

Bioinformatics Topics

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Generation

Experimental Data types include :

Sequences

- Typically [Next-Generation DNA Sequencing \(NGS\)](#).

3D Protein Structures - [X-ray crystallography](#) or

[Nuclear magnetic resonance spectroscopy \(NMR\)](#)

Gene Expression Data - [Microarrays](#)

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The Alignment of Pairs of Homologous DNA/Protein sequences.

Fundamental to most forms of DNA/Protein Sequence analysis

Substitution

Deletion

Conservation

Insertion

Putative Ancestor Sequence

... ACYG-QWRWDVI-S...
... ACFGHQWRWDVI-S...
... ACFGHQWRWDVILS...

Sequences of Homologous Proteins

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The Alignment of Families of Homologous sequences.

First, find a family of Homologous sequences.

```
... APFELVISWKLIVESPAINCDWRTENGLANDSGMLVNOWPAI ...
... APYELVISQWKLIVESNPAINKDWRTYENGLANDSGMLVNOWAI ...
... APFELVISWKLIVESNPAINCDWRTENGLANDSGMLVNOWAI ...
... APFELVISQWKLIVESNPAINCDWRTENGLANDSGMLVNOWAI ...
... APYELVISWKLIVESNPINCDWRTENGLANDRSGLINOWAI ...
... APFELVISQWKLIVESNPAINCDWRTENGLANDSGMLVNOWLI ...
... APFELVISQWKLIVESNPAINCDWRTENGLANDSGMLVNOWAI ...
... APYELVISWKLIVESNPAINCDWRTENGLANDSGMLLNOWMI ...
```

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The Alignment of Families of Homologous sequences.

Then, align by inserting “-”s representing InDels, in each sequence.

```
... APFELVIS-WKLIVES-PAINCDWRT-ENGLANDSGMLV-NOWPAI ...
... APYELVISQWKLIVESNPAINKDWRTYENGLANDSGMLV-NOW-AI ...
... APFELVIS-WRLIVESNPAINCDWRT-ENGLANDSGMLV-NOW-AI ...
... APFELVISQWKLIVESNPAINCDWRT-ENGLANDSGMLV-NOW-AI ...
... APYELVIS-WKLIVESNP-INCDWRT-ENGLANDRSGMLINOW-AI ...
... APFELVISQWKLIVESNPAINCDWRT-ENGLANDSGMLV-NOW-LI ...
... APFELVISQWKLIVESNPAIN-DWRT-ENGLANDSGMLV-NOW-AI ...
... APYELVIS-WKLIVESNPAINCDWRT-ENGLANDSGMLL-NOW-MI ...
```

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The Alignment of Families of Homologous sequences.

Next, identify the columns where **Substitutions** and/or **InDels** have been predicted.

```
... APFELVIS-WKLIVES-PAINCDWRT-ENGLANDSGMLV-NOWPAI ...
... APYELVISQWKLIVESNPAINKDWRITYENGLANDSGMLV-NOW-AI ...
... APFELVIS-WRLIVESNPAINCDWRT-ENGLANDSGMLV-NOW-AI ...
... APFELVISQWKLIVESNPAINCDWRT-ENGLANDSGMLV-NOW-AI ...
... APYELVIS-WKLIVESNP-INCDWRT-ENGLANDRSGMLINOW-AI ...
... APFELVISQWKLIVESNPAINCDWRT-ENGLANDSGMLV-NOW-LI ...
... APFELVISQWKLIVESNPAIN-DWRT-ENGLANDSGMLV-NOW-AI ...
... APYELVIS-WKLIVESNPAINCDWRT-ENGLANDSGMLL-NOW-MI ...
```

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The Alignment of Families of Homologous sequences.

Then, identify the columns where full **Conservation** has been predicted.

```
... APFELVIS-WKLIVES-PAINCDWRT-ENGLANDSGMLV-NOWPAI ...
... APYELVISQWKLIVESNPAINKDWRTYENGLANDSGMLV-NOW-AI ...
... APFELVIS-WRLIVESNPAINCDWRT-ENGLANDSGMLV-NOW-AI ...
... APFELVISQWKLIVESNPAINCDWRT-ENGLANDSGMLV-NOW-AI ...
... APYELVIS-WKLIVESNP-INCDWRT-ENGLANDRSGMLINOW-AI ...
... APFELVISQWKLIVESNPAINCDWRT-ENGLANDSGMLV-NOW-LI ...
... APFELVISQWKLIVESNPAIN-DWRT-ENGLANDSGMLV-NOW-AI ...
... APYELVIS-WKLIVESNPAINCDWRT-ENGLANDSGMLL-NOW-MI ...
```

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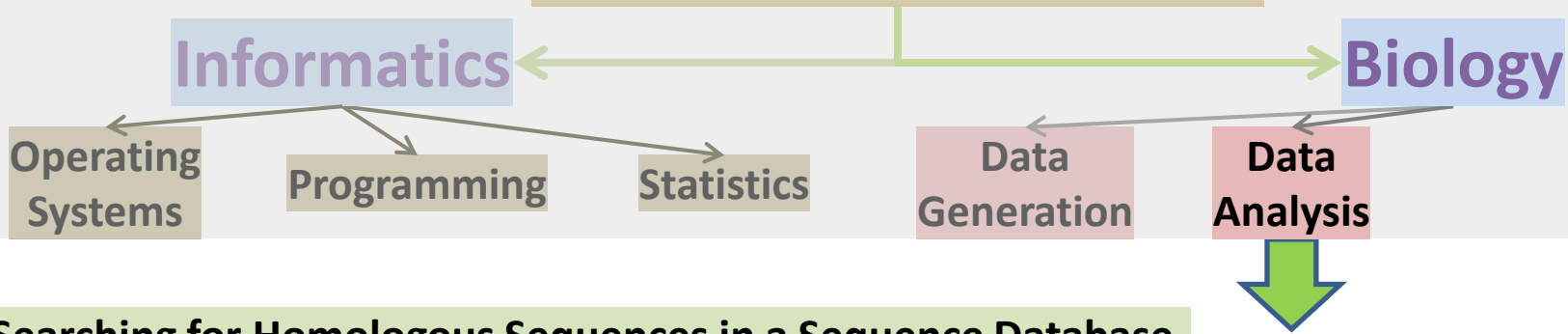


The Alignment of Families of Homologous sequences.

Finally ... Identify the Glorious Message!!!!.

... APFELVIS-WKLIVES-PAINCDWRT-ENGLANDSGMLV-NOWPAI ...
... APYELVISQWKLIVESNPAINKDWRTYENGLANDSGMLV-NOW-AI ...
... APFELVIS-WRLIVESNPAINCDWRT-ENGLANDSGMLV-NOW-AI ...
... APFELVISQWKLIVESNPAINCDWRT-ENGLANDSGMLV-NOW-AI ...
... APYELVIS-WKLIVESNP-INCDWRT-ENGLANDRSGMLINOW-AI ...
... APFELVISQWKLIVESNPAINCDWRT-ENGLANDSGMLV-NOW-LI ...
... APFELVISQWKLIVESNPAIN-DWRT-ENGLANDSGMLV-NOW-AI ...
... APYELVIS-WKLIVESNPAINCDWRT-ENGLANDSGMLL-NOW-MI ...

Bioinformatics Topics



Searching for Homologous Sequences in a Sequence Database.

Database searching is the most common Bioinformatics process by far.

Database searching is pairwise comparison repeated many times.

Non-optimal comparison methods are essential for practical reasons.

A list of matches, ordered by the improbability of occurring just by chance is generated.

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Searching for Homologous Sequences in a Sequence Database.

Database searching seeks "Similarity". Users seek "Homology".

Query	KLYPLRPQTPEPPPPPPPPPLPAAPPQP
Similarity	+L P +P P P PP P PP PP+P
Database Entry	RLTPPQPLMMPPRPTPPTPLPPATLTVPPRP

Homology?

Or 2 proteins including a lot of Prolines??

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Searching for Homologous Sequences in a Sequence Database.

Database searching seeks “Similarity”. Users seek “Homology”.

Query	TCTCCATTCGTAAAAAAAAAAAAAAAAAAAA
Similarity	
Database Entry	TCTTCCATTTGTAAAAAAAAAAAAAAAAAAAA

Homology?

Or 2 unrelated mRNAs??

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Searching for Homologous Sequences in a Sequence Database.

Database searching seeks “Similarity”. Users seek “Homology”.

Query

TTAGCAAGATCAGCCCTAACTCGGCATCTT

Similarity

| | | | | | | | | |

Database Entry

CTTGCGCGCTCTGTCTTGACGAGACACTTA

Homology?

A very unconvincing alignment!!

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Searching for Homologous Sequences in a Sequence Database.

Database searching seeks “Similarity”. Users seek “Homology”.

	T	T	A	G	C	A	A	G	A	T	C	A	G	C	C	T	A	A	C	T	C	G	G	C	A	T	C	T	T	
Query	L	A	R	S	C	L	T	R	H	L																				
Similarity	L	A	R	S	C	L	T	R	H	L																				
Database Entry	L	A	R	S	C	L	T	R	H	L																				
	C	T	T	G	C	G	C	T	C	T	T	G	A	C	G	A	G	A	C	A	C	T	T	A						

Homology?

Probable --- a perfect protein match??

In all circumstances – always align at the protein level wherever possible.

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Searching for simple sequence patterns Sequences in DNA.

Largely a matter of finding short sequences within longer ones.

Computationally trivial.

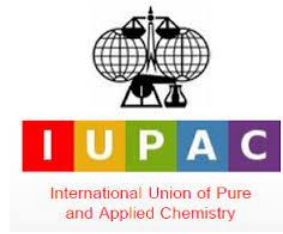
A concrete example is required:

[Restriction Mapping](#)

Detecting [Restriction Enzyme Recognition Sites](#)
is complicated by their redundancy.

Few Recognition Sites can be simply defined
using only the codes **A**, **C**, **G** and **T**.

The solution is to use the [Nucleotide Ambiguity Codes](#) defined by [IUPAC](#).



Unambiguous site (EcoRI):

G/AATC

Ambiguous site (PpuMI):

RG/GWCCY

Cut here

And here

TTAGCAAGATCAGGACCTACTCGGCATCTTCCTGGGTCCC

RGGWCCY

<u>Code</u>		<u>Meaning</u>
A		A
C		C
G		G
T/U		T/U
M	`aMino`	A C
R	`puRine`	A G
W	`Weak`	A T
S	`Strong`	C G
Y	`pYrimidine`	C T
K	`Keto`	G T
V	`not T`	A C G
H	`not G`	A C T
D	`not C`	A G T
B	`not A`	C G T
N	`aNy`	A C G T

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Searching for simple sequence patterns Sequences in Proteins.

Patterns can be derived manually to represent conserved regions of MSAs

Simple where conservation is 100%

```
... CQVLNPYYHWGQCGGIGWSGPTVCASGTT ...  
... CQYSNDYYHWGQCGGIGWSGCKTCTSGTT ...  
... CHVLNPYYQWGQCGGIGWTPSTTCASPYT ...  
... CSTLNPYYVWGQCGGIGWSGPTNCAPGSA ...  
... CVYSNDYYVWGQCGGIGWSGPTCCASGST ...
```

WGQCGGIGW

Pattern

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Searching for simple sequence patterns Sequences in Proteins.

Not so easy where conservation is less than perfect

An Amino Acid Alphabet including all ambiguities is not practical!

The solution is a simple syntax for ambiguous amino acid sequences.

```
... CQVLNPPYYHWKQCGGLGWSGPTVCASGTT ...
... CQYSNDYYHWGQCPGIGWSGCKTCTSGTT ...
... CHVLNPPYYQWAQCFVGVWTPSTTCASPYT ...
... CSTLNPYYVWLVQCYGIGWSGPTNCAPGSA ...
... CVYSNDYYVWAQCGGVGWSGPTCCASGST ...
```

W{P}QCxG[LIV]GW

Pattern

NOT a P

Anything

L or I or V

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Searching for simple sequence patterns Sequences in Proteins.

Match Here
Feature Site?

... GGSKFAWD GMYDKLRMLMRLWLQCKGVGWRTSFTQEQIEALEKEFERRQASNTPSHPI ...

W{P}QCxG[LIV]GW

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Searching for simple sequence patterns Sequences in Proteins.

Simple Protein patterns are of limited precision.

Only highly conserved regions can be described usefully.

Patterns cannot weight possibilities by frequency.

F
F
F
Y
F
F
F
F

[FY] --- or --- [F_Y] ?

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Searching for simple sequence patterns Sequences in Proteins.

Simple Protein patterns are of limited precision.

Patterns do not reflect commonly accepted substitutions.

F
F
F
F
F
F
F
F
F
F
F

--- or --- [F_Y] ?

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Searching for Protein properties with better models.

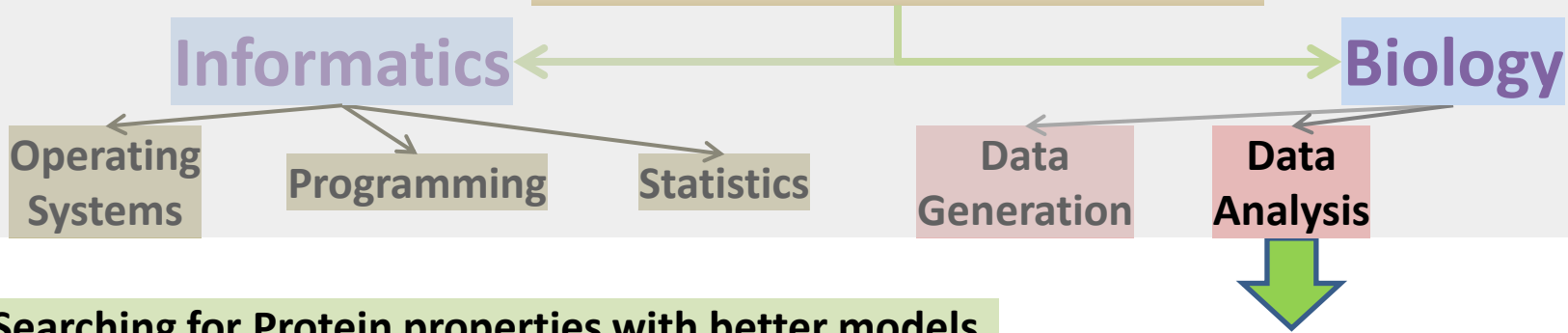
Again, start with an MSA of instances of the feature to be modelled.

Create a “suitable” representation of the relevant portion of MSA

Compare the model along other protein sequences was illustrated for simple patterns.

Where matches are detected, the corresponding protein property is likely to occur.

Bioinformatics Topics



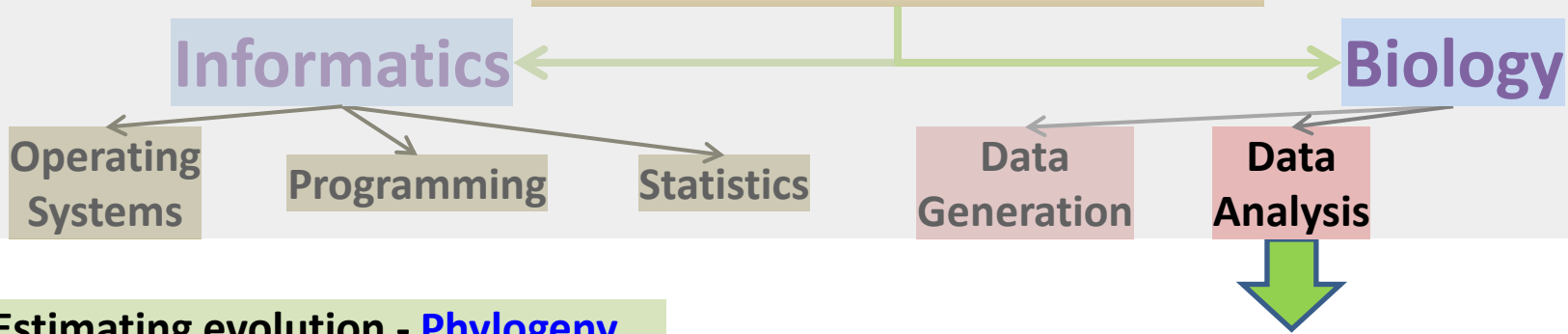
Searching for Protein properties with better models.

A variety of simple models have been developed (e.g. [Position Weight Matrices](#)) for a number of purposes, including:

- [Gene discovery in bacteria genomes \(DNA\)](#)
- [Early versions of 2D protein Structure Prediction](#)
- [Transmembrane Alpha Helix prediction](#)
- [TATA box Detection \(DNA\)](#)
- [Helix-Turn-Helix \(HTH\) Prediction](#)
- [Prediction of Coiled Coils](#)

The most powerful and prolific current profiles are [Hidden Markov Models \(HMMs\)](#)

Bioinformatics Topics



Estimating evolution - [Phylogeny](#).

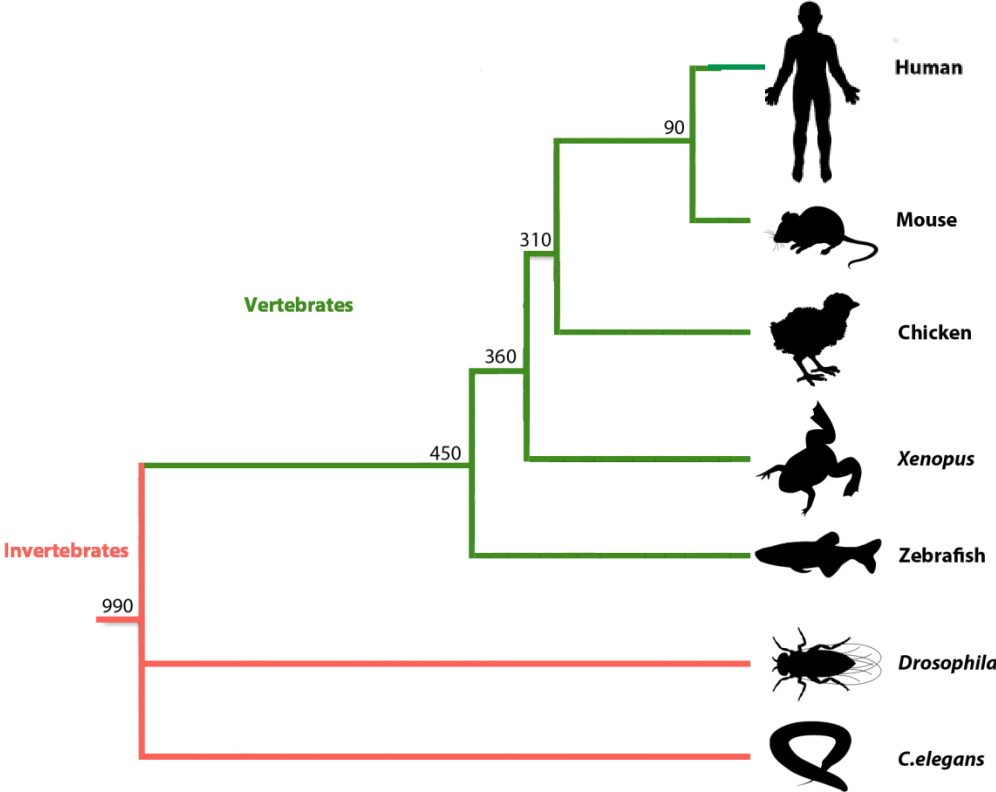
Broadly, the estimation of evolutionary history from available evidence.

“Evidence” does not have to be a carefully crafted MSA of Orthologous sequences from a range of organisms.

However, in the context of Bioinformatics, it invariably is.

Typically, conclusions of Phylogenetic analysis are represented as Evolutionary Trees.

Which are very Beautiful!!



My personal preference is for trees that place ME as far away from a MOUSE as possible!!!!

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Estimating evolution - [Phylogeny](#).

Phylogeny is another example of an analysis based on MSAs.

One very effective Phylogenetic strategy is to seek an answer to the question:

“What is the most probable Evolutionary Tree, given I believe this MSA to be perfect?”

Reinforcing how central is the role of Statistics in Bioinformatics.

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Protein structure prediction. Secondary Structure.

Essentially predicting the locations of Alpha Helices, Beta Sheets and Turns.

Modern methods employ Machine Learning to generate Artificial Neural Networks.

That is profiles computed by “learning” from observation of examples.

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Protein structure prediction. Secondary Structure.

Better predictions are obtained from MSA data than from individual protein sequences.

General principle being, the more information offered, the more reliable the prediction.

Some systems will automatically generate an MSA if offered a solitary protein sequence.

Prediction will be based on the MSA, computed by iterative database searching.

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Protein structure prediction. Tertiary Structure.

Predicting Tertiary Structure directly from Primary Structure is not currently practical.

De novo protein structure prediction requires better algorithms and more computing power.

Homology modelling requires a reliable Tertiary Structure for a homologous protein.

Tertiary Structure for a protein is predicted by comparison with the homologous structure.

Homology modelling is hampered by low volumes and uneven spread of available structures.

And now ... Once again ... Your turn!
Some issue for consideration, discussion and reaction

The Bioinformatics topics mentioned here do not constitute a comprehensive list. What would suggest is missing ... in order of importance?

The term algorithm was mentioned once or twice. There are slightly differing definitions. Pick the one you like best and justify your selection.

Define the three terms Homologue, Paralogue and Orthologue, being ever assiduous to ignore offensive American misspellings!

The is but one basic strategy for computing Pairwise Alignments that is considered optimal. However, this strategy can be implemented to compute either [Global Alignments](#) or [Local Alignments](#).

Just informally, [how do these two possibilities differ?](#)

Generally speaking, would you compute MSAs using a Global or a Local approach? Briefly justify your choice.

Generally speaking, would you conduct Database Similarity searches using a Global or a Local approach? Briefly justify your choice.

“Sequence alignment only makes sense for sequences representing Homologous entities”

A profound observation made by the ever sagacious David Philip Judge whilst sipping an eventide cup of [Tesco](#)’s very cheapest tea in the penthouse suite of his Ivory Tower (personal communication, 2016.06.10).

Consider and comment upon this fundamental truth.

“A Multiple Alignment of Homologous sequences which were a mixture of Orthologues and Paralogues would not be suitable as input data for [Phylogenetic analysis](#) ”

Another deep one from DPJ

Consider and comment upon this further pearl of enlightenment.

In the course of the dialogue for this presentation, there was mention of “Accepted Substitutions”, more formally referred to as “Accepted Point Mutations”, or ... if you enjoy clumsy for the sake of a pronounceable acronym, “[Point Accepted Mutation](#)” (PAM).

How would you informally define an “Accepted Point Mutation”?

The [Extended syntax for ScanProsite](#) is the most common syntax used for protein pattern definition. [ScanProsite](#) being the program for searching the of the [Prosite database](#). Prosite was first created way back in the 1980s and, initially, was composed exclusively of protein patterns.

There is no great value, at this stage, to be entirely familiar with this very simple syntax. However, from the hints in this presentation and a quick glance at the appropriate web pages, can you interpret the pattern?

C{P}x(3,7) [FY] (2) Wx(2) [VIL]

In the slides preceding, [Protein Domains](#) and [Protein Sequence Motifs](#) were mentioned with rather sparse explanation.

Define both of these terms and describe simply the [difference between them](#).

In the slide notes, there is mention of Position Weight Matrices (PWMs).

Can you say, simply, what a Position Weight Matrix might be and how it might be used?

What obvious property does a PWM possess that is lacking in a simple sequence pattern (or consensus sequence)?

The best secondary structure programs are reckoned to be around 80% accurate.

It is further suggested that 80% is about as good as it is possible to achieve.

Stated simply, why would you suppose that 100% accuracy might be unobtainable?

Hint: Do you think that two human experts, given the very best evidence of Tertiary Structure, would also agree upon the exact amino acid positions where an Alpha Helix starts and finishes?

Homology Modelling is mentioned in the slides as a method for predicting tertiary structure when structure(s) of protein(s) homologous to the query protein are available. The process involves aligning the query protein with the known structure, using the known sequence as a guide.

It is also possible to predict Tertiary Structure when, known structures thought to be appropriate exist, but only for sequences that **ARE NOT HOMOLOGOUS**. In such cases, the Primary Sequence corresponding to the known structure will be of little assistance.

Tricky eh!? What are the name(s) for those types of method? **ONLY** if you can do so **VERY** simply. Say a few words to say how they overcome the lack of a homologous sequence.

It was noted in the slides that often different Protein Feature searches often do not exactly agree.

It is common for two services to agree upon the presence of a domain, but not upon its precise start and end positions within a protein.

Would you find this to be worrying? Surprising? If not, why not?

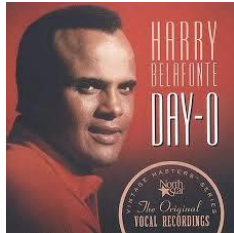
End of Part 2

BREAK!

More to come I fear ... but time for a swift cup of tea perchance?

Maybe time for a short jig? The whistling of a merry tune?

Or, mayhap, a delving into the melodic possibilities of youtube?
There be much good stuff there ... I offer you a few of my favourites.



Once fully refreshed Click on mon braves!

