Science - 2 Assignment 4 Page: 11 Maimeesh Nanayern Troos 2020/0/074 Applications powerful tool for many fields such as phylogenetic reconstruction, illumination of functionality important regions, and prediction of higher order structures of proteins and RNAS eseful in bioinformatics for identifying sequence similarity a producing phylogenetic freez , & Landon trapely models but the sequence producing phylogenetic trees developing homology models of protein structures MSA carries more into than Pouruse alignment it is because they show conserved regions with a protein family which are of structural & functional importance. MSA is basically an extension of purvice alignment to incorporate more than two sequences at a time. Q2) sum-of-poirs score

Multiple sequence similarity suggests a common surderlying structure of the govtein, a common function, or a common evolutionary source.

For B measurement of the quality of alignment we analyze the scoring function which is used.

After getting aligned sequences, we sum up the score in all columns.

Som-of-pairs is one such swring system. It is defined on columns & is the sum of all parrishe scores of the symbols in the column.

DRAWBACKS LALTERNATIVE

becomes easily brased if the sequences from the same family are over-represented in the justification for the score.

-> Alternative & to use weighted sop.

(3) steps involved in Progressive alignment approach

Step 1 . parrise alignments - calculate distance Step 2!

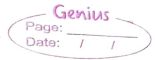
Comstruct a phylogenetic tree, cluster closely related sequences.

(rested neighbor joining tree (guide tree))

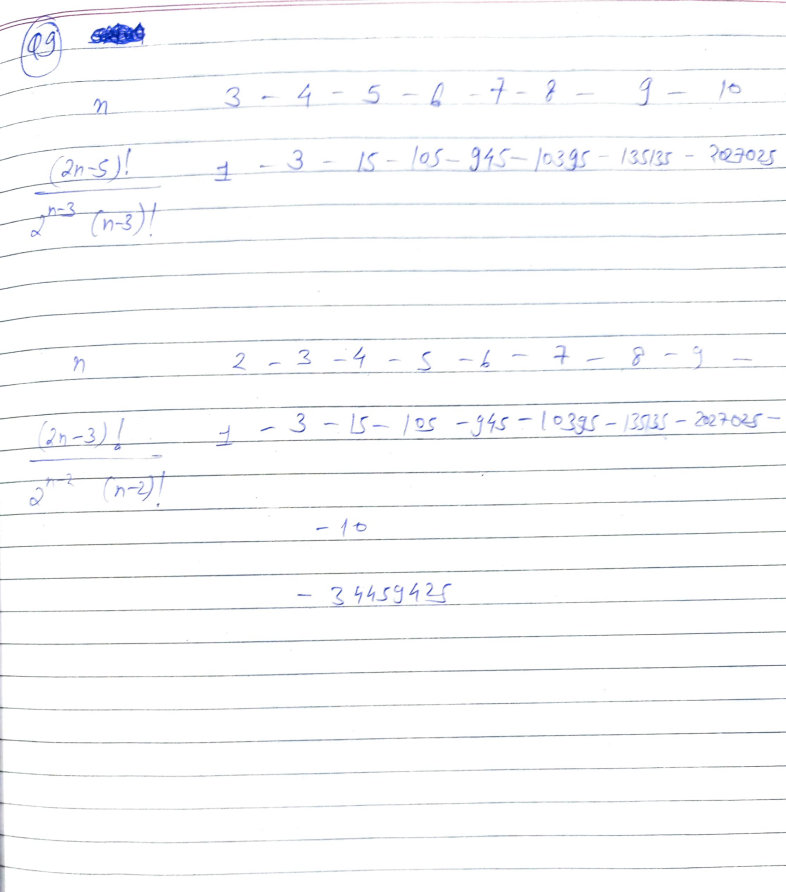
Progressive alignment: Align following the Step3:

closely related sequences first,

Dechy depends on the very first closely related sequences used for constructing the multiple alignment. DRAWBACKS More distantly related the sequences are more errors will get propogated through the alignment. De greedy approach results in officiency of the algorithm at the cost of accuracy. Sold Fixes to these Drawbacks for DI -> if the sequences align well there will be fewer errors De -> Using Bayesian methods such as Hidden Markov models (AMMS) may be useful. Other solutions: Stochastic or Herative methods



We have to align N sequences of length 50 g and knowing that a single pairwise companison fakes I second & the alignment of 4 of those segmences takes 109 seconds, I me have 5 billion years. 5 x 10 12 years = \$ 15768 x 10 seconds $(2L)^{N-2} = 10^{2N-4} = 15768 \times 10^{-16}$ QN-9) log (0 = log (15768) × 1016 N = log (1576x10(6) + 4 - [12.0988] = 12 seguinees 5 billion yrs, we can align 12 sequences.



a) [7,7,3] b) ST2, T3, T5} C) ET, 1763, ET2, T3, T55 d) { T, T2, T3, T4, T5, T6 } 74, 76

MSA of 6 species site: 3 Speciel asc Jast de 3 GCAT ATAG 4 GAAC