

# **IB16S**

## Introductory Bioinformatics

12-16 December 2016

(Second 2016 run of this Course)

## Basic Bioinformatics Sessions

Practical 2: Pairwise Sequence Alignment

### **Sensitive Pairwise Alignment**

The purpose of this exercise is to look at some aspects of **Pairwise Sequence Alignment** using the most accurate methods available.

As hopefully has been discussed, sequences can be aligned using a **global** strategy, in which the two sequences being aligned are assumed to be homologous from end to end, or using a **local** approach, in which the sequences are assumed to just have homologous region(s).

#### **Global Pairwise Sequence Comparison**

First the **global** approach. In a previous exercise, you already have used the **blast** facility at the **NCBI** to perform crude pairwise alignment. **blast** also offers a sensitive option, so maybe that would be a good place to start.

So, once more to the NCBI home page (http://www.ncbi.nlm.nih.gov/). From there chose BLAST from the

Popular Resources list. Scroll down to the Specialized searches section and chose the

Global Align
Option.
Compare two sequences across their entire span (Needleman-Wunsch)

A choice of settings for **Nucleotide** or **Protein** alignment is offered. As we are going to investigate the alignment of DNA sequences, the default choice is fine. For the first sequence, browse for the file **pax6\_genomic.fasta**, which you created when looking at **Ensembl**. It contains the region of **Chromosome 11** containing the entire **PAX6** gene (with a few extra base pairs either end).

To specify the second sequence, you could load the file **pax6\_mrna.fasta**, but just typing the corresponding **Accession** code in the appropriate box seems far more sophisticated, so that is what I chose to do.

Open the **Algorithmic Parameters** section, and see that they are as one might expect. The defaults are fine here as the alignment to be computed is trivial (given the way **blast** will go about the task), so anything not outrageous should work.

Ask to Show results in a new window and then click on the Align button.

After some significant Rollin' and Tumblin' blast will proclaim its lyrical conclusions. First examine the Dot Matrix View. This sort of representation has rather gone out of fashion in recent years. A shame, I say, this picture represents such a succinct summary of what should be expected of the textual alignment(s) that are the "real" detailed output of this sort of program.

How would you interpret this picture?

What do the diagonal(ish) lines represent?

What are the gaps in between the lines?

Which axis represents the genomic sequence and which the mrna?

	Nucleotide Protein			
	-	alignment	of two n	nucleotide sequences 🕢
	Enter accession number, gi, or FASTA sequence		Clear	Query subrange 😡
	Enter accession number, gi, or FASTA sequence		Clear	From To
	Or, upload file  Browse pax6_genomic.fasta  Job Title	•		
	Enter a descriptive title for your BLAST search	Ð		
,	Enter Subject Sequence Enter accession number, gi, or FASTA sequence		Clear	Subject subrange  From
				То
	Or, upload file Browse No file selected.	•		
	Align   ☑ Show results in a new window			
	☐ Algorithm parameters  Scoring Parameters			
,	Match/Mismatch Scores			
	Gap Costs Existence: 5 Extension: 2			
	Align Show results in a new window			



perfect alignments.

You must know what these aligned regions represent by now?

But, just in case:

What do you suppose these regions represent?\_

How many are there and do they correspond nicely to the lines of the Dot Matrix View?"

How many exons would you say this mrna has?

If one was to forgive the strange "bits" at the start, would you say blast seems to have done a reasonable job here?

I think I would.

The final alignment section even has a poly A tail!

Query	24301	TTTTGTGTAGTTCTGGCACAATATGGAAAATCAACTTACTCTTTCAGAGTTTGAGAGAAC	24360
Sbjct	1045	AGTTTGAGAGAAC	1057
Query	24361	${\tt CCATTATCCAGATGTGTTTGCCCGAGAAAGACTAGCAGCCAAAATAGATCTACCTGAAGC}$	24420
Sbjct	1058	CCATTATCCAGATGTGTTTGCCCGAGAAAGACTAGCAGCCAAAATAGATCTACCTGAAGC	1117
Query	24421	AAGAATACAGGTACCGAGAGACTGTGCAGTTTCACACTTTGTGATTCATACCATTTGTCT	24480
Sbjct	1118	AAGAATACAGGTA	1130
Query	24481	${\tt TTCCTAGAGACAGAGGTGCTTGTACAGAGTACTATTTATT$	24540
Sbjct			
Query	24541	${\tt AAGGTTCAGTCTGCTAAATGCTCTGCTGCCATGGGCGTGGGGGGGG$	24600
Sbjct			
Query	24601	${\tt CCAAGGTGGGGCTGGGCTCGACGTAGACACAGTGCTAACCTGTCCCACCTGATTTCCAGG}$	24660
Sbjct			
Query	24661	TATGGTTTTCTAATCGAAGGGCCAAATGGAGAAGAAGAAAAAACTGAGGAATCAGAGAA	24720
Sbjct	1131	TGGTTTTCTAATCGAAGGGCCAAATGGAGAAAAAAGAGAAAAACTGAGGAATCAGAGAA	1188
Query	24721	${\sf GACAGGCCAGCAACACCCTAGTCATATTCCTATCAGCAGTAGTTTCAGCACCAGTGTCT}$	24780
Sbjct	1189	GACAGGCCAGCAACACCTAGTCATATTCCTATCAGCAGTAGTTTCAGCACCAGTGTCT	1248
Query	24781	ACCAACCAATTCCACAACCCACCACCACCGGGTAATTTGAAATACTAATACTACGAATCAA	24840
Sbjct	1249		1278
Query	24841	${\tt TGTCTTTAAACCTGTTTGCTCCGGGCTCTGACTCTCACTCTGACTACTGTCATTTCTCTT}$	24900
Sbjct			
Query	24901	GCCCTCAGTTTCCTCCTTCACATCTGGCTCCATGTTGGGCCGAACAGACACAGCCCTCAC	24960
Sbjct	1279	TTTCCTCCTCACATCTGGCTCCATGTTGGGCCTAACAGACACAGCCCTCAC	1330
Query	24961	${\tt AAACACCTACAGCGCTCTGCCGCCTATGCCCAGCTTCACCATGGCAAATAACCTGCCTAT}$	25020
Sbjct	1331	AAACACCTACAGCGCTCTGCCGCCTATGCCCAGCTTCACCATGGCAAATAACCTGCCTAT	1390
Query	25021	GCAAGTAAGTGCGGCTGGTGGCCTGCATAACCCAGGCCCCAGAGAAGTGAGGAGTGG	25080
Sbjct	1391	 GCAA	1394
Query	25081	${\tt CTCAGGGCCTGCGGACCTCATTGGCTGTGTCTGCACCCTTGAGAGCTTTTCGCACTACAG}$	25140
Sbjct			
Query	25141	TGATTGGCTTGACCAGTCAAGTCGGAGACAGTCAATCCCATCACTTTTAAGTGATTGACT	25200
Sbjct			

Sbjct Query Sbjct 1547 Sbjct 1604

Wonderful, but it is not safe to assume that just selecting any service that claims to do a sensitive global pairwise alignment will just work for any pair of sequences. I fact, pretty though it appears, the alignment blast has generated is not as entirely logical as it might first seem. For example, consider:

How might the gap around **24,500** in the genomic sequence been positioned more intelligently?

#### **Practical 2: Pairwise Alignment**

Next, try aligning the same two sequences with another program Global Alignment (implementing the same algorithm) at the EBI.

Go Sequence Alignment **Pairwise** (http://www.ebi.ac.uk/Tools/psa/). From there, select the Nucleotide option for the Global Alignment program Needle. Needle implements the best global pairwise algorithm faithfully.



Tuesday 6 December 2016

Global alignment tools create an end-to-end alignment of the sequences to be aligned. There are separate forms for protein or nucleotide sequences.

Needle @ (EMBOSS)

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

Nucleotide 🔌 Nucleotide

Load up the first sequence from pax6 genomic.fasta.

Load up the second sequence from pax6 mrna.fasta.

Click on the **More options** button to see what parameters you can set. They should be as you might expect. The defaults are fine for the first run.

Click on the **Submit** button to get **Needle** into action.

11	22101	TCTTGTAACACAATGTGGCCCGCTGCACGCCTCAAGAGAATCCT	22144
HUMOCLHMB	1	TATCGATAAGTT	12
11	22145	TTTGTTGTCCGCGCTCATTGTAGCCTCAAAAT-TCTGCCCACGAAAGTTT	22193
HUMOCLHMB	13	TTTTTTTTATTGTCAATCTCTG	34
11	22194	GCCAACGCTCCTGCCCCAGGAGTTTAATAGTTTCCCTTACTCGCGGGGCA	22243
HUMOCLHMB	35	.      .       . .   . .      TCTCCT-TCCCAGGAATCTGAGGATTGCTCTTACACAC	71
11	22244	TTGTGCAGCGCTGAAAAGCAGCCCCTCGCTATTCAAGTGTTGGTGGTCA-	22292
HUMOCLHMB	72	-  -  -  -  -	95
11	22293	TCTCAATAG-ATCTCCAAGGGCCCATATGGTGGCCAGTGCCGATGAA	22338
HUMOCLHMB	96	.    .    AACTCTCACCAGCAACTCC	114
11	22339	TCCGCCTGTTTAAATGGGGGAGAAAGTTGGGGTTTTAAAACAT	22381
HUMOCLHMB	115	ACACCGTCATTTCAAACCATTGTGGTC	147
11	22382	TTCAAAGTTCCTGAAAAGATCCCACT	22407
HUMOCLHMB	148	TTCAAGCAACAACAGCAGCACAAAAAACCCCAACCAAACAAAACTCTTGA	197
11	23546	${\sf TACCTTGGGAATGTTTTGGTGAGGCTGTCGGGATATAATGCTCTTGG}$	23592
HUMOCLHMB	804	ATGTTGAACGGCAGACCGGAAGCTGG	830
11	23593	AGTTTAAGACTACACCAGGCCCCT-TTTGGAGGCTCCAAGTTAATCCA	23639
HUMOCLHMB	831	GGTATCCGGG	856
11	23640	AATTTCTCTTACCATCCTATTCTTTTTGTTCCAGATGGCTGCCAGC	23685
HUMOCLHMB	857	GACTTCGGTGCCAGGCCAACCTACGCAAGATGGCTGCCAGC	897
11	23686	AACAGGAAGGAGGGGGAGAATACCAACTCCATCAGTTCCAACGGAGAA	23735
HUMOCLHMB	898		947
11	23736	GATTCAGATGAGGCTCAAATGCGACTTCAGCTGAAGCGGAAGCTGCAAAG	23785
HUMOCLHMB	948		997
11	23786	AAATAGAACATCCTTTACCCAAGAGCAAATTGAGGCCCTGGAGAAAGGTG	23835
HUMOCLHMB	998	AAATAGAACATCCTTTACCCAAGAGCAAATTGAGGCCCTGGAGAA	1042
11	23836	${\tt ATAGAGTTTTTCAAAGTAGAGAAGCAGTAAATCAAAGTAAATGCCACATC}$	23885
HUMOCLHMB	1043		1042
11	24636	${\sf TAACCTGTCCCACCTGATTTCCAGGTATGGTTTTCTAATCGAAGGGCCAA}$	24685
HUMOCLHMB	1125		1153
11	24686	ATGGAGAAGAAGAAAAACTGAGGAATCAGAGAAGACAGGCCAGCAACA	24735
HUMOCLHMB	1154		1203
11	24736	CACCTAGTCATATTCCTATCAGCAGTAGTTTCAGCACCAGTGTCTACCAA	24785
HUMOCLHMB	1204	CACCTAGTCATATTCCTATCAGCAGTAGTTTCAGCACCAGTGTCTACCAA	1253
11	24786	CCAATTCCACAACCCACCACCACGGGTAATTTGAAATACTAATACTACGA	24835
HUMOCLHMB	1254		1277
11	24836	ATCAATGTCTTTAAACCTGTTTGCTCCGGGCTCTGACTCTCACTCTGACT	24885
HUMOCLHMB	1278		1277
11	24886	ACTGTCATTTCTCTTGCCCTCAGTTTCCTCCTTCACATCTGGCTCCATGT	24935

24936 TGGGCCGAACAGACACAGCCCTCACAAACACCTACAGCGCTCTGCCGCCT

25036 TGGTGGTGGCCTGCATAACCCAGGCCCCAG--AGAAGTGAGGAGTGGCTC 25084 AGGGCCTGCGGACCTCAT----TGGCTGTGTCTG--CACCCTTGAGAGC ||| |||||||||||||||||||||||||||||-cct----cctcatactcctgcatg---ctgcccacc

Well! Nothing like as convincing as the alignment blast produced!

Alignment does not even begin until over 22,000 base pairs along the genomic sequence. Even then it is not convincing, as in wrong, if we accept the results already obtained from blast as a fair approximation of the truth.

There are some well aligned regions after genomic position 23,600.

Then a resumption of chaos after **25,000** or so.

How many convincingly aligned regions did you see?

How many did you expect?\_

Clearly, this alignment is not correct. Can you explain why?

IIIMOCI HMB

HUMOCLHMB

HUMOCLHMB

11

11

I assume you have all read the lucid answers to the question above? If so, I am confident you will agree that there are 3 ways to get an answer, similar to that generated by **blast**, from the tools offered at the **EBI**. They are:

Make gap penalties so cheap that Needle will have no excuse not to avoid gaps anywhere they are needed. This works if you use a gap opening penalty of 1.0 (the lowest allowed by both the program and the web interface) and a gap extension penalty of **0.0**, allowed by the program but not by the EBI web interface!! The lowest value the web interface allows is **0.0005**, which really should be sufficiently small, but proveably is not. The most important question being "Why would a web interface restrict a program's capabilities other than to prevent excessive resource use?". I have no answer for that one, I will just petulantly include my successful low gap alignment (made without a web interface) in your **Backup Results** directory and retire with self righteous hauteur!

Actually, using gap penalties to suit huge gaps that are really introns, will only work when the exons are so similar (as here) that any gap penalties will work for their alignment. Generally, you need to pick gap penalties to optimise exon alignment. So this is a very horrible way to "fix" the situation anyway.

- Tell Needle to penalise the gaps it puts at either end of the alignment in the same way it penalises gaps it puts in the middle. By default, end gaps are free!! Which is not very logical here.
- Use **Stretcher**, which uses essentially the same algorithm as **Needle**, except, it also applies a bit of common sense (heuristics, if you like). Stretcher takes a look at the sequences before it starts to do any serious computation. It identifies any "good regions" (all 12 exon matches in this case) and then says "OK, I am definitely having those, how best can I deal with the rest?". In essence, Stretcher does a quick Dot Matrix View before it starts and so only goes to work when it has a pretty good idea what the answer should look like It works in this case, but not always. Stretcher is faster than Needle but does not necessarily generate the highest scoring alignment. Stretcher works in a fashion far closer to the way a human would work, which has to be good! Well, usually anyway.

Tools > Pairwise Sequence Alignment > EMBOSS Stretcher

So, try the Needle with penalised End Gaps approach by returning to the Needle launch page from your results. You should find the two sequences are still selected, so you should only have to click on More Options again and change the END GAP PENALTY field from false to true.

	STEP 2 - S	et your p	airwise ali	gnment op	otions					
,	MATRIX				GAP OPE	EN	GAP EXTEND		OUTPUT FORMAT	
	DNAfull			*	10	*	0.5	▼	pair	*
	END GAP P		END GAP			PEXTEND				
	true	▼	10	▼	0.5	•				

Click on the **Submit** button and **Needle** will be on the road again.

How many matching regions are there this time?\_\_\_\_\_

Is the count **now** roughly as you would expect?

Finally, check that **Stretcher** works as expected.

Go again to the Pairwise Sequence Alignment EBI page (http://www.ebi.ac.uk/Tools/psa/).

From there, select the Nucleotide option for the Global Alignment program Stretcher.

Load up the sequences exactly as for Needle.

Take a look at the parameters and see there is nothing unexpected hiding there.

Set Stretcher sequence rope stretching.

How do you feel about the results this time?

How do you think **blast** achieve the correct results without any fuss?

Pairwise Sequence Alignment (NUCLEOTIDE)  EMBOSS Stretcher calculates an optimal global alignment of two sequences using a modification of the classic dynamic programming algorithm which uses linear space.
This is the form for nucleotide sequences. Please go to the <u>protein</u> form if you wish to align protein sequences.
STEP 1 - Enter your nucleotide sequences
Enter or paste your first nucleotide sequence in any supported format:
Or, upload a file: Browse pax6_genomic.fasta
AND  Enter or paste your second nucleotide sequence in any supported format:
7_1
Or, <u>upload</u> a file: Browse pax6_mrna.fasta
STEP 2 - Set your pairwise alignment options
MATRIX GAP OPEN GAP EXTEND OUTPUT FORMAT
DNAfull   T   16   T   4   T   pair   T   T   T   T   T   T   T   T   T
STEP 3 - Submit your job
Be notified by email (Tick this box if you want to be notified by email when the results are available)
Submit

#### Pairwise Sequence Comparison using Specialised Software

None of the alignments generated thus far have been entirely correct.

By persuading the general global alignment software to treat huge gaps (i.e. the introns) in some sort of special manner, a reasonable answer was obtained. However, the general software could not know that something more than just **Substitutions** and **Indels** were at issue here. Consequently, it stood no chance of dealing with the intron/exon boundaries sensibly.

The solution is not to fiddle around with the parameters of the general tools. Aligning **mRNA**s with **Genomic** sequence is simply not "General Alignment". It is an example of a problem that is sufficiently particular to require specialised software for an optimal solution.

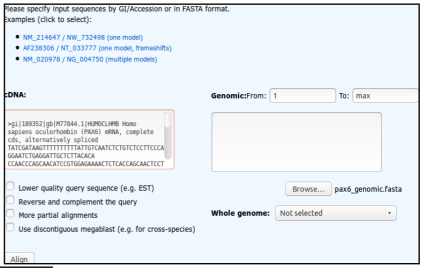
There is a program in the **EMBOSS** package (the same collection of programs as **Needle** and **Stretcher**), called **est2genome**, which is specifically designed for the alignment of cDNA/mRNA and genomic sequences. **est2genome** (and similar programs) may assume much more about the sequences to be aligned than can a general purpose alignment program. Gaps representing introns can be placed far more accurately if they are **known** to represent introns. Programs such as **est2genome** seek the highly conserved bases that occur at intron/exon boundaries, **C/T** rich intronic regions, **polyA** regions and **Stop/Start** codons to assist its detection of exons and gene structures.

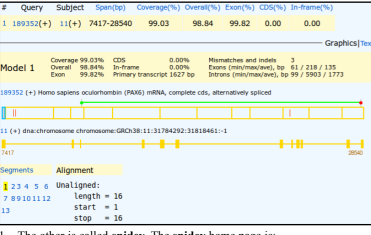
**est2genome** is a fine program, but there are two other options offered at the **NCBI** in America that do the same job, I think, somewhat more nicely. Of these, I choose the program called **splign** for this exercise on the grounds it is the more sophisticated service<sup>1</sup>. To investigate, go to the home of **splign** at:

#### http://www.ncbi.nlm.nih.gov/sutils/splign

Click on the Online button. In the Genomic section, Browse to upload pax6 genomic.fasta.

In the cDNA section, paste the sequence pax6\_mrna.fasta. Where cDNA and Genomic sequences share exons that are nearly identical, splign uses the comparison algorithm megablast (default). Where exons are less similar (e.g. when the cDNA and Genomic sequences are from different organisms) the more sensitive option discontinuous megablast, is a better choice<sup>2</sup>. Note the option to compare your cDNA with a Whole genome (including Human). Today, the default options are fine. Click the Align button.





Your results will appear showing the cDNA split into 12 sections (the predicted exons) corresponding to 12 regions of the genomic sequence indicated by yellow rectangles. A 13<sup>th</sup> region of 16 base pairs is displayed and declared to be **unaligned**. These are the 16 mystery base pairs at the start of this particular mRNA that **Needle** and **Stretcher** had trouble treating sensibly also. I wonder what they are?

Any theories?

1 The other is called **spidey**. The **spidey** home page is:

http://www.ncbi.nlm.nih.gov/IEB/Research/Ostell/Spidey/

It is pretty straight forward to use, just follow your nose. **est2genome**, **splign** and **spidey** should all give the same answer for this simple alignment and very similar answers for all such problems.

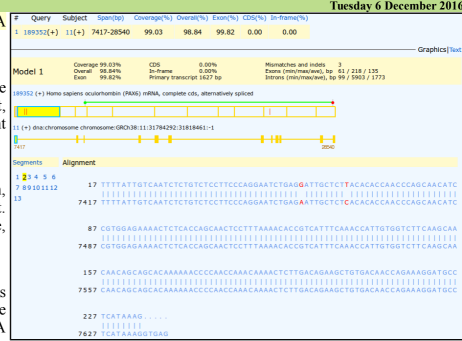
Why this is so will be considered later when we look at the database searching program **blast**.

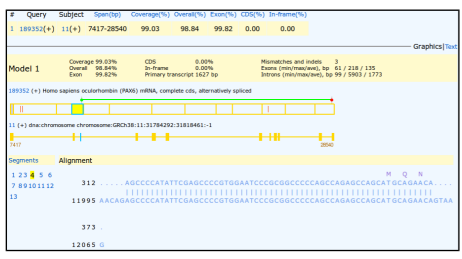
Practical 2: Pairwise Alignment
Click on the first exon section of the cDNA
display.

Here there shows two **substitutions**. These were also apparent in the successful **blast**, **Needle** and **Stretcher** alignments. You might have spotted them?

Though these are in a non-coding region, they could easily still be very significant. However, for the purposes of this exercise, let us assume they are not.

The **Start** (green) and **Stop** (red) codons delimiting the **CoD**ing **S**equence (**CDS**) are illustrated by the bar above the cDNA display.





Click on the exon including the green **Start** codon (the 3<sup>rd</sup>).

The first coding exon is now displayed with translation of the mRNA where appropriate.

The statistics at the top of the display include the claim that there are 3 discrepancies (Mismatches and Indels) between the cDNA and Genomic sequences.

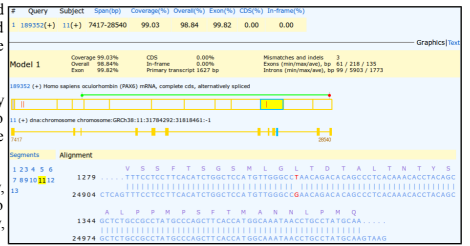
Two of these are the **substitutions** we have already seen in the first exon of the cDNA. The third is indicated by the red bar in the **10**<sup>th</sup> exon of the **cDNA** display.

#### Click on the 10<sup>th</sup> exon section of the cDNA display.

The third difference, a substitution, should be clear to see. Given it changes the coded protein, this substitution is likely to be the most significant.

Irritatingly, in the extreme! **spline** only translates the mRNA. So one has to work to discover the alternative suggested by the Genomic sequence.

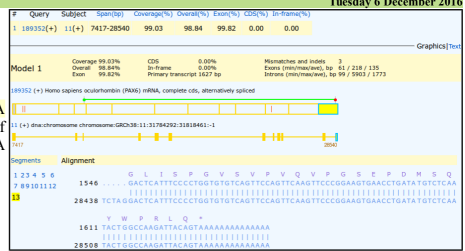
Vital if we were really doing this seriously, but for an exercise, it is fine to relax. I do not intrude on real life much and **it**, largely, leaves **me** untouched in grateful response.

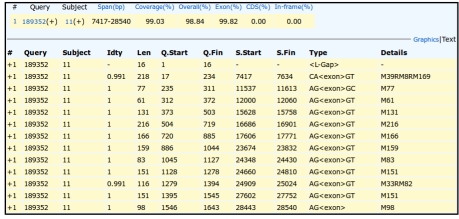


What is the amino acid corresponding to the mutated position in the **Genomic** sequence?

What are the **Genomic** and **mRNA** base positions corresponding to the mutation at amino acid position 33?\_

Click on the last exon section in the cDNA display. You should now see the final exon of the cDNA with the **Stop** codon and polyA region.





Finally, click on the **Text** link to view the textual summary of the **splign** results.

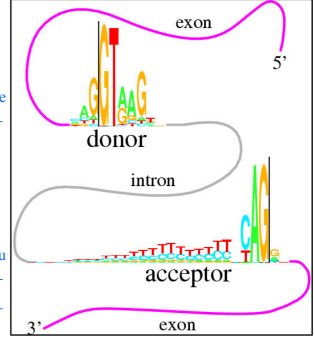
How do you interpret the **Details** column for exons 1 and 10?

Where is the 3<sup>rd</sup> substitution in the mRNA?

Where is the 3<sup>rd</sup> substitution in the Genomic Sequence?

Compare the predicted **splign** intron/exon boundaries with the conservation suggested by the logo<sup>3</sup>?

What deviation(s) from the model suggested by the logo can you see?\_\_\_\_\_



3 The original label for this very nice graphic is:

This figure shows two "sequence logos" which represent sequence conservation at the 5' (donor) and 3' (acceptor) ends of human introns. The region between the black vertical bars is removed during mRNA splicing. The logos graphically demonstrate that most of the pattern for locating the intron ends resides on the intron. This allows more codon choices in the protein-coding exons. The logos also show a common pattern "CAGIGT", which suggests that the mechanisms that recognize the two ends of the intron had a common ancestor. See R. M. Stephens and T. D. Schneider, "Features of spliceosome evolution and function inferred from an analysis of the information at human splice sites", J. Mol. Biol., 228, 1124-1136, (1992).

#### Sensitive Local Pairwise Sequence Comparison

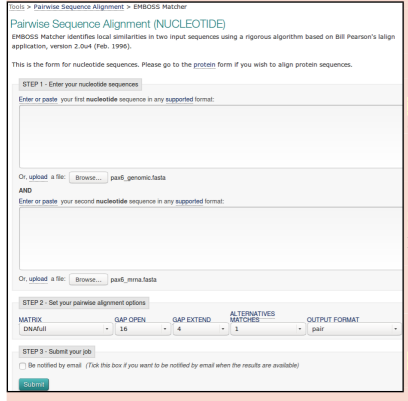
Finally, a swift look at sensitive local pairwise sequence alignment. You have already used blast to do a local pairwise alignment in the last Practical, when you aligned the two human genomic sequencing contigs that covered the PAX6 location in Chromosome 11. blast did not use a sensitive approach however, nothing subtle was required

for that particular alignment.

For a more accurate alignment, return to the Pairwise Sequence Alignment EBI page (http://www.ebi.ac.uk/Tools/psa/).

From there, select the Nucleotide option for the Local Alignment program Matcher.

Water or LALIGN would also be fine options, but I declare the nucleotide Matcher (EMBOSS) option of **Matcher** to be choice of the day.



Local Alignment

Local alignment tools find one, or more, alignments describing the most similar region(s) within the sequences to be aligned. There are separate forms for protein or nucleotide sequences.

Water @ (EMBOSS)

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

Nucleotide

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

Nucleotide

LALIGN @

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

Nucleotide

Load up the Genomic and mRNA sequences as you did for Needle.

Click on the More options button to see what parameters you can set. They should be as you might expect. The defaults are fine for the first run.

Click on the Submit button to get Matcher into Matchbox mode.

After due consideration of all the possibilities, **Matcher** will enrich your screen with its conclusions.

But, only one alignment? A good one, covering the highest scoring region of all those considered, but it cannot be the whole story, which must tell the tale of 12 exons! Here is but one.

In common with most local alignment programs, by default **Matcher** will only show you the single best local alignment between two sequences.

A good reason to have a **Dot Matrix View** to inform one of roughly what to expect, which is not one miserable alignment in this case.

	11	16670	CACTTCCCCTATGCAGGTGTCCAACGGATGTGTGAGTAAAATTCTGG	16716
•	HUMOCLHMB	485		534
t	11	16717	${\tt GCAGGTATTACGAGACTGGCTCCATCAGACCCAGGGCAATCGGTGGTAGT}$	16766
,	HUMOCLHMB	535	GCAGGTATTACGAGACTGGCTCCATCAGACCCAGGGCAATCGGTGGTAGT	584
	11	16767	AAACCGAGAGTAGCGACTCCAGAAGTTGTAAGCAAAATAGCCCAGTATAA	16816
	HUMOCLHMB	585	AAACCGAGAGTAGCGACTCCAGAAGTTGTAAGCAAAATAGCCCAGTATAA	634
,	11	16817	GCGGGAGTGCCCGTCCATCTTTGCTTGGGAAATCCGAGACAGATTACTGT	16866
	HUMOCLHMB	635	GCGGGAGTGCCCGTCCATCTTTGCTTGGGAAATCCGAGACAGATTACTGT	684
	11	16867	CCGAGGGGGTCTGTACCAACGATAACATACCAAGCGTAAGTTCATTGAGA	16916
)	HUMOCLHMB	685	CCGAGGGGGTCTGTACCAACGATAACATACCAAGCGTGTCATCAATAAAC	734
t	11	16917	ACATCTGCCCTCCCTGCC 16934	
	HUMOCLHMB	735	-     -  -    -  AGAGTTCTTCGCAACCTGGC 754	

Of course, it is also miserable biologically! Matcher fails to align the exons accurately for all the same reasons that the **Needle** failed to represent the *biological* reality.

So, what can one do but try again! By returning to the **Matcher** launch page from your results. You should find the two sequences are still selected, so you should only have to click on More Options again and set the ALTERNATIVE MATCHES field 20.

is the most sensible cautious choice.

STEP 2 - Set your pairv	wise alignment options						
MATRIX	GAP OPEN	GAP EXTE	ND	ALTERNATIVES MATCHES		OUTPUT FORMAT	
DNAfull	▼ 16	₹ 4	٧	20	٧	pair	*

24655 TCCAGGTATGGTTTTCTAATCGAAGGGCCAAATGGAGAAGAAGAAAAA

1222

24804

Actually, as you know there are only 12 exons. And that some might well be close enough to be included in the same alignment, you do not need to go as high as 20. However, the web interface restricts choice (WHY!?) such that this

HUMOCLHMB

11 24705 CTGAGGAATCAGAGAAGACAGGCCAGCAACACCCTAGTCATATTCCTAT HUMOCLHMB Click on the Submit button and Matcher will trust 24755 CAGCAGTAGTTTCAGCACCAGTGTCTACCAACCAATTCCACAACCCACCA 11 and obey. **HUMOCLHMB** 11 24805 CACCGGGTAATTTGAAATACTAATACTACGAATCAATGTCTTTAAACCTG |||||| 1273 CACCGG------HUMOCLHMB At the top of your output will be some nice believable 24855 TTTGCTCCGGGCTCTGACTCTCACTCTGACTACTGTCATTTCTCTTGCCC local alignments, some involving more than one exon. **HUMOCLHMB** 

**Matcher** tries to make each alignment as long as it can, stopping only when, to stretch the alignment any further would involve the alignment score deceasing due to the necessity for gap penalties.

```
1272
                                                  24854
                                                   1278
                                                  24904
          24905 TCAGTTTCCTCCTTCACATCTGGCTCCATGTTGGGCCGAACAGACACAGC
                                                  24954
           HUMOCLHMB
          24955 CCTCACAAACACCTACAGCGCTCTGCCGCCTATGCCCAGCTTCACCATGG
                                                  25004
          HUMOCLHMB
                                                   1374
          25005 CAAATAACCTGCCTATGCAA 25024
           1375 CAAATAACCTGCCTATGCAA
HUMOCLHMB
```

```
# Aligned_sequences: 2
 1: 11
# 2: HUMOCLHMB
# Matrix: EDNAFULL
 Gap penalty: 16
 Extend_penalty: 4
 Length: 58
 Identity:
                 39/58 (67.2%)
 Similarity:
                39/58 (67.2%)
                 6/58 (10.3%)
 Gaps:
# Score: 83
#
11
               2353 GCTGGACGCCACCCGGCGCCAGA--GCCGGGC---CTGAGGAGCGGGGTC
                                                                         2397
                ||.|||.||||||||...||| |...|| ||...|| ||...||| 425 GCCGGACTCCACCCGGCAGAAGATTGTAGAGCTAGCTCAC-AGCGGGGCC
HUMOCLHMB
                                                                          473
11
               2398 TGGCCGGG 2405
                474 CGGCCGTG
# Aligned_sequences: 2
# 2: HUMOCLHMB
# Matrix: EDNAFULL
 Gap_penalty: 16
# Extend_penalty: 4
#
# Length: 46
 Identity:
                31/46 (67.4%)
 Similarity:
                31/46 (67.4%)
  Gaps:
                 1/46 ( 2.2%)
# Score: 83
#
```

Go to far down the list of alignments and you will realise what a literal interpretation **Matcher** has of its duties.

You asked for **20** alignments?

So here are the best 20 alignments and it is entirely up to you to decide where "silly" begins.

Not too difficult in this case I suggest.

Why do you suppose your aligned exons are not presented in the correct positional order?

11417 ACAGTTTGACTGAGCCCTAGATGCATGTTTTTT-CCTGAGAGTGA 11461 

#### THE END

#### **Model Answers to Questions in the Instructions Text.**

#### **Notes:**

For the most part, these "**Model Answers**" just provide the reactions/solutions I hoped you would work out for yourselves. However, sometime I have tried to offer a bit more background and material for thought? Occasionally, I have rambled off into some rather self indulgent investigations that even I would not want to try and justify as pertinent to the objective of these exercises. I like to keep these meanders, as they help and entertain me, but I wish to warn you to only take regard of them if you are feeling particularly strong and have time to burn. Certainly not a good idea to indulge here during a time constrained course event!

Where things have got extreme, I am going to make two versions of the answer. One starting:

#### Summary:

Which has the answer with only a reasonably digestible volume of deep thought. Read this one.

The other will start:

#### Full Answer:

Beware of entering here! I do not hold back. Nothing complicated, but it will be long and full of pedantry.

This makes the Model answers section very big. <u>BUT</u>, it is not intended for printing or for reading serially, so I submit, being long and wordy does not matter. Feel free to disagree.

From your investigations of Global Alignment:

What do you suppose these regions represent?

**Exons** 

#### How might the gap around **24,500** in the genomic sequence been positioned more intelligently?

**blast** has positioned a gap in this region merely to maximize the overall alignment score. There is more than one way of achieving this simple goal. However, if it were to be recognized that the gap to be positioned was to represent an intron, then one of the arithmetically equivalent options becomes far more attractive than the others. This "best" option is not the one chosen by **blast**, which is forgiveable as **blast** had nor reason to expect an intron and was not written to understand the properties of introns anyway.

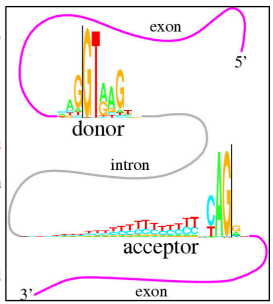
The alignment chosen for this region by **blast** was:

Genomic	24421	AAGAATACAGGTACCGAGAGACTGTGCAGTTTCACACTTTGTGATTCATACCATTTGTCT	24480
mRNA	1118	AAGAATACAGGTA	1130
		• • •	
Genomic	24601	CCAAGGTGGGGCTGGGCTCGACGTAGACACAGTGCTAACCTGTCCCACCTGATTTCCAGG	24660
mRNA			
Genomic	24661	TATGGTTTTCTAATCGAAGGGCCAAATGGAGAAGAGAAG	24720
mRNA	1131	TGGTTTTCTAATCGAAGGGCCAAATGGAGAAGAAGAAAAAACTGAGGAATCAGAGAA	1188

Shifting the gap 3 places to the left neither changes the size of the gap nor the perfection of the alignment either side of the gap and so does not affect the alignment score.

However, it does mean the gap begins with an **GT** and ends with a **AG** which is what one might expect if it were known that the gap represented an intron. I include the beautiful **Intron/Exon** logo. As you might gather, I rather like this one.

So, if **blast** was a little better informed, the improved alignment would have been:



Genomic	24421	AAGAATACAG <u>GT</u> ACCGAGAGACTGTGCAGTTTCACACTTTGTGATTCATACCATTTGTCT	24480
mRNA	1118	AAGAATACAG	1130
		• • •	
Genomic	24601	CCAAGGTGGGGCTGGGCTCGACGTAGACACAGTGCTAACCTGTCCCACCTGATTTCCAGG	24660
mRNA		G	
Genomic	24661	TATGGTTTTCTAATCGAAGGGCCAAATGGAGAAGAAGAAAAACTGAGGAATCAGAGAA	24720
mRNA	1131	TATGGTTTTCTAATCGAAGGGCCAAATGGAGAAGAAGAAAAAACTGAGGAATCAGAGAA	1188

This is the alignment that one might expect from any program customized to align **mRNA** with **Genomic** sequence, as you will see in the fullness of time.

How many convincingly aligned regions did you see?

4

#### How many did you expect?

12, as that was how many **blast** found, not including the silly ones at the beginning.

The 4 that were found correspond the the 4 diagonal lines grouped together in the **Dot Matrix View** made by **blast**.



#### Clearly, this alignment is not correct. Can you explain why?

This alignment algorithm only wishes to maximise an alignment score. It sees <u>ALL</u> the high scoring exon regions, however, as the gaps between many of the exons (introns that is) are so long that the penalties for representing them correctly are greater than the gain achieved by the inclusion the extra exons in the alignment. Arithmetically, it is better to align all the exons either side of the 4 exons that were aligned sensibly, in the biologically improbably fashion shown. Arithmetically the best alignment, biologically ridiculous!

This behaviour is exaggerated because this program regards the enormous gaps in has suggested at the start and end of the alignments as "free". Some global alignment programs (including this one if you ask politely, as you will see) offer the option of penalising the ends gaps in the same way as for internal gaps. Normally, not penalising end gaps is sensible as it allows for the sequences to have slightly different lengths. In this case, penalising end gaps will result in a far better alignment.

Had you used **stretcher** (also offered by the **EBI**) you would have got a much improved answer in this case (but not necessarily in generally). This is because **stretcher** works in a way far closer to the way an informed human might think. **stretcher** does not mindlessly insist of the highest alignment score. Instead, it looks for all the high scoring regions (i.e. all the exons) and then computes the best way to link them together. The result is a far more convincing alignment, but not the arithmetically best scoring answer.

#### How many matching regions are there this time?

Were you to trawl though your textual output carefully (or simply take my immaculate word for it), you would find 12 perfectly (or nearly so) aligned regions, implying 12 exons.

To be pedantic, the nicely aligned regions do not match the exons exactly (as has been discussed), but well enough to claim definite evidence for the number of exons. 12 is good enough for me.

#### Is the count **now** roughly as you would expect?

Yes, exactly the same as **blast** predicted in the first place. More exons that **17** might have been a surprise as that is how many the gene record for **PAX6** at the **NCBI** suggested. Any given transcript may have less than **17** exons or exactly **17** exons, but not more than **17** exons if the heroes of the **NCBI** are not mistaken.

#### How do you think **blast** achieve the correct results without any fuss?

The only way blast could have got the right answer, as it did, would be to use one of the strategies listed previously. **blast** did not use the horrible idea of making gaps super cheap! Not only is that a disgustingly dirty trick, but **blast** actually declares that it is using quite sensible gap penalties.

Leaving **penalising end gaps** and/or using the same sort of heuristics employed by **stretcher**. I would strongly suspect **blast** uses a **stretcher** approach. After all, **blast** has clearly already identified all the "promising regions" in order to construct its **Dot Matrix View**. Also the **stretcher** strategy is similar to that of all **blast** searches (discussed in the next Practical). Finally, **blast** is often used to align very long DNA sequences to detect very strongly similar large regions. This is exactly what the faster (if less pure) **stretcher** approach is all about.

From your investigations comparing mRNA/cDNA with genomic DNA:

#### What is the amino acid corresponding to the mutated position in the **Genomic** sequence?

The top sequence is the mRNA. **spline** is kind enough to explicitly inform us that the "mutated" codon, CTA, will be expressed a

Leucine.

So, why not translate the **Genomic** sequence also **spline**?! Easy enough to look up. But I resent having to do so!

From this rather beautiful representation of the **Genetic Code**, I conclude:

mRNA CTA  $\rightarrow$  Leucine (L) Genomic CGA  $\rightarrow$  Arginine (R)

I checked, and this does not appear to be a substitution that is associated with any "interesting" phenotype.

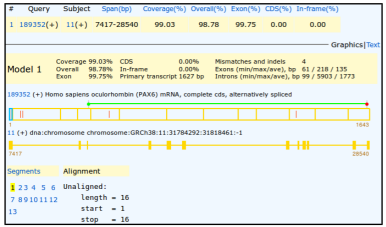
There is no real reason why it should. We did not pause to find out anything about the mRNA downloaded from the NCBI, The annotation is particularly unrevealing by itself (it is in **Backup Files** if you really want to check).

Let us simply assume it is a benign Accepted Point Mutation (PAM). Yes indeed, that feels comfortable. Not so very tricky this Science stuff after all what!

What are the Genomic and mRNA base positions corresponding to the mutation at amino acid position 33?

Remember the **Natural variation** at amino acid position **33**? You looked at it in passing during the course of the first exercise. It is a major cause of **Aniridia**. An **Alanine** mutated to a **Proline** at the end of a **Helix** vital to the **DNA Binding** function of the **PAX6** protein.

			1
	Natural variant i (VAR_008694)	29	I → S in AN. # 1 Publication ▼
1	Natural variant i (VAR_003811)	29	I → V in AN. # 1 Publication ▼
į	Natural variant i (VAR_008695)	33	A → P in AN.   ¶ 1 Publication ▼
•	Natural variant i (VAR_008696)	37 – 39	Missing in AN. # 1 Publication #
١	Natural variant i (VAR_008697)	42	$I \rightarrow S$ in AN; mild. $\P$ 1 Publication $\checkmark$
	Natural variant i (VAR_008698)	43	S → P in AN. # 1 Publication ▼
	Natural variant i (VAR_003812)	44	R → Q in AN. # 1 Publication ▼



**Spline** shows alignments for all exons and from those alignments the answer to this question is thus clearly available. To make finding the right spot in the alignment to study easier, I ran **Spline** again with an edited version of the **mRNA** (saved as **pax6\_mrna\_edited.fasta** amongst your cheat files) against the same **Genomic** sequence. Had there been a suitable **mRNA** sequence in the databases, I would have used it for the exercise, but there is not.

You should be able to clearly see the extra mutation is in the  $5^{th}$  exon.

Focussing on the 5<sup>th</sup> exon, the substitution is clear. Using the same methods as were used for the previous question, it is easy to confirm that the variation at amino acid position 33<sup>4</sup> amounts to:

#### **Affected Patient protein:**

 $CCT \rightarrow Proline(P)$ 

#### **Canonical protein:**

 $GCT \rightarrow Alanine(A)$ 

Squinting madly, you can also discover that the variation base positions are:

### 

#### **Affected Patient mRNA:**

Base position  $459 \rightarrow C$ 

#### **Wild Type Genomic DNA:**

Base position  $15714 \rightarrow G$ 

In case you were wondering, chasing these values around is a little more than tragic pedantry. You will need this information later when you investigate **Primer Design**. No need to take notes, I will remind you of what you need when the time comes. Here I just want to show how the values could be determined, if you had to. Not difficult, just tedious!

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<sup>4</sup> Proving beyond reasonable doubt that that substitution is exactly at amino acid position 33 requires a little more counting, dividing by 3 and subtracting the number you first thought of. For now, just trust me? I really am more honest than I look.

How do you interpret the **Details** column for exons 1 and 10?

#### Summary:

The **Details** column shows the alignments of each exon in a compressed format described in the **spline** documentation as illustrated.

)	
	11. Alignmer
5	transcript

Alignment transcript represents full details of the alignment in a form of a string composed of characters 'M', 'R', 'I' and 'D' where each character corresponds to an elementary command (Match, Replace, Insert or Delete) needed to transform the query segment into the subject segment. The string is encoded with RLE.

The majority of the exon alignments are trivial.

#	Query	Subject	Span(bp)	Coverag	ge(%)	Overall(%)	Exon(%)	CDS(%)	In-frame(	%)	
1	189352(+)	11(+)	7417-28540	99.0	03	98.84	99.82	0.00	0.00		
											Graphics Text
#	Query	Subject	Idty	Len	Q.Sta	art Q.	Fin S	.Start	S.Fin	Туре	Details
+1	189352	11	-	16	1	16	-		-	<l-gap></l-gap>	-
+1	189352	11	0.991	218	17	23	4 74	417	7634	CA <exon>GT</exon>	M39RM8RM169
+1	189352	11	1	77	235	31	1 1	1537	11613	AG <exon>GC</exon>	M77
+1	189352	11	1	61	312	37	2 1	2000	12060	AG <exon>GT</exon>	M61
+1	189352	11	1	131	373	50	3 1	5628	15758	AG <exon>GT</exon>	M131
+1	189352	11	1	216	504	71	9 10	6686	16901	AG <exon>GT</exon>	M216
+1	189352	11	1	166	720	88	5 1	7606	17771	AG <exon>GT</exon>	M166
+1	189352	11	1	159	886	10	44 23	3674	23832	AG <exon>GT</exon>	M159
+1	189352	11	1	83	1045	11	27 24	4348	24430	AG <exon>GT</exon>	M83
+1	189352	11	1	151	1128	12	78 24	4660	24810	AG <exon>GT</exon>	M151
+1	189352	11	0.991	116	1279	13	94 24	4909	25024	AG <exon>GT</exon>	M33RM82
+1	189352	11	1	151	1395	15	45 23	7602	27752	AG <exon>GT</exon>	M151
+1	189352	11	1	98	1546	16	43 28	8443	28540	AG <exon></exon>	M98

For example:

For Exon 2, spline informs us M77, meaning "There are 77 bases aligned and they all Match perfectly".

For Exon 4, spline informs us M131, meaning "There are 131 bases aligned and they all Match perfectly".

The only 2 interesting entries are those were there are some disagreements. That is, the entries for Exons 1 and 5, which, following the documentation, I translate thus:

#### Exon 1 – M39RM8RM169

An alignment of 218 bases, the first 39 of which Match perfectly (M39), there then follows an Replacement (R), a further 8 Matched bases(M8), a second Replacement (R) all finished off with 169 Matched bases (M169).

#### Exon 10 - M33RM82

An alignment of 116 bases, the first 33 of which Match perfectly (M33), there them follows a Replacement (R) and a further 82 Matched bases(M82).

Its a pity there are no Insertions (I) and Deletions (D), but this was the best mRNA I could find.



A point of pedantry to commence. From a different example, which included **InDels**, I got the display illustrated.

The exon was reported as: M53IM5IM43

This implies that the choice of Insertion (I) or Deletion (D) is made to describe the type of variation required to transform the cDNA (Query) sequence into the genomic (Subject). Hence the two InDels displayed here are considered to be Insertions.



Not that it is a vital issue, but I would have thought the other way around was more logical? That is, to consider the **genomic** sequence as the **reference** against which a particular **mRNA** might vary. In other words, what we see here would surely be more relevantly recorded as "This **mRNA/cDNA** has two **D**eletions relative to the **genomic** sequence which, presumably, attempts to represent the norm in the general population"? Just the reflection of an irretrievable pedant, but I am right, nevertheless!!!

In the documentation (see illustration in the **Summary** answer) it enigmatically states "The string is encoded with **RLE**.". Just in case, **RLE** stands for **Run-length encoding** which is succinctly defined by **Wikipedia**. In a nutshell, it is a very simple form of data compression that recognizes that:

#### 

can be compressed to:

60X

which has to be very effective for any data that has runs of identical characters of significant length. This is certainly the case here where one would expect long stretches of **M**s in most alignments. Of course, life would get tricky if the data included numeric characters, but that is not an issue here<sup>5</sup>.

I think it worth mentioning, that this way of representing an alignment is a simplification of CIGAR format. This format is used for SAM (Sequence Alignment Map) and BAM (Binary Alignment Map, exactly the same as

SAM, except compressed) files. You will be engulfed in SAM/BAM files if you ever do any Next Generation Sequencing (NGS).

CIGAR: CIGAR string. The CIGAR operations are given in the following table (set '\*' if unavailable):

Op BAM Description

M 0 alignment match (can be a sequence match or mismatch)

I 1 insertion to the reference

D 2 deletion from the reference

So, straight from the SAM/BAM Format Specification I copy the table of CIGAR enlightenment.

Op	BAM	Description
M	0	alignment match (can be a sequence match or mismatch)
I	1	insertion to the reference
D	2	deletion from the reference
N	3	skipped region from the reference
S	4	soft clipping (clipped sequences present in SEQ)
H	5	hard clipping (clipped sequences NOT present in SEQ)
P	6	padding (silent deletion from padded reference)
=	7	sequence match
X	8	sequence mismatch

- H can only be present as the first and/or last operation.
- S may only have H operations between them and the ends of the CIGAR string.
- For mRNA-to-genome alignment, an N operation represents an intron. For other types of alignments, the interpretation of N is not defined.
- Sum of lengths of the M/I/S/=/X operations shall equal the length of SEQ.

Note, in particular, the extended range of **Op**erators and the different meaning associated with the operator 'M'. The operators '=' and 'X' are such that any 'M' is either an '=' or and 'X' but never both. Which leaves one pondering when one might use 'M' in preference to either an '=' or an 'X'?

The Wikipedia article shows how this complication might be overcome.

There may or may not be some justification for calling the format CIGAR, but if there is, I have no idea what it might be.

Where is the  $3^{rd}$  substitution in the mRNA? Where is the  $3^{rd}$  substitution in the Genomic Sequence?

**spline** makes one work quite hard to answer this one! Unless I am missing something.

From the alignment of Exon 10, the exon including the 3rd Replacement, with a bit of squinting, it can be confirmed that the 3<sup>rd</sup> Replacement is at:



Base pair position 1,312 of the mRNA

Base pair position 25,042 of the genomic sequence

It might also have been relevant to ask which amino acid position corresponded to the Replacement. To discover this one would need to look at the alignment of **Exon 3**, where the coding begins.

More squinting, and I conclude the A of the ATG representing the initial Methionine of the protein coding region is at position 363. That is, the 5' UTR ends at position 362. So the Replacement is at:

Base position 1312 - 362 = 950 of the protein coding region of the mRNA.

As 950 / 3 is 316 remainder 2, the Replacement is at codon position 2 of the 317<sup>th</sup> amino acid of the protein.

Cannot help thinking that **spline** might have helped a bit more here?

I also reflect that I cannot fully recall why I wanted to know where the mutation was, especially given we have decided to reject any chance that it might be a mutation of consequence. Oh well, some things a man must do, just because they are there to be done!!

Time to move on ... without checking my arithmetic. Bound to be right, I used to be a mathematics teacher you know! Several lifetimes ago.

#### **Postscript:**

After the passage of many months, I now recall why I obsessed as to the position of this amino acid substitution. I wondered if it was in the region of one of the major domains of this protein. If it was, it might increase its chances of being significant?

Well, it is not. In the last exercise, we discovered that:

The Paired-box domain is between positions 4 and 128 (Consensus isoform) or 4 and 142 (isoform 5a).

The Homeo-box domain is between 214 and 266 (Consensus isoform) or 228 and 280 (isoform 5a).

So the **Substitution**, at position **317**, is in a relatively neutral region and so, maybe, less likely to be of great consequence?

Compare the predicted **splign** intron/exon boundaries with the conservation suggested by the logo?

What deviation(s) from the model suggested by the logo can you see?

You may have gathered, I rather like this logo, although I rather think it is leading me to make the same point a trifle to often?

The logo is in almost 100% agreement with the predictions of spline.

As you will have noted previously, when looking at the **Ensembl** predictions of exons locations of a similar transcript of the **PAX6** human gene (previous Practical), there is a single exception.

Type

<L-Gap>

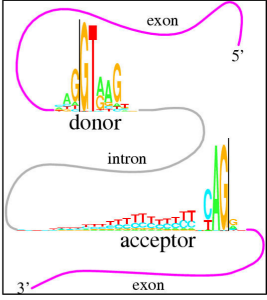
CA<exon>GT

AG<exon>GC

AG<exon>GT

The easiest way to show this in the **spline** output is to look at the **spline** text output again.

The **Type** column records the type of all the **<exon>** alignments it predicts. It also records **2 flanking intron base pairs**.



It is clear that the only time the spline prediction deviates from the model suggested by the logo is at the end of the  $2^{nd}$  exon. Here there is GC rather then GT. Well, nothing is perfect!

From your investigations of Local Alignment:

Why do you suppose your aligned exons are not presented in the correct positional order?

To **Matcher**, the logical order in which to present the alignments is that governed by quality rather than position. So, the highest scoring alignment, rather than the first exon alignment, will be at the top of the list. I think this is generally logical. Once again, the program **splign**, knowing it was looking for an ordered set of exons, was more specifically logical.

DPJ - 2016.12.06