages are permitted except $N_i \rightarrow (N_i)$ and $(N_i) \rightarrow (N_i)$.

The interpretation of these rules may be seen in Figure 3. The first rule states that, given a particular ancestral nucleotide, there is no option but to link that nucleotide to the identical nucleotide in the descendant nodal set if that descendant nucleotide was placed there in the preliminary phase of the reconstruction. It is the operation of this rule that causes there to be only one valid complete linkage for Figure 3 when the ultimate ancestral form is a U. Another way of phrasing this rule is: A nucleotide not in parentheses is the only valid terminal point of a link which originates from the same nucleotide in its immediate ancestor.

When the first rule does not apply, then and only then does the second rule apply. As a consequence, except for the descent to the A at node 3, all ancestral A's are permitted to descend to any nucleotide in the nodal set of its immediate descendant.

None of the exceptions to rule II is shown in Figure 3. One may illustrate the exceptions though by imagining two cases. If the nodal sets were correct in Figure 3 except for the addition of a C to nodal set 4, that G could link to C or U in nodal set 1 but not to (A) because $N_i \rightarrow (N_i)$ is forbidden.
For the two links from $^8A$, $P[L(\wedge A)] = P(\wedge A)/2 = 0.25$. The descendent nodal nucleotides have probabilities associated with the sum of the probabilities for the links that terminate upon them. Thus $P(\wedge A) = 0.25$ and $P(U) = 0.25 + 0.5 = 0.75$. Note that the sum of the probabilities for the nucleotides in any given nodal set must equal one and the same is true for the sum of the probabilities for the links between any two consecutive nodes. Figure 4 depicts the probabilities associated with every link and every nodal nucleotide shown in Figure 3. Also shown, by dotted lines, are the probabilities associated with those links to a final descendant where a nucleotide replacement was required. The result is that four necessary nucleotide replacements are distributed among those that form a part of a most parsimonious tree as follows: $A \rightarrow C$, $2/3$; $A \rightarrow G$, $1/6$; $A \rightarrow U$, $7/12$; $C \rightarrow U$ and $G \rightarrow U$, $1/12$; $U \rightarrow A$, $3/4$; $U \rightarrow G$ and $U \rightarrow C$, $5/6$.

The following relationships may be noted as indicating the "reasonableness" of this procedure. (I) C and G are each present in only a single taxonomic unit and are given only low probabilities of having existed at ancestral nodes. That is to say, where a phenotype is singular we would expect it to have arisen since the most recent bifurcation. (II) A, which has no representatives among the descendants of nodes 1 and 2 is given only a small probability of being the ancestral form at these nodes. The probability of the A, C and G nucleotides representing the ancestral form in nodes 1 and 2 for the case where all complete linkages are equiprobable would be more than three times as great as those shown in Figure 4.
DISCUSSION

A method for reconstructing ancestral sequences was first presented by Fitch and Margoliash (1967). That early method was also based upon the concept of minimum evolution but was less formal in that all possible solutions might not necessarily be recognized. Moreover, in order to decide among several equally parsimonious alternatives, the expected influences of selection were invoked. For example, Figure 2f shows one of three possible ways of accounting for the descent (on the topology given) of the four nucleotides presented. Clearly, this requires the same replacement, C→A, to have occurred twice. We may consider this pair of replacements to represent a parallel fixation. If, however, the two lower ancestral nodes are set equal to A rather than C while leaving the ultimate ancestral node at C, then we are indicating that a distant C→A replacement was subsequently followed by an A→C replacement. We may consider this pair to represent a back mutation (fixation). We believe that selection was more likely to discover the utility of a mutation in two closely related, and possibly contemporaneous, lines of descent than that a mutation once found beneficial should then become deleterious relative to its previous ancestral form. This remains our belief to the extent that selection is the operative mechanism. However, in view of the recent speculation about the possibility of neutral mutations (Kimura, 1968; Smith, 1968; Arneheim and Taylor, 1969; King and Jukes, 1969; O’Donald, 1969; Corbin and Uzzell, 1970), it is perhaps best not to exclude those back mutations for which parallel fixations represent an equally parsimonious solution. Actually, the procedure in this paper produces weights suggesting that back mutations are less likely on statistical grounds since the probability of the C→A replacement in the descent to the penultimate ancestor in Figure 2f is only 0.25.

Finally, it should be pointed out that the procedure presented here has not been mathematically proved. Counter examples have been diligently sought but all attempts to find a more parsimonious solution to the one(s) obtained by this method or to find equally parsimonious solutions not obtained by this procedure have failed. Nevertheless, in the absence of rigorous proof of the method’s validity, a medico in of judicious reserve is not totally unwarranted. The procedure is presented in this state in order to invite mathematicians to bring their talents to bear upon this problem. One additional caveat is in order when translating amino acid sequences into codon sequences, namely, a selection must be made between the codons A.G.(CU) and U.C. (ACGU) for serine, A.G.(AG) and C.G. (ACGU) for arginine, and U.U.(AG) and C.U.(ACGU) for leucine. Any attempt at total ambiguity leads to such cases as (AU).(CC).(ACGU) for serine. Unfortunately, this implies the possibility that A.C.A. (threonine), U.G.U. (cysteine) or others are present when in fact only serine is present. This in turn could lead to others errors. A computer procedure is available for translating amino acids into codons that selects the codons for serine, arginine and leucine that are most likely to give the fewest mutations.

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REFERENCES


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