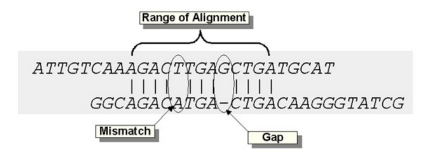
Alignment

<u>Similarity, Identity & Homology</u>: Sequence similarity can simply be a mathematical distance between two sequences given events such as insertions, deletions and substitutions. Sometimes the percent of matching nucleotides or amino acids is referred to as the percent similarity, sometimes, incorrectly called percent homology, and sometimes as referred to as identity. In some cases similarity is used to refer to shared properties (e.g. charge or hydrophobicity) even when nucleotides are different. Even NCBI is inconsistent in the use of these terms and you will need to consider the context when you see them. When identity or similarity refers to matching nucleotides or amino acids, and the amount is 25% or more, this suggests possible shared function.

Generally alignment is when two or more sequences (bases, amino acids, proteins, etc.) are matched in a pairwise, either globally (two sequences matched over their whole length) or locally (some subset of the sequences matched while other regions are not expected to match).

In the simplest model this is the "Edit distance" or the minimal number of events/edits required to transform one sequence into another.

Example: to go from acctga to agcta: accgta <<[substitution]>> agctga <<[deletion]>> agcta The edit distance = 2.



S= \sum (identities, mismatches) - \sum (gap penalties)

Score = Max(S)

Image: http://www.ncbi.nlm.nih.gov/books/NBK62051/

BLAST (Altschul et al. 1990). Basic local alignment search tool. For example, a gene is newly identified and function understood in *Drosophila*, a researcher can BLAST the database of the human genome to look for similar gene sequences.

Very basic description of BLAST

1. Uses short segments of sequence to find other sequences that contain the same set.

- 2. Does "ungapped" alignment extending from the matched subsequence regions to find high-scoring matches
- 3. Does a rapid gapped alignment to select and rank close matches

Alignment for phylogenetic analysis. Practical issues:

For phylogenetic analysis we need to establish character homology (aligned columns in the matrix) and primary homology hypotheses of character states.

Two problems- how to find alignments and how to choose.

For two sequences, i.e. pairwise alignment, of length **n**, if no gaps are allowed then there is one or few optimal alignments. If gaps are allowed, i.e. there is sequence length variation, then... (2n)!/(n!)2 e.g. **n**=50 then 1029 alignments. Enumeration is not an option! We need heuristic searches based on optimality and scoring. For phylogenies, pairwise comparison is not sufficient. What must be done is multiple sequence alignment, a global solution for the whole data matrix or primary homology for the characters (columns) in the matrix.

```
Taxon1 CTTCCGAATTTGGCT
Taxon2 ACTCGATTGCCT

Minimize substitutions
ACTTCCGAATTTGG-CT
||| ||| ||| ||
ACT--CGA--TTG-CCT

Minimize ind/dels
ACTTCCGGAATTTGGCT
|||* **|||*||
ACTC----GATTGCCT
```

Alignment really attempts to balance the amount of in/dels with the amount of base substitution, normally based on some cost differential. Of course it is possible to account for all differences by inserting enough gaps (trivial alignment).

Gaps are needed, but don't really existpoint mutation
unequal crossing over during meiosis
DNA slippage during replication
retrovirus insertions
movement of sequences in the genome (e.g. translocation or transposition)

Very brief overview of some methods

Dynamic Programming and global alignment: (Needleman-Wunsch) underlies or is part of most alignment methods. See tutorial at http://www.sbc.su.se/~pjk/molbioinfo2001/dynprog/dynamic.html

Simultaneous alignment- Simultaneous multiple alignments synchronise the information of all input sequences in a hyperspace lattice, e.g. so-called exact alignment algorithms using the divide-and-conquer

(DCA) strategy (Tönges et al. 1996). In part it cuts down the input sequences at carefully chosen positions to align in segments. Current algorithms cannot handle large/complex data sets.

Progressive alignment- As in Clustal W(X) the most prominent program for progressive alignment strategies.

- 1. All sequences are compared to each other (pairwise alignments)
- 2. A dendrogram is constructed, describing the approximate groupings of the sequences by similarity.
- 3. Final multiple alignment uses the guide tree

Basically, the multiple alignment is created by iteratively aligning sequences from the input to an already partially constructed solution. Obviously, the order is a crucial point in this method as uses a sort of UPGMA tree-based alignment order and requires sequence weighting.

It could be argued that it doesn't make sense to determine alignment order with one optimality criterion (e.g., phenetics) and then analyze the alignment later with another (e.g., parsimony, ml) but to re-align on a parsimony tree derived from the first alignment to get an "improved" alignment may be circular.

Progressive, consistency-based alignment- These strategies are incorporating "local signals" into global alignment construction.

Iterative, segment-based alignment- One example is DIALIGN, which iteratively collects local similar segments, which can be merged into a common multiple alignment. Iteration continues until no more local signals can be found or until all positions are aligned. Recent benchmarks have shown that this strategy can even handle long sequences of a low overall similarity. No explicit gap cost or input trees.

Direct optimization - POY (Varón et el. 2010)-The correspondences among homologues are determined and evaluated simultaneously with transformations. Treespace is explored without a static alignment.

- -Single process of alignment and tree construction.
- -Insertion and deletion events are counted are real events (transformations) as opposed to being implied by the pattern as in multiple sequence alignments.

Eliminates inconsistent treatment of data between alignment and tree construction steps. Tree-alignment methods like this are ones that simultaneously deals with base changes and insertion/deletion events on the tree, with simultaneous estimation of the parameters of change (including rates in insertions and deletions, which is what in the parsimony world is referred to as gap costs, or rate matrices for nucleotide change (e.g., TV/TS ratios)).

Direct optimization and iterative-pass optimization strive to construct HTU sequences such that the overall cladogram is of minimal length. This is done through modified two and three dimensional string-matching, respectively.

Fixed-states optimization and search-based optimization draw optimal HTU sequences from a pool of predetermined sequences. This can be a small or large collection of possible sequences. Dynamic programming is used to identify the best HTU sequences and determine cladogram length.

Adjustments:

Manual or by eye- For very simple data this may be sufficient, however, it violates any criterion of repeatability as there is no obvious costs matrix. The counter argument is that the aligned matrix can be made available.

Purging "bad" data or scoring variable regions as single characters. Frequently used get around problems in hard to align sections is the elimination of gap heavy regions in alignments. Exactly which columns should be eliminated (left-right boundaries) is subjective and obviously they may have an impact on the results (otherwise why bother).

Alternatively, the variable region can be converted into a character in each taxon and scored. This has all the problems above and adds another layer of difficulty in determining how to code the states.

Nucleotide alignments of coding sequences informed by amino acids. This works well and can easily be implemented in commonly used software such as Mesquite.

Alignments informed by consideration of secondary structure-

- 1. Does not solve the problem of nucleotide homology.
- 2. Determination of secondary structure is not simple and not unambiguous. Generally the actual pattern of bonding is probabilistic and depends on the minimization of free energy and the thermodynamic stability of the resulting structure. Programs explicitly designed to model secondary structure are not very realistic (yet) in terms of the actual cell environment and might find multiple, equally probable models. In phylogenetic studies, secondary structure is typically inferred by aligning with a sequence of "known" secondary structure, although the basis of that knowledge remains uncertain and applicability to the study taxa is unclear in many cases, but this is heading in the right direction.
- 3. There might be reasonable to expect selective pressures to apply to secondary structure interactions (that is, requirements of compensatory changes), it is unclear just how relevant those interactions are compared to selective pressures applied at other structural levels.

Detail your methods!

However you derive your alignment it is necessary to explain clearly how it was done. Cite the software, its version number and setting. If you make manual edits the aligned matrix should be deposited.

Reference cited:

Altschul, S.F., W. Gish, W. Miller, E.W. Myers, and D.J. Lipman. 1990. Basic local alignment search tool. J Mol Biol 215(3):403-10.

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Varón, A., L. S. Vinh, W. C. Wheeler. 2010. POY version 4: phylogenetic analysis using dynamic homologies. Cladistics, 26:72-85.