



Table containing all sequences (species) as rows and homologous positions as columns

Main Criteria for building a multiple sequence alignment			
Criterion	Meaning		
Structure similarity	Amino acids that play the same role in each structure are in the same column. Structure superposition programs are the only ones that use this criterion.		
Evolutionary similarity	Amino acids or nucleotides related to the same amino acid (or nucleotide) in the common ancestor of all the sequences are put in the same column. No automatic program explicitly uses this criterion, but they all try to deliver an alignment that respects it.		
Functional similarity	Amino acids or nucleotides with the same function are in the same column. No automatic program explicitly uses this criterion, but if the information is available, you can force some programs to respect it or you can edit your alignment manually.		
Sequence similarity	Amino acids in the same column are those that yield an alignment with maximum similarity. Most programs use sequence similarity because it is the easiest criterion. When the sequences are closely related, structure, evolutionary and functional similarities are equivalent to sequence similarity.		

Main applications of multiple sequence alignments			
Application	Procedure		
Extrapolation	A good multiple alignment can help convincing you that an uncharacterized sequence is really a member of a protein family.		
Phylogenetic analysis	If you carefully chose the sequences to include in your multiple alignment, you can reconstruct the history of these proteins.		
Pattern Identification	By discovering very conserved positions you can identify a region that is characteristic of a function (in proteins or in nucleic acid sequences).		
Domain identification	It is possible to turn a multiple sequence alignment into a profile that describes a protein family or a protein domain. You can use this profile to scan databases for new members of the family.		
DNA regulatory elements	You can turn a DNA multiple alignment of a binding site into a weight matrix and scan other DNA sequences for potential similar binding sites.		
Structure prediction	A good multiple alignment can give you an almost perfect prediction of your protein secondary structur for proteins or RNA. Sometimes it can also help building a 3-D model.		
PCR analysis	A good multiple alignment can help you identifying the less degenerated portions of a protein family		
nsSNP	Identify the nsSNP that are the most likely to alter the function		

- Move from gene homology to homology at the nucleotides and residues:
 - Input: a set of (homologous) sequences
 - Output: an alignment of every pair of sequences that is consistent $(x_i \leftrightarrow y_j \text{ and } x_i \leftrightarrow z_k \text{ implies } z_k \leftrightarrow y_j)$

Where's the problem?

```
TACAT
TCCAGT
TCAGT
```

pairwise alignments (Needleman-Wunsch)

TACA-T TACA-T TCCAGT TCCAGT T-CAGT

pile them up: they are consistent!

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TAGT

TACA-T

TACAT

TCCAGT

TCCAGT

TAG-T

T--AGT

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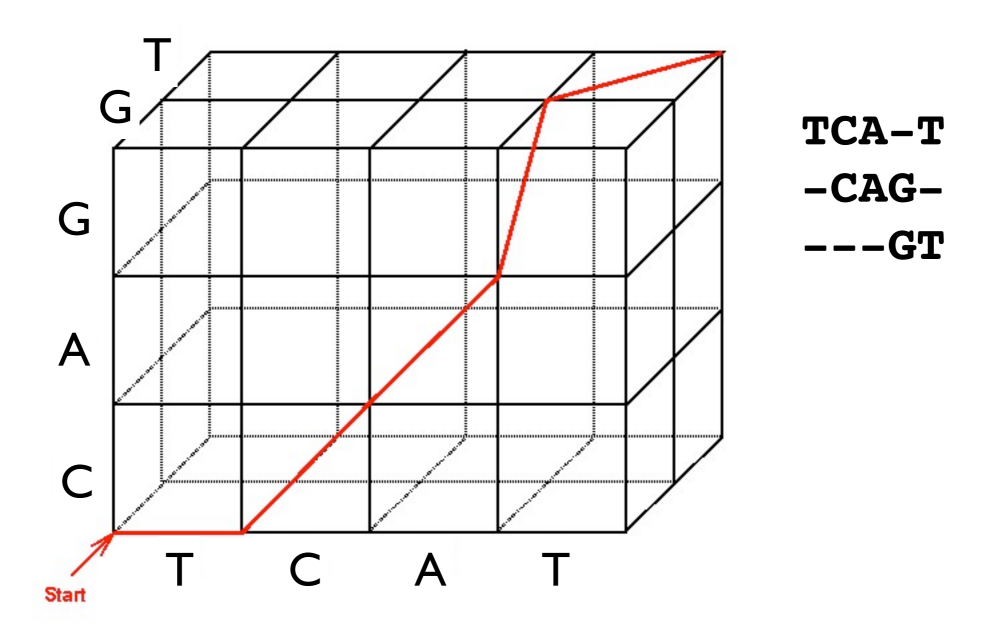
TACA-T
TCCAGT
T-CAGT

TACAT
TCCAGT
TAGT

TACA-T TACAT TCCAGT
TCCAGT TAG-T T--AGT

TACA-T TACA-T
TCCAGT Or TCCAGT ...
TAG--T TCCAGT ...

Multi-dimensional dynamic programming



Dynamic programming: extend Needleman-Wunsch to $n_1 \times n_2 \times n_3$ table

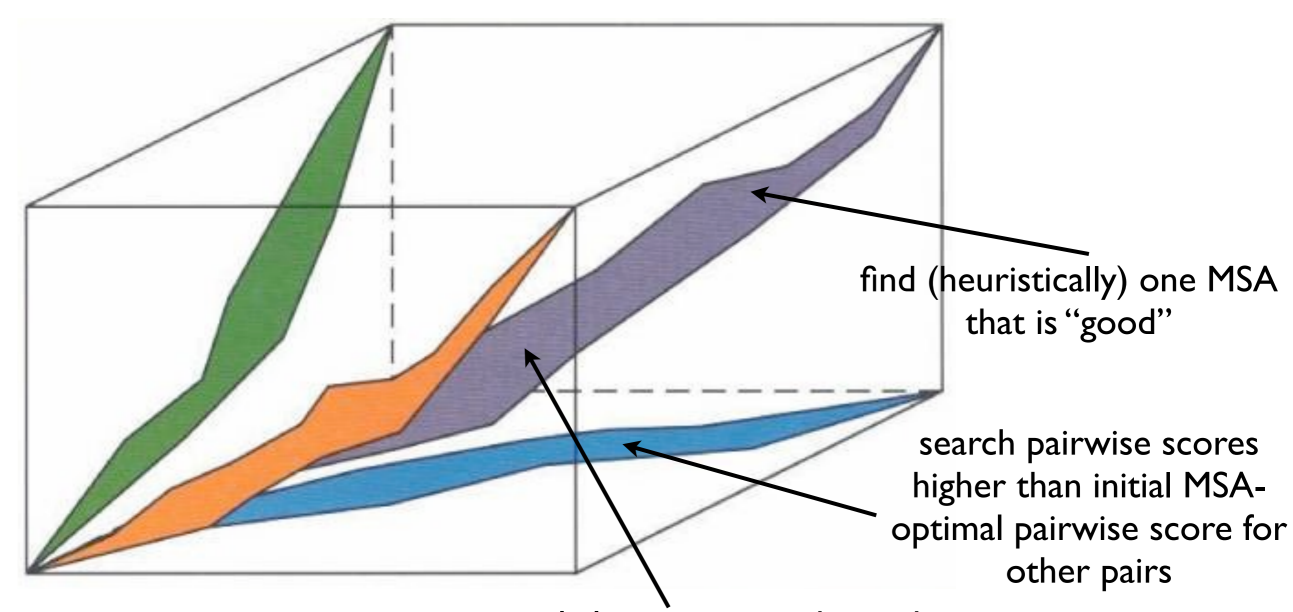
We want to optimize the MSA, but what is the score? <u>Version I: Sum of pairs (SP) score</u>

$$S(MSA) = \sum_{i} S(col_i)$$
,

score each column independently, then sum them all

$$S(\operatorname{col}_i) = \sum_{j < k} M(x_{ji}, x_{ki})$$
 . ith column in alignment All pairs of sequences

Multi-dimensional dynamic programming



optimal alignment must be in the intersection of the pairwise bounds

Restrict table to a narrow "tube" around initial find

Version I: Sum of pairs (SP) score

Problem with the probabilistic interpretation of scores:

$$M(x_{ji}, x_{ki}) + M(x_{ji}, x_{\ell i}) + M(x_{ki}, x_{\ell i})$$

$$= \log p(x_{ji}x_{ki}) + \log p(x_{ji}x_{\ell i}) + \log p(x_{ki}x_{\ell i})$$

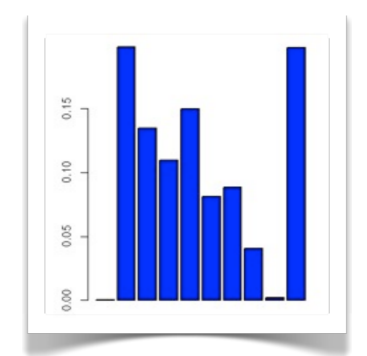
$$-2\log q(x_{ji}) - 2\log q(x_{ki}) - 2\log q(x_{\ell i})$$

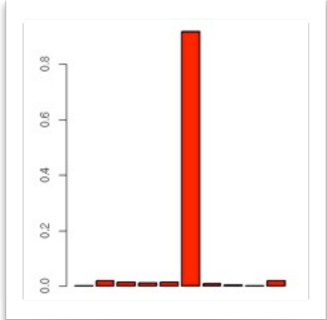
$$\neq \log \frac{p(x_{ji}x_{ki}x_{\ell i})}{q(x_{ji})q(x_{ki})q(x_{\ell i})},$$

Common improvement uses weighted means of pairs scores

Version 2: Entropy

- Entropy H(p) is a measure of how flat (or peaked) is a probability distribution
- peaked = $0 \le H \le \log 2(N)$ = flat





$$H(p) = -\sum_{k=1}^{N} p(k) \log_2(p(k))$$

Version 2: Entropy

- We can score a MSA by the sum of the column entropies
- Best alignment has minimum entropy

$$p_i(n) = \frac{\#\{x_{ji} = n\}}{\#\{x_{ji} \neq -\}},$$

 $n = A, C, G, T.$

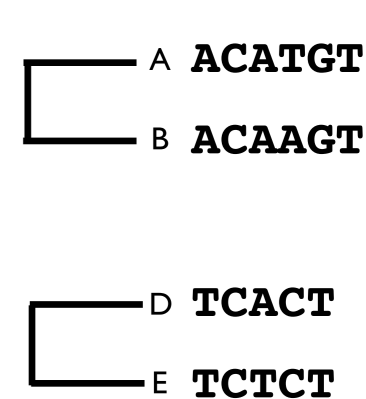
- Strategy: compute all pairwise optimal alignment scores
- Build a guide tree by "Neighbour-Joining" (NJ):
 - join the two nearest items in a tree node
 - replace the pair by one item in the list, its pairwise scores are the maximum of the two scores
 - iterate these two operations
- Construct the MSA in the order of the tree
 - when aligning a sequence to a previous MSA, do what is called a profile alignment

Pairwise alignment scores

	В	C	D	Ш
A	9	-	2	2
	В	4	2	0
		С	3	0
			D	7
				E

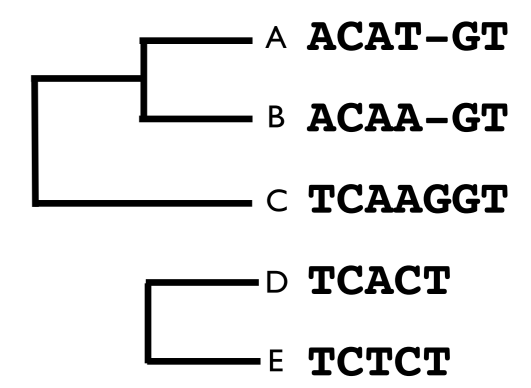
Pairwise alignment scores

	В	C	D	Е
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	В	4	2	0
		C	3	0
			D	7
				E



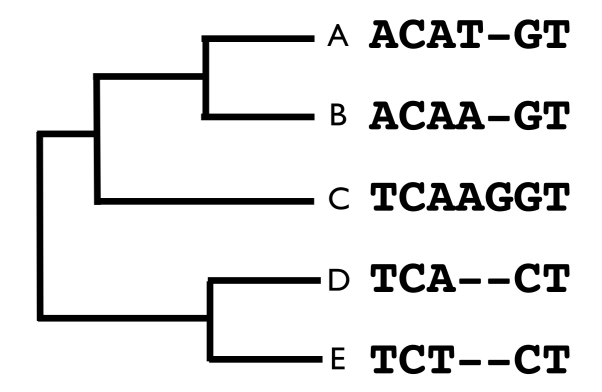
Pairwise alignment scores

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Pairwise alignment scores

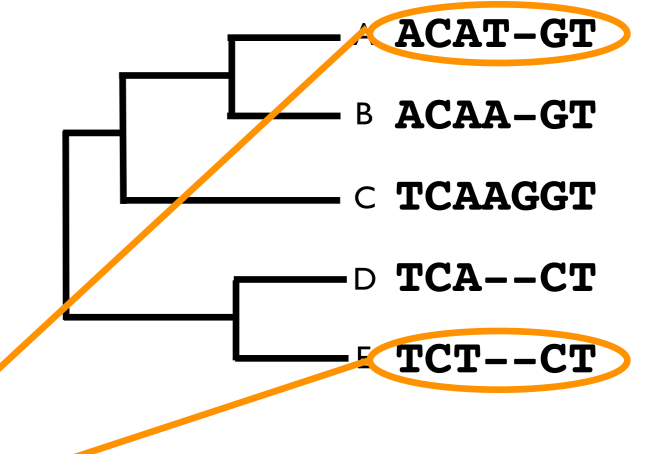
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Pairwise alignment scores

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			D	7
				Ε

ACATGT
ACAAGT
TCAAGGT
TCACT
TCTCT



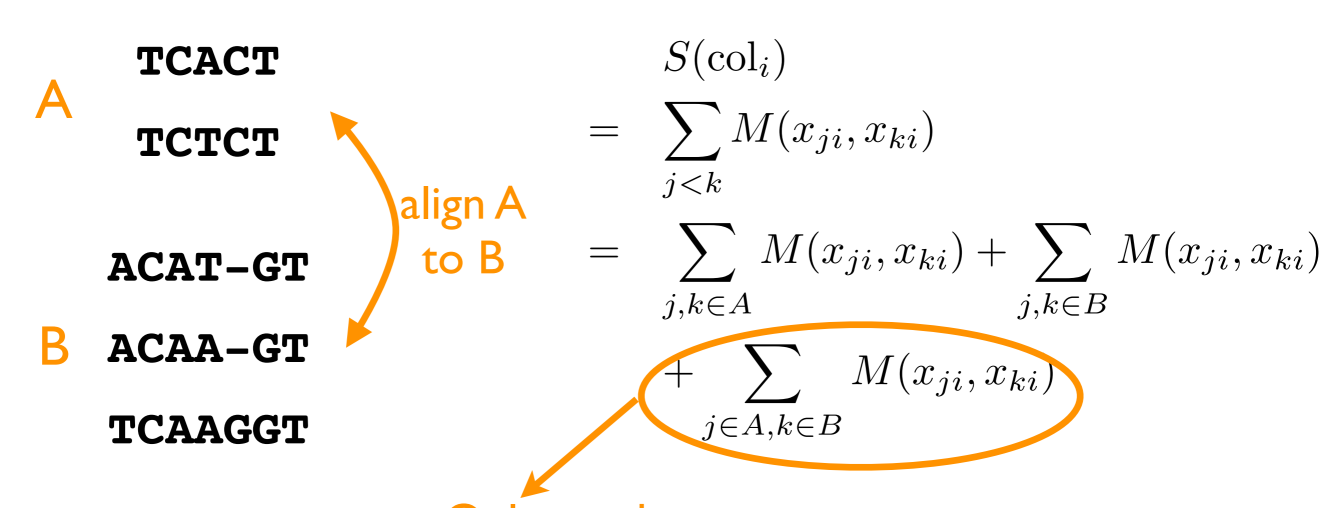
Best pairwise alignment

ACAT-GT

TC-T-CT

Alignment to a profile

A MSA can be seen as a "profile": a nucleotide distribution at each position

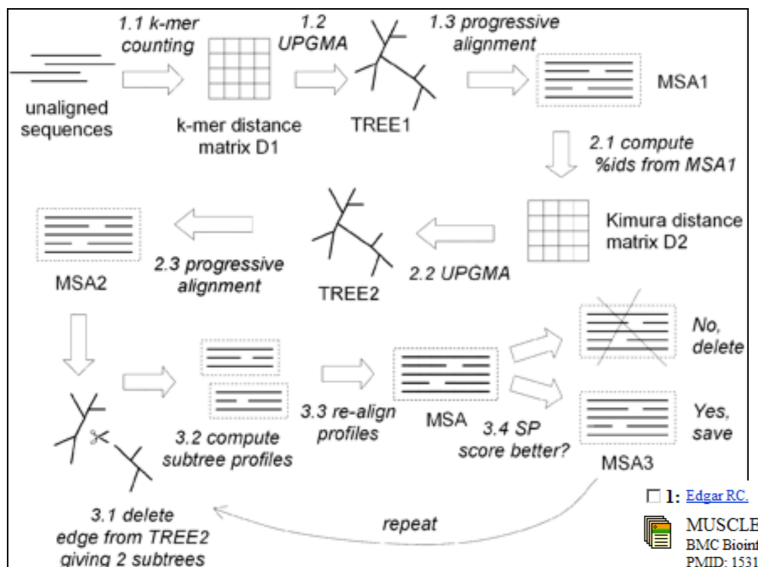


Only need to optimize this sum

Requires M(-,-) = 0

Consistency post-processing

Because of the above scoring scheme, earlier alignments can never be modified by later sequence additions



MUSCLE: a multiple sequence alignment method with reduced time and space complexity. BMC Bioinformatics. 2004 Aug 19;5(1):113. PMID: 15318951 [PubMed - indexed for MEDLINE]

Purpose \Leftrightarrow Solution

	MUSCLE	MAFFT	PROBCONS	T-COFFEE	CLUSTALW
Dist Based Phylogeny	+++	+++	++	++	++
ML or MP Phylogeny	++	+++	+++	+++	++
Profile Construction	++	+++	+++	+++	++
3D Modeling	++	++	++	+++	+
Secondary Structure P	+++	+++	++	++	++