International Bioinformatics Workshop on Molecular Epidemiology and Phylodynamics

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Multiple Sequence Alignment

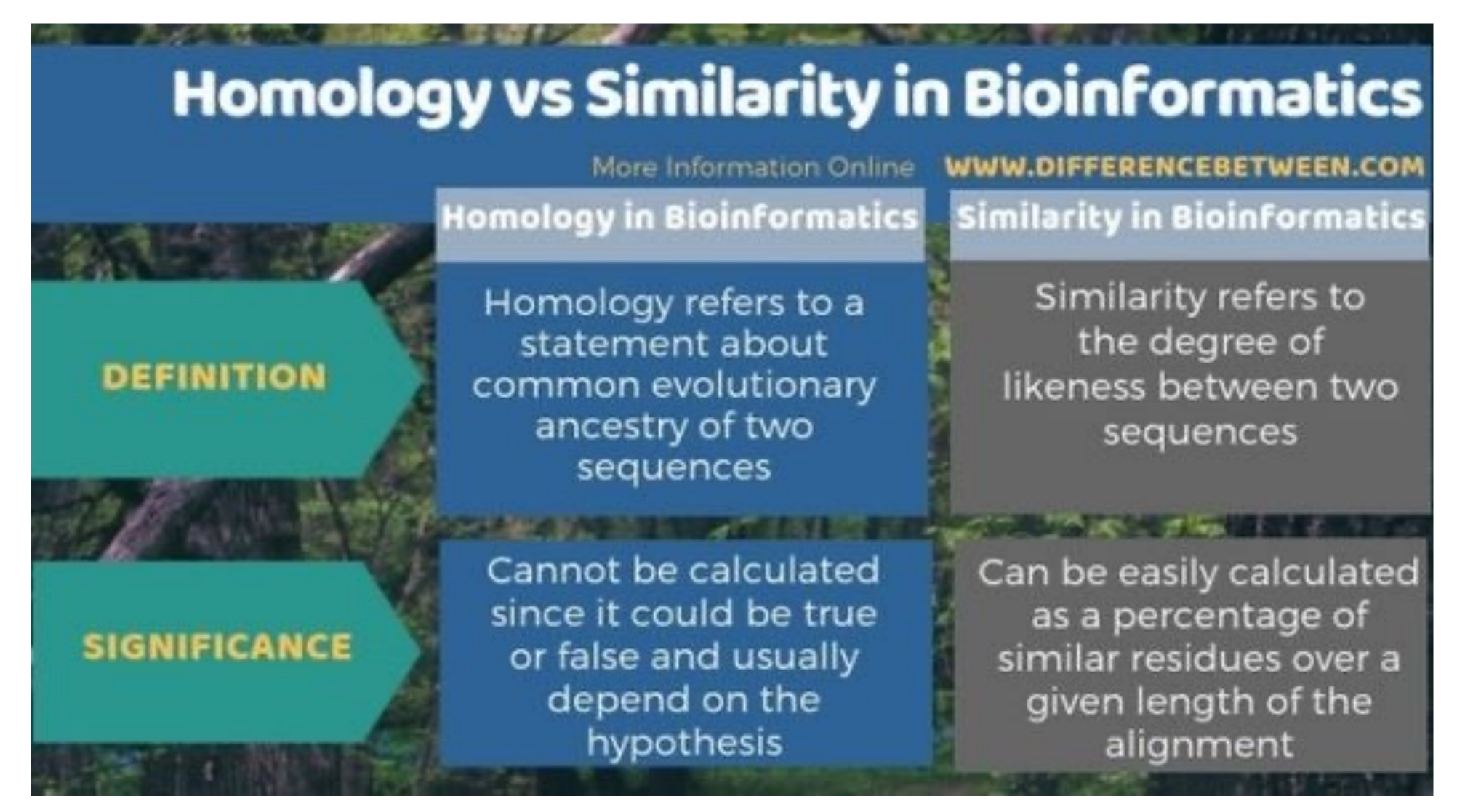
Learning Objectives

- Understand the algorithms behind multiple sequence alignment.
- Perform multiple sequence alignments using tools
 - MAFFT
 - EMBOSS

Outline

- 1. Homology v.s. Similarity
- 2. Global Alignment v.s. Local Alignment
- 3. Pairwise Alignment v.s. Multiple Sequence Alignment
- 4. Alignment methods
 - Longest common subsequence
 - Time Complexity
- 5. Sum of pairs for evaluating an alignment
- 6. Hands-on session

Homology v.s. Similarity



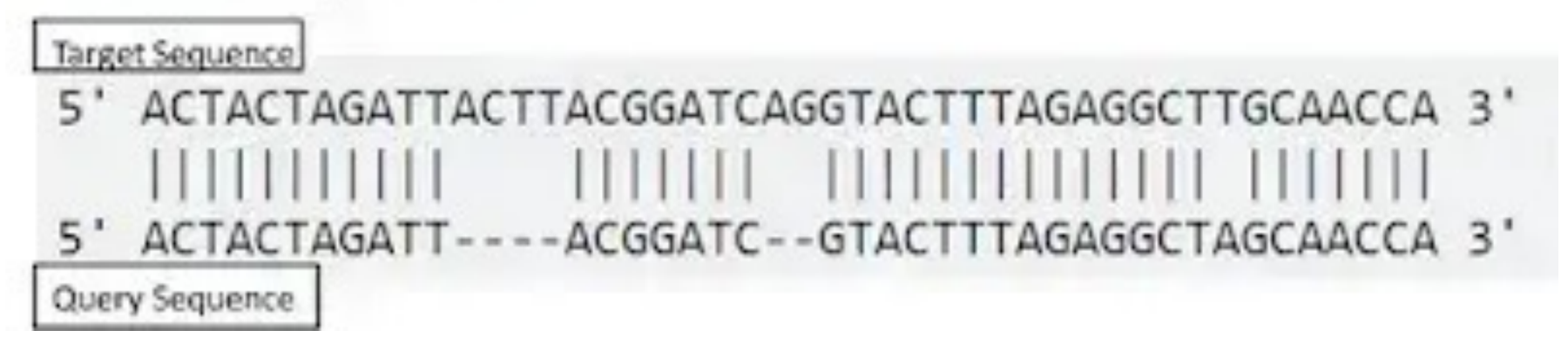
Global Alignment v.s. local Alignment

Global Alignment	Local Alignmennt
Purpose: Align the entire sequences	Find local regions with high similarity
Contains all nucletoides (or amino acids)	Align a substring of query to a substring of target
If two sequences are quite similar	Find stretches of sequences without considering the rest of sequence
Suitalbe for aligning two closely related sequences	Suitable for aligning more divergent sequences
Needleman-Wunsch algorithm (EMBOSS neelde)	Smith-Waterman algorithm (EMBOSS water)

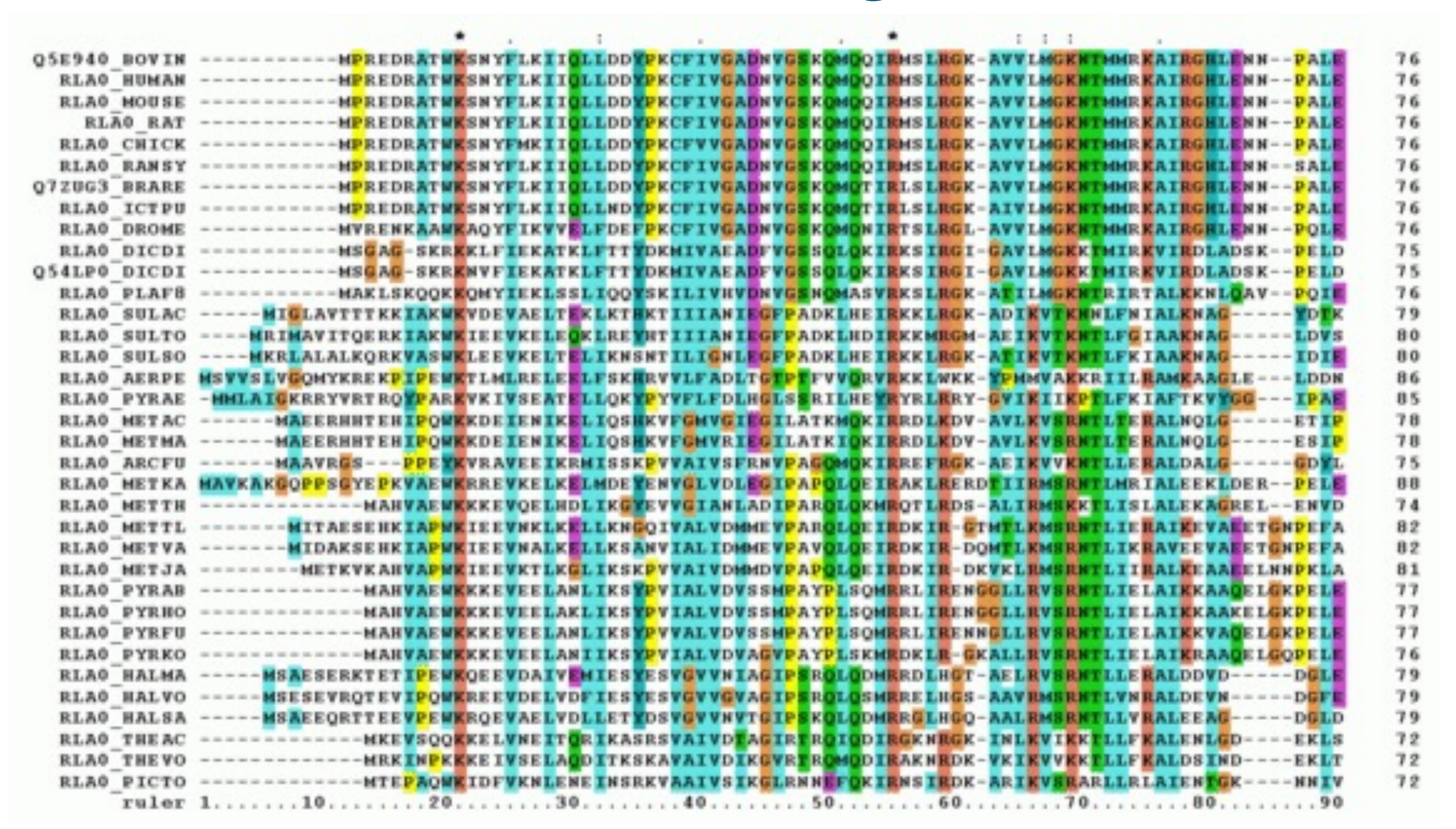
Local Alignment



Global Alignment



Multiple Sequence Alignment

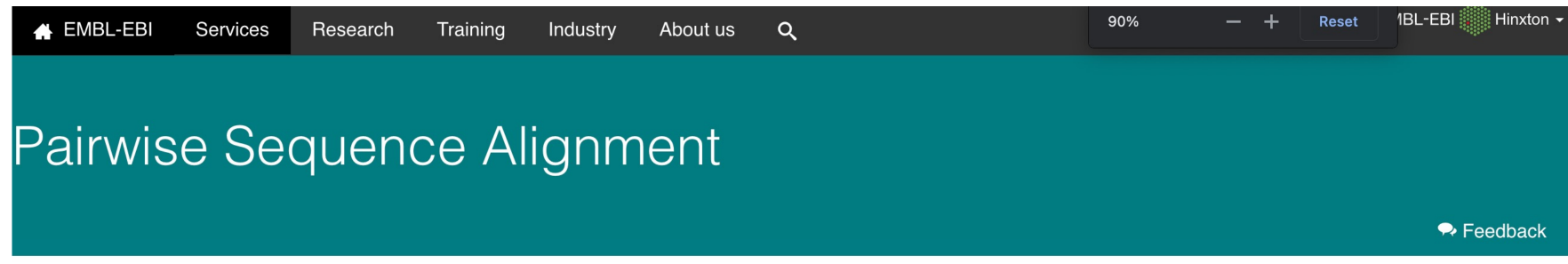


Pairwise Alignment v.s. Multiple Sequence Alignment

Pairwise Alignment	Multiple Sequence Alignmennt
two sequences	Three or more
Global or local alignment	Global
Comparatively simple algorithm	Complex sophisticated algorithm
Needleman-Wunsch algorithm (EMBOSS neelde), Smith-Waterman algorithm (EMBOSS water), and so on	Progressive alignment construction, Iterative methods, and so on
Applications: (1) find out conserved regions between two sequences; (2) similarity searches in a database	Applications: (1) find out regions of variability or conservation in a family; (2) similarity searches between a newly sequence and an existing gene family; (3) phylogenetic analysis

Online tools

https://www.ebi.ac.uk/Tools/psa/; https://www.ebi.ac.uk/Tools/msa/



Tools > Pairwise Sequence Alignment

Service Announcement

The new Job Dispatcher Services beta website is now available at https://wwwdev.ebi.ac.uk/Tools/jdispatcher. We'd love to hear your feedback about the new webpages!

Pairwise Sequence Alignment is used to identify regions of similarity that may indicate functional, structural and/or evolutionary relationships between two biological sequences (protein or nucleic acid).

By contrast, Multiple Sequence Alignment (MSA) is the alignment of three or more biological sequences of similar length. From the output of MSA applications, homology can be inferred and the evolutionary relationship between the sequences studied.

Global Alignment

Global alignment tools create an end-to-end alignment of the sequences to be aligned.

Needle (EMBOSS)

EMBOSS Needle creates an optimal global alignment of two sequences using the Needleman-Wunsch algorithm.

Launch Needle

Stretcher (EMBOSS)

EMBOSS Stretcher uses a modification of the Needleman-Wunsch algorithm that allows larger sequences to be globally aligned.

Launch &Stretcher

GGSEARCH2SEQ

GGSEARCH2SEQ finds an optimal global alignment using the Needleman-Wunsch algorithm.

Launch *ggsearch2seq

Local Alignment

Local alignment tools find one, or more, alignments describing the most similar region(s) within the sequences to be aligned. They are can align protein and nucleotide sequences.

Water (EMBOSS)

EMBOSS Water uses the Smith-Waterman algorithm (modified for speed enhancements) to calculate the local alignment of two sequences.

Matcher (EMBOSS)

EMBOSS Matcher identifies local similarities between two sequences using a rigorous algorithm based on the LALIGN application.

Launch Matcher

LALIGN

LALIGN finds internal duplications by calculating non-intersecting local alignments of protein or DNA sequences.

Launch **\LALIGN**

SSEARCH2SEQ

SSEARCH2SEQ finds an optimal local alignment using the Smith-Waterman algorithm.

Launch Assearch2seq

Genomic Alignment

Genomic alignment tools concentrate on DNA (or to DNA) alignments while accounting for characteristics present in genomic data.

GeneWise

GeneWise compares a protein sequence to a genomic DNA sequence, allowing for introns and frameshifting errors.

¾Launch GeneWise

The tools described on this page are provided using Search and sequence analysis tools services from EMBL-EBI in 2022

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Multiple Sequence Alignment

Feedback

Tools > Multiple Sequence Alignment

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Multiple Sequence Alignment (MSA) is generally the alignment of three or more biological sequences (protein or nucleic acid) of similar length. From the output, homology can be inferred and the evolutionary relationships between the sequences studied.

By contrast, Pairwise Sequence Alignment tools are used to identify regions of similarity that may indicate functional, structural and/or evolutionary relationships between two biological sequences.

Clustal Omega

New MSA tool that uses seeded guide trees and HMM profile-profile techniques to generate alignments. Suitable for medium-large alignments.

Launch Clustal Omega

Cons (EMBOSS)

EMBOSS Cons creates a consensus sequence from a protein or nucleotide multiple alignment.

♣Launch EMBOSS Cons

Clustal Omega

New MSA tool that uses seeded guide trees and HMM profile-profile techniques to generate alignments. Suitable for medium-large alignments.

▲Launch Clustal Omega

Cons (EMBOSS)

EMBOSS Cons creates a consensus sequence from a protein or nucleotide multiple alignment.

♣Launch EMBOSS Cons

Kalign

Very fast MSA tool that concentrates on local regions. Suitable for large alignments.

▲Launch Kalign

MAFFT

MSA tool that uses Fast Fourier Transforms. Suitable for medium-large alignments.

♣Launch MAFFT

MUSCLE

Accurate MSA tool, especially good with proteins. Suitable for medium alignments.

♣Launch MUSCLE

MView

Transform a Sequence Similarity Search result into a Multiple Sequence Alignment or reformat a Multiple Sequence Alignment using the MView program.

▲Launch MView

T-Coffee

Consistency-based MSA tool that attempts to mitigate the pitfalls of progressive alignment methods. Suitable for small alignments.

▲Launch T-Coffee

WebPRANK

The EBI has a new phylogeny-aware multiple sequence alignment program which makes use of evolutionary information to help place insertions and deletions.

Try it out at WebPRANK.

Alignment methods

Longest Common Subsequence

To find a longest common subsequence between two strings.

S1: TAGTCACG

S2: AGACTGTC

LCS: AGACG

a computer programming technique where an algorithmic problem is first broken down into sub-problems, the results are saved, and then the sub-problems are optimized to find the overall solution — which usually has to do with finding the maximum and minimum range of the algorithmic query.

https://www.spiceworks.com/tech/devops/articles/what-is-dynamic-programming/

Equation

$$c_{i,j} = \max \begin{cases} c_{i-1,j-1} + 1 & \text{if } a_i = b_j \\ c_{i-1,j} + 0 & \text{if } a_i \neq b_j \\ c_{i,j-1} + 0 & \text{if } a_i \neq b_j \end{cases}$$

Go to this website and input sequences

```
ⓒ Fork this ► Run
                                                              Language C++ V 1 1
 1 #include <iostream>
 2 #include <iomanip>
 3 using namespace std;
 5 int **table;
 6 string s1,s2;
 8 void find_LCS()
        int i,j,n,m;
        char x;
        n = s1.length();
        m = s2.length();
       // 宣告 2D-array by size of 2 strings
        table = new int*[n+1];
        for (int i=0; i \le n; i++) table[i] = new int[m+1];
       // 將第一排第一列都先歸零
        for (int i=0;i<=n;i++) table[i][0] = 0;
        cout<<"str2: ";</pre>
        for (int j=0; j<=m; j++)</pre>
21-
            table[0][j] = 0;
                                                                    //****顯示比較字
            if (j==0) cout<<"___"; else cout<<setw(3)<<s2[j-1];</pre>
        cout<<endl<<"str1 |"; for (int j=0;j<=m;j++) cout<<" 0"; //****顯示比較字
                                                                    //****顯示比較多
        cout<<endl;</pre>
  G 0 ^1 +2 <2 <2 <2 +2 <2 <2
      ^1 +2 ^3 ^4 ^4 +5 <5 <5
```

https://www.onlinegdb.com/Hkx1yiw2v

$$c_{i,j} = \max \begin{cases} c_{i-1,j-1} + 1 & \text{if } a_i = b_j \\ c_{i-1,j} + 0 & \text{if } a_i \neq b_j \\ c_{i,j-1} + 0 & \text{if } a_i \neq b_j \end{cases}$$

Step 1

S2 G

G 0 0

Ğ	U				
Т	0				
C	0				
A	0				
C	0				

Step 2

	S2	Α	G	A	C	Т	G	Т	C
S1	0	0	0	0	0	0	0	0	0
Т	0	0	0	0	0	1	1	1	1
A	0								
G	0								
Т	0								
C	0								
A	0								
C	0								
G	0								

$$c_{i,j} = \max \begin{cases} c_{i-1,j-1} + 1 & \text{if } a_i = b_j \\ c_{i-1,j} + 0 & \text{if } a_i \neq b_j \\ c_{i,j-1} + 0 & \text{if } a_i \neq b_j \end{cases}$$

Step 3

S2 G G **S1** A G

Step 4

	S2	A	G	A	C	Т	G	Т	C
S1	0	0	0	0	0	0	0	0	0
Т	0	0	0	0	0	1	1	1	1
A	0	1	1	1	1	1	1	1	1
G	0	1	2	2	2	2	2	2	2
Т	0								
C	0								
A	0								
C	0								
G	0								

$$c_{i,j} = \max \begin{cases} c_{i-1,j-1} + 1 & \text{if } a_i = b_j \\ c_{i-1,j} + 0 & \text{if } a_i \neq b_j \\ c_{i,j-1} + 0 & \text{if } a_i \neq b_j \end{cases}$$

Step 5

	S2	A	G	A	C	Т	G	Т	C
S1	0	0	0	0	0	0	0	0	0
Т	0	0	0	0	0	1	1	1	1
Α	0	1	1	1	1	1	1	1	1
G	0	1	2	2	2	2	2	2	2
Т	0	1	2	2	2	3	3	3	3
C	0								
Α	0								
C	0								
G	0								

...

$$c_{i,j} = \max \begin{cases} c_{i-1,j-1} + 1 & \text{if } a_i = b_j \\ c_{i-1,j} + 0 & \text{if } a_i \neq b_j \\ c_{i,j-1} + 0 & \text{if } a_i \neq b_j \end{cases}$$

Final

	S2	Α	G	Α	С	Т	G	Т	C
S1	0	0	0	0	0	0	0	0	0
Т	0	0	0	0	0	1	1	1	1
A	0	1	1	1	1	1	1	1	1
G	0	1	2	2	2	2	2	2	2
Т	0	1	2	2	2	3	3	3	3
C	0	1	2	2	3	3	3	3	4
A	0	1	2	3	3	3	3	3	4
C	0	1	2	3	4	4	4	4	4
G	0	1	2	3	4	4	5	5	5

S1:TAGTCACG

S2: AGACTGTC

LCS: AGACG

Alignment

TAGTCAC-G--

-AG--ACTGTC

Time Complexity

The number of sequences is k

K-dimensional table

Each dimension is sequence length

Each entry depends on 2^k-1 adjacent entries

Complexity: O(2^kn^k)

NP (nondeterministic polynomial time)-complete problem

$$c_{i,j} = \max \begin{cases} c_{i-1,j-1} + 1 & \text{if } a_i = b_j \\ c_{i-1,j} + 0 & \text{if } a_i \neq b_j \\ c_{i,j-1} + 0 & \text{if } a_i \neq b_j \end{cases}$$

Sum-of-pairs for evaluating an alignment

Alignment #1

$$s_1 = ATTCGAT$$
 $s_2 = -TT-GAG$
 $s_3 = AT--GCT$

$$\sum_{i < j} scoring(S_i, S_j)$$

- $s(S_i \text{ and } S_i) = 1 \text{ when match}$
- $s(S_i \text{ and } S_i) = -1$ when mismatch
- $s(S_i \text{ and } S_i) = -2 \text{ when gap}$

For the alignment, the pairwise alignment score of position 7:

$$score(s_1, s_2) = -1$$

 $score(s_2, s_3) = -1$
 $score(s_1, s_3) = 1 \Rightarrow SP \ score = -1$

Alignment #2

$$s_1 = ATTCGAT s_2 = -TT-GA-G$$
 $s_3 = AT--GCT-$

$$\sum_{i < j} scoring(S_i, S_j)$$

- $s(S_i \text{ and } S_i) = 1 \text{ when match}$
- $s(S_i \text{ and } S_i) = -1$ when mismatch
- $s(S_i \text{ and } S_i) = -2 \text{ when gap}$

For the alignment, the pairwise alignment score of position 7 and 8:

$$score(s_1, s_2) = -2-2$$

$$score(s_2, s_3) = -2+1$$

$$score(s_1, s_3) = 1+1 \Rightarrow SP score = -3$$

Hands-on session

>1 ACTACTAGATTACTTACGGATCAGGTACTTTAGAGGCTTGCAACCA

>2 TACTCACGGATGAGGTACTTTAGAGGC

- 1. Copy and save as two sequence files (1.fas and 2.fas) (using Sublime app)
- 2. Install EMBOSS by using conda (https://anaconda.org/bioconda/emboss)
- 3. Perfrom Needle and Water to compare global and local alignment
 - \$ water 1.fas 2.fas 1-2.water -auto
 - \$ needle 1.fas 2.fas 1-2.needle -auto