Sequence analysis

Sequence analysis

- Next generation sequencing methods have generated large amounts of sequencing data
- Bioinformatics uses information from protein and DNA/RNA sequences to derive information about living organisms
- Most bioinformatic approaches are based on sequence comparisons

Sequence comparison/alignment

You have two sequences THISSEQUENCE and THATSEQUENCE

```
THISSEQUENCE
|||||||||
THATSEQUENCE
```

Yet another sequence THATISASEQUENCE

```
or THIS SEQUENCE
|||| ||||||
THATISASEQUENCE
```

Which aligment is the best?- Aligment scoring

- Simplest way is % idendity
- Scoring matrices e.g. Blosum62:

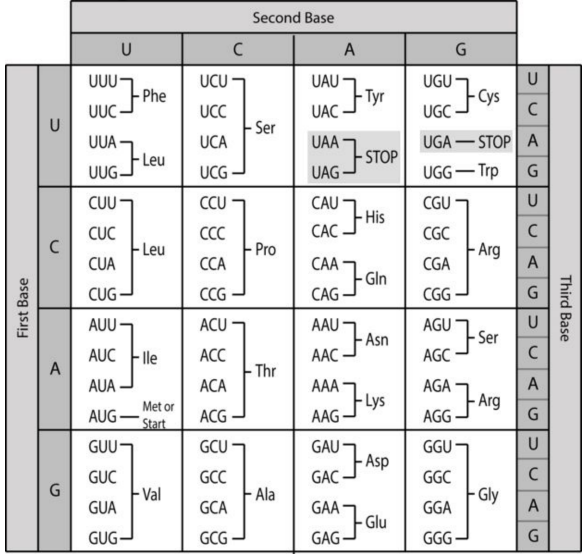
Sequence comparison/alignment

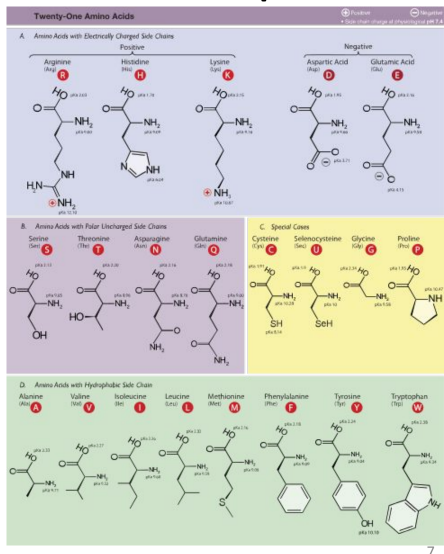
Scoring the first two sequences

```
T H I S S E Q U E N C E
T H A T S E Q U E N C E
5 8-1 1 4 5 5 0 5 6 9 5
Summ is 52
```

Substitution matrices

• "Various types of substitution matrices have been used over the years. Some were based on theoretical considerations, such as the number of mutations that are needed to convert one amino acid into another, or similarities in physicochemical properties. The most successful, however, use actual evidence of what has happened during evolution, and are based on analysis of alignments of numerous homologs of well-studied proteins from many different species." The genetic code – denoted as RNA triplets





Sequence homology – sequences derived from a common ancestor

 Sequences derived from a common ancestor during evolution share sequence similarities, how much is dependent on mutation rate and evolutionary distance

 Sequence homology can imply a common function or structure of a protein

 Sequence identity above 30% over a whole protein sequence is considered a good indication for homology

Blosum and PAM matrices

 PAM uses substitution frequencies from known closely related sequences - PAM matrix number indicates evolutionary distance thus increases with evolutionary distance and sequence divergence

• Blosum use mutation data from highly conserved local regions of sequence — Blosum matrix number refers to percentage identity thus decreases with evolutionary distance and sequence divergence

What about gaps?

- The likelyhood of insertion and deletion mutations is lower than that of point mutations and often result in frame shifts thus gaps receive a penalty the standard is -11
- Insertion and deletions can have different length thus and it is likely that not just a single amino acid got deleted gap extensions thus receive a lower penalty -1

Exercise: Sequence comparison/alignment

• Score the aligment of these two sequences

Local and global alignment

Global Alignment

Target seguence	1 ATCGTTGACGCACAAACACACTCTTCCAAGACCACCACATGCTGAGGTGT	50
Target sequence		36
Query sequence		41
	Local Alignment	
Target sequence	1 ATCGTTGACGCACAAACACACTCTTCCAAGACCACCACATGCTGAGGTGT	50
Query sequence		18

Basic local aligment search tool = Blast

- blastn: nucleotide blast
- tblastn: translated nucleotide blast
- blastp: protein blast
- Whatch NCBI tutorial:

https://digitalworldbiology.com/BLAST/slide1.html