

Earlham Institute summer school on bioinformatics

25-29 July 2016

Basic Bioinformatics Sessions

Practical 4: Multiple Sequence Alignment

Thursday 7 July 2016

Multiple Sequence Alignment

Here we will look at some software tools to align some protein sequences. Before we can do that, we need some sequences to align. I propose we try all the human **homeobox** domains from the well annotated section of **UniprotKB**. Getting the sequences is a trifle clumsy, so concentrate now! There used to be a much easier way, but that was made redundant by foolish people intent on making the future ever more tricky!!

So, begin by going to the home of **Uniprot**:

http://www.uniprot.org/

Choose the Advanced option of the Search button.

First specify that you are only interested in **Human** proteins. To do this, set the first field to **Organism [OS]** and

Set the second field selector to **Reviewed** and

Term to Human [9606].

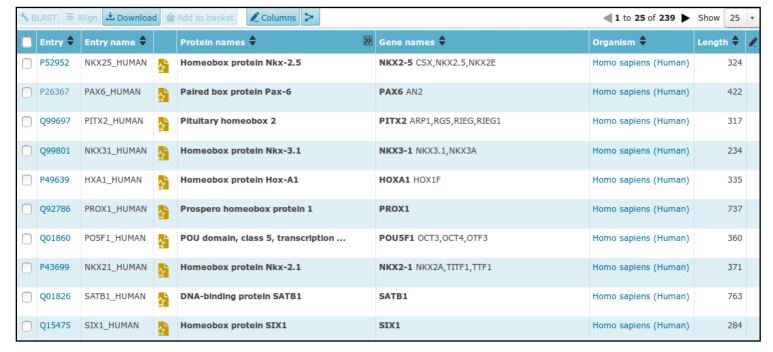
the corresponding Term to
Yes (that is, choose to find
only SwissProt entries).

Click on the button to request a further field selection option. Set the new field to Function. Set

the type of Function to DNA binding. Set the Term selection to Homeobox.

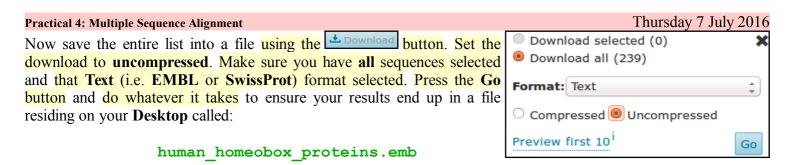
From previous investigations, you should be aware that a **Homeobox** domain is **generally 60** amino acids in length. To avoid partial and/or really weird **Homeobox** proteins, set the **Length** range settings to recognise only **homeobox**s between **50** and **70** amino acids long.

Leave the Evidence box as Any assertion method, one does not wish to be too fussy! Address the uthority to get the search going.



A fine miscellany of sequences will assemble upon you screen. Most seem to declare themselves in possession of a **Homeobox** or two (including **PAX6 HUMAN**), so I suggest a declaration of success.

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```
P52952; A8K3K0; B4DNB6; E9PBU6;
     01-OCT-1996, integrated into UniProtKB/Swiss-Prot.
     01-OCT-1996, sequence version 1.
    06-JUL-2016, entry version 173.
RecName: Full=Homeobox protein Nkx-2.5;
     AltName: Full=Cardiac-specific homeobox;
     AltName: Full=Homeobox protein CSX;
     AltName: Full=Homeobox protein NK-2 homolog E;
GN
OS
OC
OC
OX
RN
RP
     Name=NKX2-5; Synonyms=CSX, NKX2.5, NKX2E;
     Homo sapiens (Human).
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
     Mammalia; Eutheria; Euarchontoglires; Primates; Haplorrhini;
     Catarrhini; Hominidae; Homo
     NCBI TaxID=9606;
     [1]
     NUCLEOTIDE SEQUENCE [MRNA] (ISOFORM 1).
     TISSUE=Heart;
     PubMed=8900537
     Turbay D., Wechsler S.B., Blanchard K.M., Izumo S.;
     "Molecular cloning, chromosomal mapping, and characterization of the
     human cardiac-specific homeobox gene hCsx.";
     Mol. Med. 2:86-96(1996).
```

Take a swift look at the file you have just created. Your neat list of **Human Homeobox** sequences will have transformed into a flood of **many SwissProt** format **UniProtKB** entries. Ugly, but what is required.

Search (Control F) for the term DNA BIND.

It should occur many times (at least once per sequence) in the Feature Tables and most often refer to a **Homeobox** region.

In the **DNA_BIND** Feature Table entries, the position of the **Homeobox**es are recorded and will be used by the next program to isolate the sequence of the **Homeobox**es.

	FT	CHAIN	1	374	Pre-B-cell leukemia transcription factor
	FT				4.
	FT				/FTId=PRO 0000049241.
	FT	DNA BIND	210	272	Homeobox; TALE-type.
	FT				{ECO:0000255 PROSITE-ProRule:PRU00108}.
	FT	VARIANT	169	169	V -> I (in dbSNP:rs8108180).
r	FT				/FTId=VAR 059355.
•	FT	VARIANT	177	177	M -> V (in dbSNP:rs8108981).
ı	FT				/FTId=VAR 059356.
	FT	VARIANT	283	283	T -> M (in a colorectal cancer sample;
	FT				somatic mutation; dbSNP:rs376647012).
	FT				{ECO:0000269 PubMed:16959974}.
	FT				/FTId=VAR 036439.
بد	FT	CONFLICT	368	368	I -> T (in Ref. 1; BAG53471).
	FT				{ECO:0000305}.
l	sQ	SEQUENCE	374 AA;	40854 MW	; B9CE8BE93D0B7ABC CRC64;
		MAAPPRPAPS	PPAPRRLI	TS DVLQQI	MAIT DQSLDEAQAR KHALNCHRMK PALFSVLCEI
-		KEKTVVSIRG	IQDEDPPI	DAQ LLRLDNI	MLLA EGVCRPEKRG RGGAVARAGT ATPGGCPNDN
		SIEHSDYRAK	LSQIRQI	HS ELEKYE	QACR EFTTHVTNLL QEQSRMRPVS PKEIERMVGA
		IHGKFSAIQM	QLKQSTCE	EAV MTLRSRI	MADA RRKRRNFSKQ ATEVLNEYFY SHLNNPYPSE
		EAKEELARKG	GLTISQVS	NW FGNKRI	RYKK NMGKFQEEAT IYTGKTAVDT TEVGVPGNHA
		SCLSTPSSGS	SGPFPLPS	SAG DAFLTLE	RTLA SLQPPPGGGC LQSQAQGSWQ GATPQPATAS
		PAGDPGSINS	STSN		
	//				

Now to extract from the whole protein sequences you have saved in a file, the sequences of just the **Homeobox** domains. One way of doing this (possibly not the best), is to use an **EMBOSS** package program called **extractfeat**. This can be found in many places, including the Bioinformatics server at **Wageningen** in the Netherlands. Go to:

http://emboss.bioinformatics.nl/

aligncopy
aligncopypair
biosed
codcopy
cutseq
degapseq
descseq
entret
extractalign
extractfeat

Find the program extractfeat (in the EDIT section), and set it going.

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Tractical in Manapic Sequence Inflament	Input section
Use the Choose File button to upload the SwissProt format sequences from UniProtKB that you saved in the file human_homeobox_proteins.emb.	Select an input sequence. Use one of the following three fields: 1. To access a sequence from a database, enter the USA here: 2. To upload a sequence from your local computer, select it here: Browse human_homeobox_proteins.emb
Set Type of feature to extract field to DNA_BIND (Make sure you remove the "*").	3. To enter the sequence data manually, type here: Additional section Amount of sequence before feature to extract Amount of sequence after feature to extract Source of feature to display * Type of feature to extract DNA_BIND
Set Value of feature tags to extract to Homeobox* (Make sure you append the "*" to ensure hits with, for example "homeoboxes").	Sense of feature to extract (default is 0 - any sense, 1 - forward sense, -1 - reverse sense) Minimum score of feature to extract Maximum score of feature to extract Tag of feature to extract Value of feature tags to extract Homeobox*
Set the Output sequence format to SwissProt (Fasta would do, but SwissProt retains more annotation).	Output section Output introns etc. as one sequence? No Append type of feature to output sequence name? No Feature tag names to add to the description
Click on the Run extractfeat button to start extractfeat going. Many sequences of 60 amino acids (or so) in length will leap into view.	Output sequence format SwissProt Run section Email address: If you are submitting a long job and would like to be informed by email when it finishes, enter your email address here. Run extractfeat Reset
OUTPUT FILE outseq ID NKX25 HUMAN 138 197 Reviewed; 60 AA. DE [DNA contact] Homeobox protein Nkx-2.5 (Cardiac-specific homeobox) (Homeobox protein Nsx-2.5 (Cardiac-specific homeobox) (Homeobox protein Nsx-2.5 (Cardiac-specific homeobox) (Homeobox protein Nsy-2.5 (Lardiac-specific homeobox) (Homeobox protein Nsy-2.5 (Lardiac-specific homeobox) (Homeobox protein Nsx-2.5 (Lardiac-specific homeobox) (Homeobox protein Nsx-2.5 (Lardiac-specific homeobox) (Homeobox protein Nsy-2.5 (Lardiac-specific homeobox) (Homeobox protein Protein) (Oculorhombin) (Homeobox protein Protein Homeobox) (Lardiac-specific homeobox) (Homeobox protein Protein Homeobox) (Homeobox) (Homeobox	CSX) (Homeobox protein NK-2 homolog E) ITX2) (Paired-like homeodomain transcription factor 2) (RIEG bicoid-related homeobox transcription factor) (Solurshin)

Right click the button and select Save Link as... Do whatever it takes to save all your Homeobox domains into a file residing on your Desktop called:

 ${\tt homeobox_human.emb}$

HXA1_HUMAN_229_288 Reviewed; 60 AA.

[DNA_contact] Homeobox protein Hox-A1 (Homeobox protein Hox-1F)

SEQUENCE 60 AA; 7365 MW; 53E2BC59B06F544E CRC64;
PNAVRTNFTT KQLTELEKEF HFNKYLTRAR RVEIAASLQL NETQVKIWFQ NRRMKQKKRE

Finally, we have some sequences with which to investigate the multiple sequence alignment programs.

Take a look at the file you have created. You should have many human homeobox domains in SwissProt format, looking rather as they did in your browser window. Happily ClustalX, the first multiple alignment program to be investigated, accepts multiple sequence SwissProt format files as input.

ClustalX is a part of the mostly widely known family of Multiple Sequence Alignments (MSA) programs, originating in the 1980s. Until relatively recently, it was the only real option. ClustalX still has merit, although it lacks some of the sophistication of more recent programs. ClustalX runs on effectively all workstations and has a

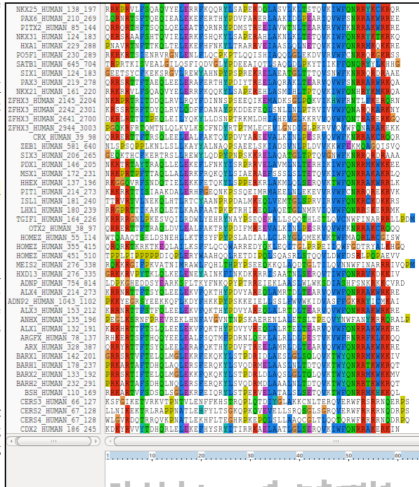
nice Graphical User Interface (GUI). A good place for us to start. It is installed on your workstations.

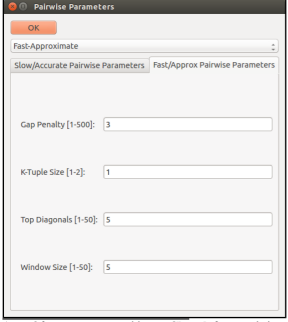
Start up the program ClustalX¹. The ClustalX Graphical User Interface (GUI) will regally mount your screen.

Select Load Sequences from the File pull down menu and load your file of homeobox domains.

The sequences will arrange themselves colourfully. Many of the **homeoboxes** are similar enough to look convincing even before alignment. Note the "Manhattan skyline" under the sequences indicating the varying degrees of conservation.

You might like to increase the **Font** size from the minute default setting designed for Hawks and Eagles, to something more comfortable. **24** works tolerably well for me.





From the Alignment pull down menu, go to the Alignment parameters menu and select Pairwise Alignment Parameters. Just for a moment, change the setting from Slow-Accurate to Fast-Approximate. Bring the corresponding parameters into view by clicking on Fast/Approx Pairwise Parameters tab².

Hopefully, we will have discussed the way **ClustalX** (and similar multiple alignment tools) work. Intuitively, it should not make a lot of difference how the initial pairwise comparison stage is conducted. However, it very often does.

Specifically for this set of proteins, as well as generally, **ClustalX** will give a noticeably better alignment if the initial pairwise alignment stage is done carefully. Accordingly, reverse your whimsical setting change by moving back from **Fast-Approximate** to **Slow-Accurate**.

Of course, you could run Clustal from websites all over the world if you wished. Specifically, it is available both at the EBI and the Bioinformatics server at Wageningen. Try it if you have time. You get the same results but will, sadly, lose the pretty interface.

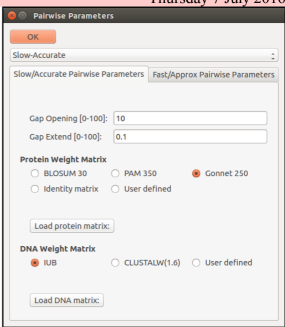
http://www.ebi.ac.uk/Tools/clustalw2/index.html

http://www.bioinformatics.nl/tools/clustalw.html

2 The **Fast-Approximate** algorithm is essential that which the database searching program **fasta** employs. Assuming we have discussed how **fasta** works, it should require no further explanation here.

Click on the Slow/Accurate Pairwise Parameters tab for a final look at the default parameters to be used. The Slow-Accurate option is essentially a version of Global Alignment algorithm we will have discussed previously. Hopefully, all the parameter options will therefore be familiar to you.

I will assume both sets of parameters at least seem familiar? If not please ask. The default **Slow/Accurate Pairwise Parameters** you now have in view are fine. Click the **OK** button to dismiss the **Pairwise Parameters** window.



Before proceeding, save the **homeobox** sequences in **FASTA** format, which will better suit the other **MSA** programs we will try. Do this by selecting **Save sequences as...** from the **File** pull down menu. **Deselect CLUSTAL format**, select **FASTA format**. Click **OK**. A file called **homeobox human.fasta** will be created. Take a look to check it is as you would expect.

Format									
	CLUSTAL format								
	GCG/MSF format								
	GDE format								
\checkmark	FASTA format								

Output Files	
CLUSTAL format	☐ NBRF/PIR format
☐ GCG/MSF format	☐ PHYLIP format
☐ GDE format	☐ NEXUS format
☐ FASTA format	

Strangely, saving your sequences in **FASTA** format convinces **clustalx** that it should now output its alignments in **FASTA** format. To prevent this, select **Output Format Options** from the **Alignments** pull down menu. Deselect **FASTA** format and select **CLUSTAL** format. Click **OK**.

From the **Alignment** pull down menu, select **Do Complete Alignment**. Accept the default names for output files and click on the **OK** button. **ClustalX** will start to think deeply and eventually come up with it view of how the **homeobox** domains should be aligned.

Not a bad first try. From an entirely non scientific, cosmetic viewpoint, the ragged ends offend a trifle, as does the gap just before position 30!

SATB1 HUMAN 645 704	TRP	RTK	ISVEALGI	L <mark>o</mark> sf	I Q D	/GLY	PDEEAI	QTL	SAQLI	LPKYT	IIKFF	QNQ R Y	YLKHHG	
SATB2 HUMAN 615 674	PRS		ISLEALGI			/GLY					IIKFF			
ZFHX3 HUMAN 2145 2204	NKRP	RTR	ITDDOLRV	RQY	F-DI	N N S	PSEEQI	KEM	ADKS	LPOKV	IKHWE	RNTLE	KERORN	
ZFHX4 HUMAN 2084 2143	FKRP	RTR	ITDDOLKI	RAY	F-D	N N S	PSEEQI	QEM	AEKS(LSOKV	IKHWE	RNTLE	KERORN	
			IENRVR G N									CNRRQ	KGKRSS	;
			IENRVR G N									CNRRQ		;
PO5F2 HUMAN 210 269	GKWR		RERRIGNS									YNRSK	MGSRPT	
	RRKK		IETNVRFA									CNRRQ		
			IETNIRVA											
			IETNIR <u>L</u> T									CNRRQ		I – – –
			I EVSVK G A											
PO3F3 HUMAN 406 465	KRKK	RTS	IEVSVK G A	LESH	FLK-	C <mark>P</mark> K	PS AQEI	TNL	ADS L	LEKEV	VRVWE	CNRRQ	KEKRMT	
PO3F1 HUMAN 339 398	KRKK	RTS	IEV G VK G A	LESH	FLK-	C <mark>P</mark> K	PSAHEI	TGL	ADS L	LEKEV	VRVWE			!
PO3F4 HUMAN 278 337	KRKK	RTS	IEVSVK G V	LETH	FLK-	C <mark>P</mark> K	PAAQEI	SSL	ADS L	LEKEV	VRVWE	CNRRQ		
			I <mark>S</mark> IAAKDA											
	KKRK		IAA <mark>PE</mark> KRS										K Q K R M K	
			IAA <mark>PE</mark> KRS											[
			IAA <mark>PE</mark> KRS										A	(F
			FTPQALEI									CNKRQ	A LK NTI	
			ETPQAIEA									CNRRQ		j
			FTVE <mark>O</mark> QRI										KMSSKN	
	ALQD		FSDR <u>D</u> LAT									GNRRR		
			FTQEQIEA											
			FSPSQAEA											
			FSAE <mark>Q</mark> LQL										K S RRQS	
			FSPVQLEQ									ONRRA		
			FSEE <mark>Q</mark> LQA											(
			FTDE <mark>Q</mark> LEA										KwrrQK	
			FTSQ <mark>Q</mark> LQE											
PITX3_HUMAN_62_121			FTSQ <mark>Q</mark> LQE										Kwrkre	
			FTSQ <mark>Q</mark> LQE											
			FTRAQLDV											
			FTRSQLDV									KNRRA		\
			FTRSQLEE											
			FTAQQLEA										KFRKKQ	
			FTAEQLEE										R <mark>w</mark> rkqa	
			FTAEQLEE											
			FTSYQLEE											
			FSTFQLEE.										KWRKRE	
ALX1_HUMAN_132_191 ARX_HUMAN_328_387			FTSL <mark>O</mark> LEE FTSY O LEE											
													AARKRE	2
PHX2A_HUMAN_90_149	0 KK I	T 1	FTSA <mark>Q</mark> LKE	K V	AE-	ТН	PDITTE	E E L	TVT) LEEAK	AOAME	ŲΝKK <mark>A</mark>	NE KRQE	
· () >	(-(111				
	1	10		20		30		40		50		60		70
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				h- 1						No. of			1 -4	

In reality, these features might be very interesting, but here I go for pretty!

So, just to investigate what is possible, select all the homeobox sequences that are causing the gap around position 30 by clicking on their names (quite a lot of them I fear). Hold the Ctrl key down to allow multiple selection.

Once you have them all, go to the Edit pull down menu and select Cut Sequences. Then select Remove Gap-Only columns from the Edit pull down menu. Nasty gap gone ... along with all scientific credibility, but ... never mind.

You could recompute the alignment scratch with the reduced sequence set, but you should end up with the same answer, of course. Just for the sake of it, select Select All Sequences from the Edit pull down menu. Then select Remove All gaps from the Edit pull down menu and confirm your intentions. You are now back where you started, but without the sequences that mess up the alignment intolerably!

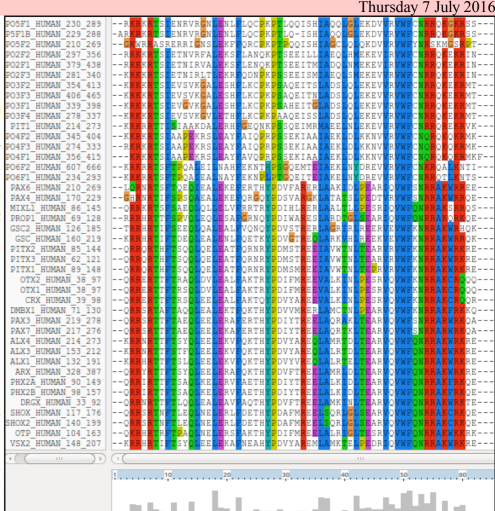
Save your filtered set of sequences in a file. From the File pull down menu select Save Sequences as... Choose **FASTA** format only.

The default file name is OK, even though it will overwrite the original sequences. I am convinced sequences eliminated would not be aligned convincingly with any of the tools we have at hand. Let us lose them! Press the **OK** button.

From the **Alignment** pull down menu, select Output Format Options and select CLUSTAL format only. Again, from the Alignment pull down menu, select **Do** Complete Alignment. Accept the default names for the output files. This will overwrite your previous efforts, but no matter. More deep thought. Well, I got back to where I was, no gaps around position 30 but still with ragged ends!

It is difficult to prove you have exactly

the same alignment as previously as the order of the MSA will be different. This order being determined by the pairwise comparison stage of the clustalx MSA computation.



You will recall from earlier that the **Prosite** pattern for a **homeobox** is the ever memorable:

This corresponds to positions 36 to 59 in my alignment. See that the "Manhatten Skyline" is encouraging in the parts of this region that matter.

Note that the profile **Tryptophan**, in position **50**, is **very COLOR** Consistent, but not quite **100%** as suggested by the **Prosite** pattern³. The **W** was even conserved in the sequences that were cosmetically removed.

Position 52 is not conserved ("-x-") according to the Prosite pattern. In the alignment segment offered here, it looks like a pretty consistent Q. However, the "Manhattan skyline" at this position is very low, suggesting that the sequences in view might not be typical of the whole alignment set. Which, upon checking they are not!

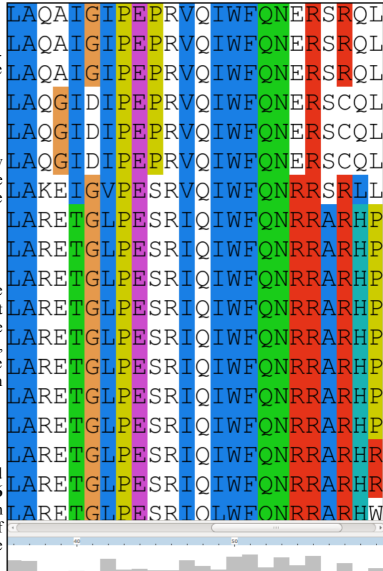
Looking through this alignment, I get the feeling I could design a better, stricter pattern for the region between 36 and 59. Possibly true, but remember the pattern in Prosite aims to represent the conservation of Homeobox domains in ALL organisms. Here we have only sequences from Human.

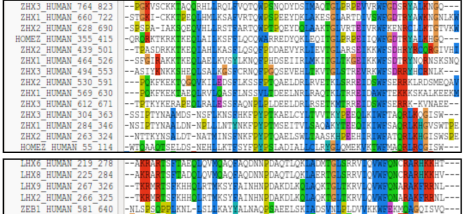
ZEB2 HUMAN 644 703

ZHX3 HUMAN 835 894

ZHX1 HUMAN 777 832

HOMEZ_HUMAN_451_510 NANGN_HUMAN_102_161





AMNME<mark>P</mark>NSDELLK

YEEDLON

1 10 20 30 40 50 60

MTHK

Of course, things are not quite so convincing throughout. If you look at the top and bottom few sequences, you will see that **ClustalX** had its moments of uncertainty.

Note, however, the consistent **W** in position **50** despite the surrounding crumble.

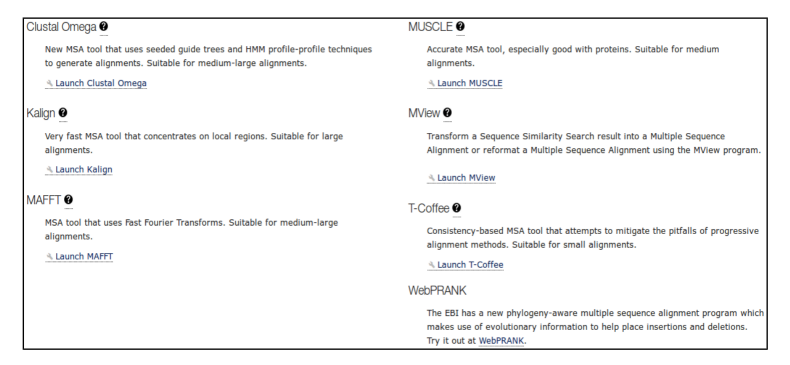
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From the "Manhattan Skyline", you can see the conservation is less than 100%. Less conserved than the F that immediately follows in fact? Look at your alignment, the "Manhattan Skyline" does not seem to reflect reality? The W is very well conserved, although the scoring matrices would regard any deviation from W as serious? I need to find out more about how the Skyline is computed.

Now to show existence of some **msa** program options available on the web. There are many. They are available from a number of server sites. An obvious place to start has to be the **EBI** page dedicated to **MSA**. Go to:

http://www.ebi.ac.uk/Tools/msa/

Offered here is a selection of popular, current generation **MSA** tools. Each is accompanied by advice to guide the choice of tool to best fit the circumstances. Each tool is provided with a link to its **Launch** interface. All the **Launch** interfaces are very consistent. Once you have run one of the **MSA** options, you should have no trouble running any of the others.



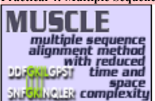
Here I intend to align again the human **homeboxes** with just one of the tools on offer. Then take a quick look at how the machine generated multiple alignment can be manually edited using **Jalview**, a program that is installed on your workstation and that you have already used as an alignment viewer when investigating **Pfam** and also **Jpred**.

Then I will invite you to try a few of the other options for yourself and see that they do not all produce the same alignment! Differences reflect not only the parameters selected, which we will have discussed, but also the particular objectives of the program selected. For example, a multiple protein sequence alignment optimal for investigating conservation of protein structure might well not be identical to one best representing protein evolution.

Used to align the **Homeobox** sequences used in this exercise, I do not expect you will see much difference between the outputs of any of these options. They will all work sufficiently on such a simple data set.

The program whose use I choose to describe carefully, leading on to a short **Jalview** exercise is **MUSCLE**. I choose thus as **MUSCLE** is now the first choice of most of the people with whom I work. Also popular are **Clustal Omega**, **MAFFT** and, for **phylogeny**, **WebPRANK**.

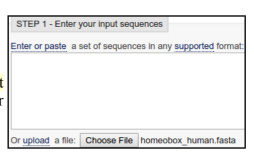
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So the plan now is to use MUSCLE⁴ to align again the homeobox sequences previously aligned with clustalX. MUSCLE works in a way similar to clustalX but it takes rather more care in the generation of the Guide Tree used to control the order of pairwise construction of the final multiple alignment⁵. Particularly for more difficult alignments, MUSCLE should do a better job than clustalX. The alignment you will generate here will certainly be different. I leave you to judge for yourselves whether it is better.

Start by requesting to Launch MUSCLE

Use the **Choose File** button to upload the file containing the **FASTA** format **homeobox** sequences, **homeobox_human.fasta**. This file should no longer included the sequences with a mess around position **30**.



STEP 2 - Set your Parameters

OUTPUT FORMAT: ClustalW

The default settings will fulfill the needs of most users and, for that reason, are not visible.

More options... (Click here, if you want to view or change the default settings.)

Take a look at the **Set your Parameters** section of the page. I find the claim that "The default settings will fulfill the needs of most users and, for that reason, are not visible" a little strange? What about the users who are not in the category "most"? I want all the control

over the programs I use that their creators deemed it sensible to make available6?

The default settings behind the More options... button are not those that affect the computation of the MSA. I confess myself confused at the lack of any meaningful options to consider? I was expecting at least the gap open and gap extension penalty options (available elsewhere, including Wageningen), plus a way to change the scoring matrix. I have inquired why things are as they are



(most recently **2016.04.17**). No practical issue here, as I intended to suggest the defaults whatever they were. Look at the range of settings for the **OUTPUT TREE** parameter. **none** is indeed the thinking persons choice, but ... one or the other (but not both?) of the **Guide Trees** that **MUSCLE** will compute can be saved if you wish⁷. You may also set the **OUTPUT ORDER** to **aligned** or ... **aligned**?



There are a number of **OUTPUT FORMATS** offered. For a quick glance at your results, both **ClustalW** or **HTML** are fine. Here I suggest it would be nice to generate an output that can be downloaded and viewed in **Jalview**⁸. The default **ClustalW** or **Pearson/FASTA** serve for this purpose. As **ClustalW** looks more like an alignment in the web page, I choose **ClustalW**⁹.

How do the options for the **OUTPUT TREE** relate to the output files of **ClustalX** and the difference between the way that **ClustalX** and **muscle** work?

Comment on how one might choose between the range of options offered for the aligned parameter?

⁴ Available from a variety of websites in addition to the **EBI**, including the Bioinformatics server at **Wageningen**: http://www.bioinformatics.nl/tools/muscle.html

⁵ As discussed, superficially at least, previously. I hope.

I have asked the **EBI** about their policy (the same for all the locally provided **MSA** options). Discussion is ongoing (2016.04.20).

A useful option if you thought it possible you might want to rerun **MUSCLE** with different parameter setting for the stages after the **Guide**Tree(s) are generated. The same possibilities exist for **ClustalX**. Of course, utterly pointless if it is impossible to control the relevant parameters
.... so I really cannot see the point of any of the **More options** section? I am open to elucidation from all/any sources.

The java alignment editor and viewer you used to look at the **Pfam** and **Jpred** alignments earlier.

But feel free to try the others. **HTML** is the default at **Wageningen**. The **Phylip** formats are the best if you are going to analyse your output further with the phylogeny programs of the **PHYLIP** package.

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After considering these enigmas, or before if you prefer, Click on the sit back to admire muscle in action.	LHX2 HUMAN 266 325 LHX6 HUMAN 219 278 LHX8 HUMAN 225 284 ZFHX3 HUMAN 2641 2700 ZFHX4 HUMAN 2560 2619 ZFHX2 HUMAN 2560 2119 ZFHX2 HUMAN 265 2124 ZFHX2 HUMAN 2944 3003 ZFHX4 HUMAN 2884 2943 LMX1A HUMAN 195 254	QRRYR-TTFTSYQLEELERAFQKTHYPDVFTREELAMRLDLTÉARVQVWFQNRRAKWRKRRIHR-TTFTSLQLEELEKVFQKTHYPDVYVREQLALRTELTEARVQVWFQNRRAKWRKRRNR-TTFTSYQLEELEKVFQKTHYPDVYAREQLALRTELTEARVQVWFQNRRAKWRKRRNR-TTFSTFQLEELEKVFQKTHYPDVYAREQLALRTDLTEARVQVWFQNRRAKWRKRRNR-TTFSTFQLEELEKVFQKTHYPDVYAREQLALRTDLTEARVQVWFQNRRAKWRTTRVR-TVLNEKQLHTLRTCYAANPRPDALMKEQLVEMTGLSPRVIRVWFQNKRCKDKTKRMR-TSFKHHQLRTMKSYFAINHNPDAKDLKQLAQKTGLTKRVLQVWFQNARAKFRTKRMR-TSFKHHQLRTMKSYFAINHNPDAKDLKQLAQKTGLTKRVLQVWFQNARAKFRAKRAR-TSFTAEQLQVMQAQFAQDNNPDAQTLQKLADMTGLSRRVIQVWFQNCRARHKAKRAR-TSFTAEQLQVMQAQFAQDNNPDAQTLQKLABMTGLSRRVIQVWFQNCRARHKDKRLR-TTITPEQLEILYGYLLDSNPTRKMLDHIAREVGLKKRVVQVWFQNTRARERDKRLR-TTITPEQLEILYRWYMQDSNPTRKMLDHIAREVGLKKRVVQVWFQNTRARERDKRLR-TTILPEQLEILYRWYMQDSNPTRKMLDCISEEVGLKKRVVQVWFQNTRARERQRRYR-TQMSSLQLKIMKACYEAVRTPTMQECEVLGEEIGLPKRVIQVWFQNARAKEK PGQRKYR-TQMSSLQLKIMKACYEAVRTPTMQECEVLGEEIGLPKRVVIQVWFQNARAKEKHKRFR-TQMSNLQLKVLKSCFNDYRTPTMLECEVLIGNDIGLPKRVVQVWFQNARAKEKHKRFR-TQMSNLQLKVLKACFSDVRTPTMQECEMLGNEIGLPKRVVQVWFQNARAKEKPKRRP-TILTTQQRRAFKASFEVSSKPCRKVRETLAAETGLSVRVVQVWFQNQRAKMK
superficially at least, similar to that offered by ClustalX .	LMX1B_HUMAN_219_278 LHX1 HUMAN_180_239 LHX5_HUMAN_180_239 LHX3 HUMAN_157_216 LHX4_HUMAN_157_216 HOMEZ_HUMAN_451_510 ZHX3 HUMAN 835_894	PKRPR-TILTTQQRRAFKASFEVSSKPCRKVRETLAAETGLSVRVVQVWFQNQRAKMKRRGPR-TTIKAKQLETLKAAFAATPKPTRHIREQLAQETGLNMRVIQVWFONRRSKERRRGPR-TTIKAKQLETLKAAFAATPKPTRHIREQLAQETGLNMRVIQVWFQNRRSKERAKRPR-TTITAKQLETLKSAYNTSPKPARHVREQLSSETGLDMRVVQVWFQNRRAKEKAKRPR-TTITAKQLETLKNAYKNSPKPARHVREQLSSETGLDMRVVQVWFQNRRAKEK -: : : : : : : : : : : : : : : : : : :
The alignment is irritatingly split into two	ZHX1_HUMAN_777_832	LGIELF ISW
sections. A nice extra parameter might have been "How wide would you like your alignment to be"? A problem with the format rather than the program, to be fair.	ZHX3_HUMAN_304_363 ZHX1_HUMAN_284_346	ISWSPE ISW VSWTPE SNS SVQ KEM LKEEKM SMEQAV AEE RGIVHI NLK NSKSNQ HGQ

At the very bottom of the page, **muscle** whines:

PLEASE NOTE: Showing colors on large alignments is slow.

```
So click the Show Colors button at the top of the page and try to live with the pain of such gross Trans-Atlantic inept spelling in a European site!!! Good Grief! They get everywhere!!
```

Well, an improvement I suppose? Colours are very useful (even slow ones) in the interpretation of alignments. Various colour schemes are used to clarify the message of alignments. Colouring can indicate shared amino acid properties not immediately evident when the letter representations differ.

```
ARX HUMAN 328 387
                              -- QRRYR-TTFTSYQLEELERAFQKTHYPDVFTREELAMRLDLTEARVQVWFQN
ALXĪ HUMAN 132 191
                              -- KRRHR-TTFTSLOLEELEKVFOKTHYPDVYVREOLALRTELTEARVOVWFONE
                              --KRRNR-TTFTSYQLEELEKVFQKTHYPDVYAREQLAMRTDLTEARVQVWFQNRRAKW
ALX4 HUMAN 214 273
                              -- KRRNR-TTFSTFQLEELEKVFQKTHYPDVYAREQLALRTDLTEARVQVWFQNRRAKW
ALX3 HUMAN 153 212
ISL1 HUMAN 181 240
                              --TTRVR-TVLNEKQLHTLRTCYAANPRPDALMKEQLVEMTGLSPRVIRVWFQNK
ISL2 HUMAN 191 250
                              --TTRVR-TVLNEKQLHTLRTCYAANPRPDALMKEQLVEMTGLSPRVIRVWFQNKRCKDI
LHX9 HUMAN 267 326
                              --TKRMR-TSFKHHOLRTMKSYFAINHNPDAKDLKQLAQKTGLTKRVLQVWFQNARAKF
LHX2 HUMAN 266 325
                              --TKRMR-TSFKHHQLRTMKSYFAINHNPDAKDLKQLAQKTGLTKRVLQVWFQNARAKF
LHX6 HUMAN 219 278
                              --AKRAR-TSFTAEQLQVMQAQFAQDNNPDAQTLQKLADMTGLSRRVIQVWFQNCRARH
LHX8_HUMAN_225
                              --AKRAR-TSFTADQLQVMQAQFAQDNNPDAQTLQKLAERTGLSR
                                                                                 RVIQVWFQNCRARH
                284
ZFHX3_HUMAN_2641_2700
ZFHX4_HUMAN_2560_2619
                              -- DKRLR-TTITPEOLEILYOKYLLDSNPTRKMLDHIAHEVGLKKRVVOVWFONTRARE
                              -- DKRLR-TTITPEQLEILYEKYLLDSNPTRKMLDHIAREVGL
                                                                                 RVVOVWFONTRARE
                              -- DKRLR-TTILPEQLEILYRWYMQDSNPTRKMLDCISEEVGL
                              -- QRRYR-TQMSSLQLKIMKACYEAYRTPTMQECEVLGEEIGLP
                                                                                (RVIOVWFONARAKE
7FHX3 HUMAN 2944 3003
                              PGOKRER-TOMTNI OLKVI KSCENDYRTPTMI ECEVI GNDTGI PKRVVOVWEONARAKE
ZFHX4 HUMAN 2884 2943
                              -- HKRFR-TQMSNLQLKVLKACFSDYRTPTMQECEMLGNEIGLPKRVVQVWFQNARAKE
LMX1A_HUMAN_195_254
                              --PKRPR-TILTTQQRRAFKASFEVSSKPCRKVRETLAAETGLSVRVVQVWFQNQRAKM
                              --PKRPR-TILTTQQRRAFKASFEVSSKPCRKVRETLAAETGLSVRVVQVWFQNQRAKM
--RRGPR-TTIKAKQLETLKAAFAATPKPTRHIREQLAQETGLNMRVIQVWFQNRRSKEI
LMX1B HUMAN 219 278
LHX1_HUMAN_180_239
LHX5_HUMAN_180_239
                              -- RRGPR-TTIKAKQLETLKAAFAATPKPTRHIREQLAQETGLNMRVIQVWFQNRRSKE
 HX3 HUMAN 157
                              -- AKRPR-TTITAKQLETLKSAYNTSPKPARHVREQLSSETGLDMRVVQVWFQNRRAKE
LHX4_HUMAN_157_216
                              --AKRPR-TTITAKQLETLKNAYKNSPKPARHVREQLSSETGLDMRVVQVWFQNRRAKE
HOMEZ_HUMAN_451 510
                              EVV - - -
                              RΔV---
ZHX1_HUMAN_777_832
                              LGIELF
HOMEZ HUMAN 55 114
                              ISW-
ZHX2_HUMAN_263_324
ZHX3_HUMAN_304_363
                              ISWSPE
ZHX1_HUMAN_284_346
                              VSWTPE
ZEB2_HUMAN_644_703
ZEB1_HUMAN_581_640
                              SNS--
                              SVQ---
NANGN HUMAN 102
ZHX1_HUMAN_569_630
                              LKEEKM
ZHX2 HUMAN 530 591
                              SMEQAV
ZHX3 HUMAN 612 671
                              AEE-
ZHX2 HUMAN 439 501
                              RGIVHI
7HX3 HUMAN 494 553
                              MLK
7HX1 HUMAN 464 526
                              NSKSNO
HOMEZ HUMAN 355 415
```

But any decoration available here is far short of what can be achieved with **Jalview**, so click on the **Download Alignment File button** to save you alignment in a file on your **Desktop** called:

homeobox human muscle.aln

10:43:40 AM

Jalview can be easily installed under all commonly used operating systems and run locally. For these exercises, I attempt to use services available freely from the INTERNET wherever possible, so let us run Jalview from the web here by first going to:

http://www.jalview.org/

and selecting the Launch Jalview Desktop link at the top of the page. Close down all the example outputs

Jalview sees fit to show you on start up. From the File pull down menu choose from File from the Input Alignment option. Locate and load the file homeobox_human_muscle.aln (you must adjust the file name

filter to included .aln files).

The default view is a trifle bland.

HXC12_HUMAN_214_273/1-60

HXD12_HUMAN_202_2617.60

HXD12_HUMAN_202_2617.60

HXD12_HUMAN_202_2617.60

HXD12_HUMAN_102_2617.60

MEOX2_HUMAN_103_2617.60

SSX2_HUMAN_202_2617.60

You could try the default colour scheme used by **ClustalX**, for example.

HAA10_HUMAN_266_325,1-60
HAD10_HUMAN_266_325,1-60
HAD2_HUMAN_185_244,1-60
HAD3_HUMAN_185_251,1-60
HAD3_HUMAN_182_251,1-60
HAD3_HUMAN_283_344,1-60
HAD3_HUMAN_283_344,1-60
HAD3_HUMAN_183_202,1-60

Now the **MUSCLE** and massaged **ClustalX** alignments look even more similar! In the nicely aligned regions at least.

Basic Bioinformatics

HXB13_HUMAN_216_275/1-60 HXD13_HUMAN_276_335/1-60 HXA13_HUMAN_322_381/1-60 HXC13_HUMAN_260_319/1-60 ATNKF AASKF EVX1 HUMAN 183 242/1-60 /X2_HUMAN_188_247/1-60 RLELEKEFHONR IOR DX4 HUMAN 173 232/1-60 CDX2_HUMAN_186_245/1-60 CDX1_HUMAN_154_213/1-60 38X2 HUMAN 247 306/1-60 BX1_HUMAN_261_320/1-60 MX1 HUMAN 241 300/1-60 ITROR I N R Q K LTRLR LTRLR LSRLR AHHN MEOX2_HUMAN_187_246/1-60 GSX2_HUMAN_202_261/1-60 SSNM GSX1_HUMAN_147_206/1-60 HXD1_HUMAN_229_288/1-60 LLELEREFASNM RLR HXAI HUMAN 229 288/1-60 LTELEKEFHFNK RAR RΕ LTELEKEFHFNK IRELEREFFFNV IRELEREFFFSV HXB1_HUMAN_203_262/1-60 HXC11_HUMAN_232_291/1-60 INKEK HXA11 HUMAN 241 300/1-60 INKEK IN. HXC10 HUMAN 268 327/1-60 EKEFLFNM RER RER RER MN LELEKEFLFNM RDR MN LFNM RDF HXD9_HUMAN_285_344/1-60 HXA2_HUMAN_143_202/1-60 HXB2 HUMAN 143 20271-60 PDX1_HUMAN_146_205/1-60 HXD3_HUMAN_194_253/1-60 DQ HXB3 HUMAN 188 247/1-60 DΩ IXA3_HUMAN_191_250/1-60 HXD8 HUMAN 197 256/1-60 HXC8_HUMAN_149_208/1-60 HXB8_HUMAN_146_205/1-60

There are many **Jalview** features that merit investigation. Have a look around if you have time. In particular, **Jalview** will compute simple phylogenetic trees for you employing a number of methods (**Calculate Tree** from the **Calculate** pull down menu). Try it, but be aware this is only sensible if you were very sure of your alignment (and have a few less sequences maybe?).

Jalview is made by the same group as produce JPred. You could send your alignment for Secondary Structure Prediction via the Web Service pull down menu, if you wished.

A very important purpose of **Jalview** is to allow users to edit alignments as well as just to view them. For example, hold down the **Shift** key, click and hold on any amino acid at the edge of a gap, slide left and right and see that you can introduce and/or alter the position of gaps. It is very important to be able to edit alignments generated by even the best of programs. As I hope has been made clear, the alignment algorithms are crude. If you know something about the sequences you are aligning it is very reasonable to suppose you can improve upon the computer's alignments. **Jalview** tries to make this possibility easy. Look through some of the other **Edit** pull down menu options, it does not matter how much you mangle your alignment, you can always make another one.

Finally, take a look at the **Jalview** "**Manhattan Skyline**" for the highly conserved **W** at position **51**. This seems better quality than **clustalX** managed? I am not sure how one can make further comment without knowing what parameters were used. Is there really an improvement? If so, is it due to the improved alogorithm or more appropriate choice of parameters?

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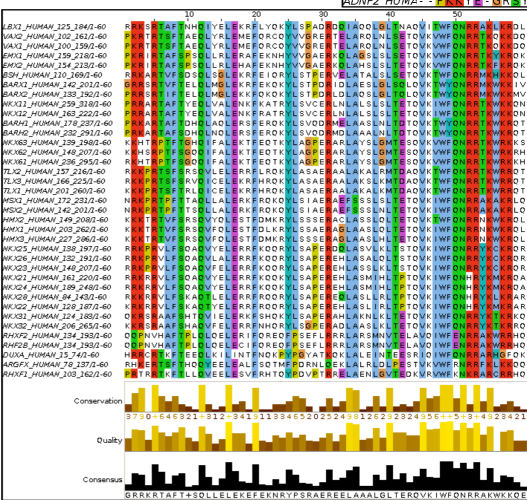
In my alignment, the **W** at position **51** was at position **50**, according to **clustalx**. This slippage to the right is due to **MUSCLE** introducing an extra gap, inspired by just one sequence at position **8**. Is this sensible? No idea ... exactly when it might be good idea to investigate the effect of lighter gap penalties? Not possible, but fear not children!! The sages of the **EBI** assure us that they have selected the correct setting for our sequences that they have never seen??? Sing praises and have faith brothers and sisters for we are in the safe hands of the blessed!!!

*ZHX3 HUMA*N - - SSI<mark>P</mark>T - YN<mark>A</mark>A *ZHX1⁻HUMA*N - - N S I <mark>P</mark> T - Y N <mark>A</mark> A *ZEB1 HUMA*N--NLS<mark>P</mark>S-Q<mark>P</mark> NANGN HUM, - - QYPEK - RLV ZHX1 HUMAN-ZHX2_HUMAN - - P<mark>OK</mark>FI ZHX3 HUMAN - - T P TKY-KFRA ZHX2 HUMAN - - TPASD - RKK ZHX3 HUMAN - - A S I Y ZHX1 HUMAN - - S F G I <mark>F</mark> *HOMEZ HUM,- -* Q<mark>RQ</mark>R<mark>K</mark>TKRK *ZHX2 HUMA*N - - S<mark>P</mark>S<mark>P</mark>A - IAK *ZHX3 HUMA*N - - <mark>PGK</mark>VS - CKK ZHX1¯HUMAN - - ST<mark>G</mark>KI - CKK *ADNP HUMA*NLD<mark>PKGHE</mark> - DD ADNP2 HUMA-

You can also **Select** and **Cut**sequences in a way similar to that you employed with **clustalx**. I could not resist it! I removed all the ugly sequences that caused the gaps at the start and finish of the alignment (just select their names and then select **Cut** or **Delete** from the **Edit** menu). I achieved the gap-free beautiful alignment illustrated.

| KAX1_HUMAN_150_159_7160_160_18M_HUMAN_130_169_7160_18M_HUMAN_130_169_7160_18M_HUMAN_160_717_160_7160_18M_HUMAN_130_717_16

Of course, **Jalview** does not compute alignments, so once I had removed all the unfortunate proteins, I had to use an **Edit** option to tidy up my meddling. I used **Remove Empty Columns** to get rid of the gap columns at the start of the alignment. The gaps at the end just melted away once the sequences that supported their presence were removed.



Science is easy! Once you remove the need for honesty that is.

If it could be done slightly more meaningfully, I would suggest you might try some of the other **MSA** tools offered by the **EBI**, to investigate the differences in the alignments computed. Any differences might be due to different parameter selection or differences in the alignment. Whilst the **EBI** refuse to share with their users how they are running the programs, however, it is hardly worth the effort. Maybe I look for a more obliging service for next time?

PSI-BLAST

This program is used to find a comprehensive set of relatives of a protein. First, **BLAST** is used to find closely related proteins. From an alignment of these proteins a general "profile" (a Position Specific Scoring Matrix -**PSSM**) is computed. A **PSSM** is very similar in concept and purpose to an **HMM** profile in that it summarises significant features present in the sequences it represents.

A query against the protein database is then run using the **PSSM**, and a larger more widely associated group of proteins is found. This larger group is used to construct another **PSSM**, and the process is repeated until no more significantly matching new sequences can be detected, or the user tires of the whole process.

You have used PSI-BLAST integrated into Jpred already and similar ideas were used to create the PFAM alignments. Here we will use PSI-BLAST explicitly at the NCBI on the Paired DOMAIN of the PAX6 protein that you saved in a file earlier. It should be possible to detect a large family of PAX domains and to eventually multiply align them generating something like the Full alignment from the PFAM database viewed earlier¹⁰.

Enter Query Sequence

To investigate **PSI-BLAST** go first to the **NCBI** Home page at:

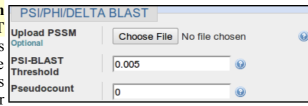
http://www.ncbi.nlm.nih.gov/

Click on the BLAST option. Select protein **BLAST** from the **Basic BLAST** section. Upload the PAX6 paired box domain sequence (stored in the file pax domain.fasta) using the appropriate **Browse** button.

Query subrange @ Enter accession number(s), gi(s), or FASTA sequence(s) (From Or, upload file Choose File pax_domain.fasta Job Title Enter a descriptive title for your BLAST search (Align two or more sequences Choose Search Set Database Non-redundant protein sequences (nr) ▼ @ Organism Exclude + Enter organism common name, binomial, or tax id. Only 20 top taxa will be shown. Exclude ■ Models (XM/XP) ■ Uncultured/environmental sample sequences **Entrez Query** Enter an Entrez query to limit search @ Program Selection Algorithm blastp (protein-protein BLAST) PSI-BLAST (Position-Specific Iterated BLAST) PHI-BLAST (Pattern Hit Initiated BLAST) ○ DELTA-BLAST (Domain Enhanced Lookup Time Accelerated BLAST)

Select PSI-BLAST from the **Program Selection** section. Leave all the others options at their default settings, particularly the option to search all the proteins available.

Before you set **PSI-BLAST** going, click on the **Algorithm** parameters link and take a look at the PSI/PHI/DELTA BLAST Upload PSSM section. Here is offered the option to use a **PSSM** from a previous run PSI-BLAST, potentially on a different database (but with the Threshold same query sequence). Accept the default that database entries Pseudocount scoring better than an Expect Threshold of 0.005 be offered for inclusion into the **PSSM** of each successive **PSI-BLAST** iteration. Remember the buttons.

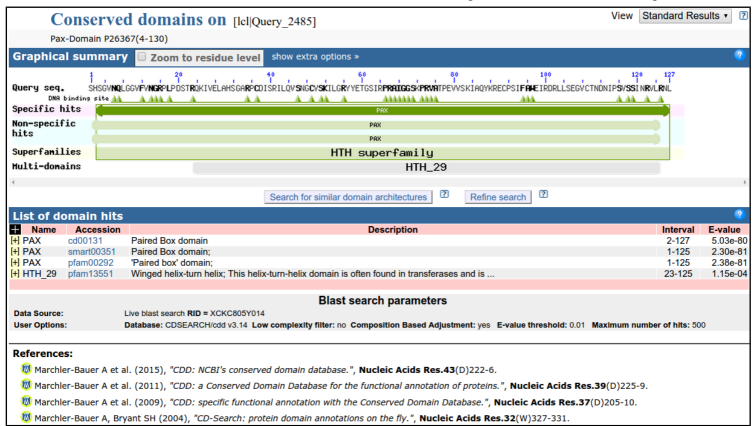


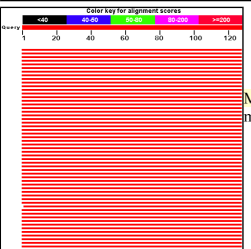
What do you suppose the choice of **Pseudocount** might influence?

Elect to Show results in a new window and then click on the button. After several moments of deep thought, **PSI-BLAST** will come back with its first set of results, at the top of which is a report that (unsurprisingly) matches have been detected between the query sequence and several domain databases. For more detail, click on the **Conserved Domains** graphic.

10:43:40 AN

SMART, Pfam and the NCBI Conserved Domains database hits are reported. None should be a surprise.





Moving back to the main **PSI-BLAST** results, you will see that there are many high quality hits covering the whole length of the query sequence.

The best **500** of these are listed.

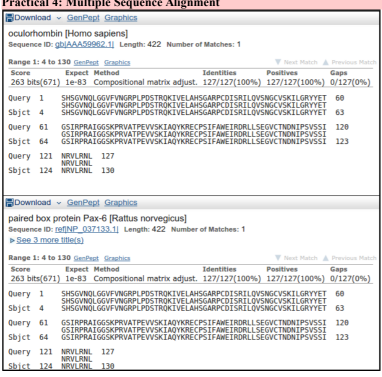
All the listed hits are selected for inclusion into the **PSSM** for the next iteration. Unless you feel strongly about any particular entry, leave them all selected.

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Sec	Sequences producing significant alignments with E-value BETTER than threshold									
Se	Select: All None Selected:0									
AT	Alignments Download <u>GenPept Graphics Distance tree of resu</u>	ılts <u>Mul</u>	tiple ali	<u>gnment</u>					Ф	
	Description	Max score	Total score	Query	E value	Ident	Accession	Select for PSI blast	to build PSSM	
	PREDICTED: paired box protein Pax-6 isoform X4 [Macaca nemestrina]	262	262	100%	1e-83	100%	XP 011722295.1	•		
	PREDICTED: paired box protein Pax-6 isoform X2 [Ursus maritimus]	263	263	100%	1e-83	100%	XP 008685073.1	•		
	oculorhombin [Homo sapiens]	263	263	100%	1e-83	100%	AAA59962.1	•		
	paired box protein Pax-6 [Rattus norvegicus]	263	263	100%	1e-83	100%	NP 037133.1	•		
	PREDICTED: paired box protein Pax-6 isoform X2 [Fukomys damarensis]	263	263	100%	1e-83	100%	XP 010638711.1	•		
	PREDICTED: paired box protein Pax-6 isoform X2 [Cavia porcellus]	263	263	100%	1e-83	100%	XP 003464531.1	•		
	PREDICTED: paired box protein Pax-6 isoform X2 [Aotus nancymaae]	263	263	100%	1e-83	100%	XP 012307699.1	•		
	PREDICTED: paired box protein Pax-6 isoform X2 [Callorhinchus milii]	263	263	100%	1e-83	100%	XP 007885973.1	•		
	PREDICTED: paired box protein Pax-6 isoform X2 [Heterocephalus glaber]	263	263	100%	1e-83	100%	XP 004851665.1	•		
	PREDICTED: paired box protein Pax-6 isoform X2 [Octodon degus]	263	263	100%	1e-83	100%	XP 004638029.1	•		
	PREDICTED: paired box protein Pax-6 [Poecilia reticulata]	261	261	100%	1e-83	98%	XP 008404092.1	•		

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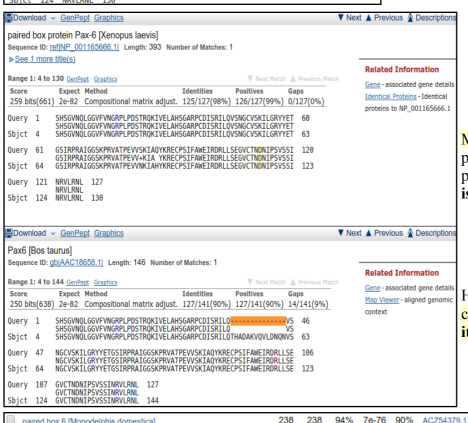
Practical 4: Multiple Sequence Alignment



Move down to the **Alignments** section of the results and you will see that many of the top hits match the query exactly.

Note that many of the top hits come from the GenPept database (roughly equivalent to the TrEMBL section of UniProtKB).

How might the inclusion of relatively poor quality sequences and the presence of so much duplication have been minimised?



Move down far enough and you will see less perfect matches, some of which involve proteins with the extra 14 amino acids of isoform 5a of PAX6 HUMAN.

Having browsed your results sufficiently, click on the Go button to Run PSI-Blast iteration 2 It is at the bottom of the hit list.

paired box 6 [Monodelphis domestica] 7e-76 ACZ54379.1 PREDICTED: paired box protein Pax-6-like isoform X1 [Acromyrmex ec 246 246 8e-76 XP 011063177.1 putative paired box protein pax-6 [Schistosoma mansoni] 254 99% 1e-75 90% CCD79466.1 putative Paired box protein Pax-6 [Operophtera brumata] 232 232 90% 1e-75 97% KOB68243.1 234 234 1e-75 89% NP 001189460.1 4 twin of eyeless [Bombyx mori] 242 242 99% 91% 4 PREDICTED: eyeless isoform X3 [Tribolium castaneum] 2e-75 XP 008192001.1 PREDICTED: eyeless isoform X2 [Tribolium castaneum] 242 242 99% 2e-75 91% XP 008192000.1 245 99% 94% 1 PREDICTED: paired box protein Pax-6-like isoform X1 [Megachile rotur 245] 2e-75 XP 012148240.1 Hypothetical protein CBG04481 [Caenorhabditis briggsae] 239 239 99% 2e-75 82% XP 002644124.1 pax6-like protein [Euperipatoides kanangrensis] 233 233 92% 3e-75 95% AGC51117.1 1 paired box protein Pax-6 [Clonorchis sinensis] 251 99% 251 3e-75 90% GAA48050.1 XP_013196296.1 4 PREDICTED: paired box protein Pax-6-like [Amyelois transitella] 231 231 91% 3e-75 92% hypothetical protein T265 09221 [Opisthorchis viverrini] 251 3e-75 90%

After a few moments, PSI-**BLAST** will return with the results of searching through the database again using the PSSM derived from the hits of the first iteration(\square ed). This time the top of the list will be predominantly filled with hits that have already been incorporated into the PSI-**BLAST PSSM**. However, look far enough down the list and you will find some new highlighted yellow.

Once more, click on the button to **Run PSI-Blast iteration 3**. That is probably enough! It took 4 iterations before there were no more new sequences suggested for inclusion into the **PSMM** when I ran this last, so if you really want to take things to their logical conclusion, it should not detain you long.

Next, move to the top of the **Descriptions** list and **Select All**. Click on the **Multiple Alignment** button. You have elected to use the **NCBI** multiple alignment program **Cobalt** to align all the **PAX** domain sequences of your final **PSI-BLAST** iteration that match with an **Expect** score better than **0.001**. In an impressively short time, your alignment will appear.

Multiple Alignment Results - Pax-Domain P26367(4-130) (127 letters)

Move past the long list of proteins that have been aligned (the easiest way is to hide the **Descriptions** view).

Multiple Alignment Results - Pax-Domain P26367(4-130) (127 letters)

XE22VT2H211 (500 seqs)

The Descriptions Select All Re-align Alignment parameters

Alignments Mouse over the sequence Identifer for sequence title

At the top of the actual alignment, set View Format to Plain Text (.... and then hide the Descriptions again??), this being the easiest format to understand in a hurry. The alignment will have very ragged ends, but the important region of 120 or so amino acids representing the PAX domain is really quite impressive. In particular, the isoform 5a insertion is very convincing.

✓ XP_003977912	52	TRQKIVELAHSGARPCDISRILQTHDAVQVLDSEKVSNGCVSKILGRYYETGSIRPRAIGGSK	114
✓ XP_009296159	26	TRQKIVELAHSGARPCDISRILQTHADAKVQVLDNENVSNGCVSKILGRYYETGSIRPRAIGGSK	90
✓ XP_003246075	54	TRQKIVELAHSGARPCDISRILQVSNGCVSKILGRYYETGSIKPRAIGGSK	104
✓ XP_012793883	41	TRQRIIELAHSGARPCDISRILQVSNGCVSKILCRYYETGSIRPKAIGGSK	91
✓ XP_005991286	25	TRQKIVELAHSGARPCDISRILQTHADAKVQVLDIQNVSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ EFX75780	37	TRQKIVELAHSGARPCDISRILQVSNGCVSKILGRYYETGSIRPRAIGGSK	87
✓ ABB43131	25	TRQKIVELAHSGARPCDISRILQVSNGCVSKILGRYYETGSIRPRAIGGSK	75
✓ ETN66652	41	TRQKIVELAHSGARPCDISRILQVVSNGCVSKILGRYYETGSIKPRAIGGSK	91
✓ XP_006128959	56	TRQKIVELAHSGARPCDISRILQ <mark>THADAKVQVLDNQN</mark> VSNGCVSKILGRYYETGSIRPRAIGGSK	120
✓ XP_010874560	44	TRQKIVELAHSGARPCDISRILQTHDDSKVQVLDNENVSNGCVSKILGRYYETGSIRPRAIGGSK	108
✓ AFJ24746	53	TRQRIVELAHSGARPCDISRILQVSNGCVSKILCRYYETGSIRPKAIGGSK	103
✓ XP_007885968	25	TRQKIVELAHSGARPCDISRILQTHADAKVQVVDNRKVSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ BAA24024	42	TRQKIVELAHSGARPCDISRILQTHADAKVQVLDSQNVSNGCVSKILGRYYETGSIRPRAIGGSK	106
✓ XP_012307695	25	TRQKIVELAHSGARPCDISRILQ <mark>THADAKVQVLDNQN</mark> VSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ CBY09679	55	TRQKIVELAHSGARPCDISRILQVVSNGCVSKILARYYETGSIKPRAIGGSK	105
✓ XP_007181079	82	TRQKIVELAHSGARPCDISRILQ <mark>THADAKVQVLDNQN</mark> VSNGCVSKILGRYYETGSIRPRAIGGSK	146
✓ CAF29075	25	TRQKIVELAHSGARPCDISRILQTHADAKVQVLDSENVSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ XP_004264009	25	TRQKIVELAHSGARPCDISRILQ <mark>THADAKVQVLDNQN</mark> VSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ XP_009184622	25	TRQKIVELAHSGARPCDISRILQ <mark>THADAKVQVLDNQN</mark> VSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ AAW24017	55	TRQKIVELAHSGARPCDISRILQVSNGCVSKILARYYETGSIKPRAIGGSK	105
✓ XP_008547741	26	TRQKIVELAHSGARPCDISRILQVSNGCVSKILGRYYETGSIRPRAIGGSK	76
✓ XP_012162452	50	TRQKIVELAHSGARPCDISRILQVSNGCVSKILGRYYETGSIKPRAIGGSK	100
✓ XP_006975926	25	TRQKIVELAHSGARPCDISRILQTHADAKVQVLDNENVSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ KDR14710	21	TRQKIVELAHSGARPCDISRILQVVSNGCVSKILGRYYETGSIRPRAIGGSK	71
✓ XP_005530321	25	TRQKIVELAHSGARPCDISRILQ <mark>THADAKVQVLDNQN</mark> VSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ ABI98847	25	TRQKIVELAHSGARPCDISRILQ <mark>THADAKVQVLDNQN</mark> VSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ XP_010794780	44	TRQKIVELAHSGARPCDISRILQTHDEVQVLDSEKVSNGCVSKILGRYYETGSIRPRAIGGSK	106
✓ NP_001103907	26	TRQKIVELAHSGARPCDISRILQVVSNGCVSKILGRYYETGSIRPRAIGGSK	76
✓ XP_010356630	25	TRQKIVELAHSGARPCDISRILQ <mark>THADAKVQVLDNQN</mark> VSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ XP_010638709	25	TRQKIVELAHSGARPCDISRILQ <mark>THADAKVQVLDNQN</mark> VSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ <u>XP_005064878</u>	25	TRQKIVELAHSGARPCDISRILQTHADAKVQVLDNENVSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ NP_038655	25	TRQKIVELAHSGARPCDISRILQTHADAKVQVLDNENVSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ XP_005401829	25	TRQKIVELAHSGARPCDISRILQ <mark>THADAKVQVLDNQN</mark> VSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ XP_004638028	25	TRQKIVELAHSGARPCDISRILQ <mark>THADAKVQVLDNQN</mark> VSNGCVSKILGRYYETGSIRPRAIGGSK	89
✓ BAM74254	32	TRQRIVELAHSGARPCDISRILQVSNGCVSKILGRYYETGSIRPRAIGGSK	82

¹¹ Much more so than the **Full** alignment offered by **PFAM**, I would contend. Although, it has to be admitted, the **Pfam** alignment included more sequences and I suspect they would have gone for a less closely homologous set of sequences. Even so ... I think the alignment illustrated here is **MUCH** more beautiful!!

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 moer back ground 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 pertenent 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. **BUT**, 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 Multiple Sequence Alignment

Clustalx

How might you have saved the need to recompute the alignment by selecting an **Edit** option other than **Remove All Gaps**?

Pretty sure using **Remove Gap-Only columns** rather than **Remove All Gaps** would have got to the second alignment in one step. I struggle to see how this could fail, but cannot convince myself it would be absolutely certain to work. Maybe some of the sequences that were removed would subtly alter the alignment calculations? Doubtful, given how crude the whole thing is. Anyway, it would have been good enough for me in "Real Life" for this data. I did not suggest this approach as we had to make a **Fasta** format file with the entire sequence set to be aligned by the alternative software as a by product of working with **clustalx**. This would not be easily possible if the **Remove Gap-Only columns** short cut was taken.

Muscle

How do the options for the **OUTPUT TREE** relate to the output files of **ClustalX** and the difference between the way that **ClustalX** and **muscle** work?

I leave this question here in the hope that one day I will be able to offer a sensible answer. First draft answer below.

Essentially, both ClustalX and MUSCLE work in two stages. First they create Guide Tree(s) (ClustalX just one, MUSCLE creates 2). Then from the Guide Tree(s) they compute the final MSA. The purpose of saving the Guide Tree(s) to a file is to enable a rerun of the second phase with new parameter settings without having to rerun the first Guide Tree generation stage.

Of course, as mentioned above, utterly pointless if there is no way to change the parameters? but that is the theory.

More investigation by me and expansion of this answer required. Discussion with EBI current (2016.04.20).

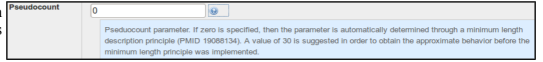
Comment on how one might choose between the range of options offered for the aligned parameter?

I cannot ... beyond suggesting it simply does not make sense? Going by what is offered at Wageningen, the choice should be between **aligned** and **input order**. i.e. the order of the original set of sequences to be aligned or the order after they have all been compared with each other and arranged into a **Guide Tree** ... or two.

Basic Bioinformatics. 20 of 23 10:43:40 AM

What do you suppose the choice of **Pseudocount** might influence?

I clicked with confidences upon the link to the help. It opined as illustrated.



I suppose the next step is to read **PMID 19088134**? There is most certainly no elucidation amongst the strangle of words offered here?

The article **Abstract** says:

"Position specific score matrices (PSSMs) are derived from multiple sequence alignments to aid in the recognition of distant protein sequence relationships. The PSI-BLAST protein database search program derives the column scores of its PSSMs with the aid of pseudocounts, added to the observed amino acid counts in a multiple alignment column. In the absence of theory, the number of pseudocounts used has been a completely empirical parameter. This article argues that the minimum description length principle can motivate the choice of this parameter. Specifically, for realistic alignments, the principle supports the practice of using a number of pseudocounts essentially independent of alignment size. However, it also implies that more highly conserved columns should use fewer pseudocounts, increasing the inter-column contrast of the implied PSSMs. A new method for calculating pseudocounts that significantly improves PSI-BLAST's; retrieval accuracy is now employed by default."

The article itself, continues in like vein how about we close our eyes and accept the defaults? I would just wonder why the whole thing does not commence with, at least an attempt, to answer the question in the forefront of my inquiry, which is .. "WHAT, in the current context, IS a pseudocount?". I do not believe it is as tricky as they appear to wish us to believe. I will try again later, when my view of the world is less storm infested.

DPJ - 2015.09.14