

# **IB16S**

## Introductory Bioinformatics

12-16 December 2016

(Second 2016 run of this Course)

## Basic Bioinformatics Sessions

Practical 4: Primer Design

### **Primer Design**

The prime intention of this exercise is to design a way to amplify a DNA fragment of reasonable size that includes a specific portion of the **PAX6** gene. The target region is that which includes the mutation you looked at earlier, that is a major cause of Aniridia. That is, the substitution that mutates the **33**<sup>rd</sup> amino acid position of the **PAX6** human protein. I remind that the details you discovered earlier are as follows:

Affected Patient: 33rd amino acid position mRNA Base position

Proline (P) 459

Wild Type: 33<sup>rd</sup> amino acid position Genomic DNA Base position

**Alanine** (A) 15714

The isolation, amplification and analysis of the target region of the genome could be affected by using restriction enzymes. In this case, there is more than one restriction enzyme whose cut site is dependant upon the mutation and so would produce a differing set of restriction fragments when used with the DNA of **Aniridia** affected patients to that normally expected. As long as those differences were course enough to be detected by a Restriction Fragment Length Polymorphism (RFLP) experiment. Software exists to select enzymes to isolate a chosen region of genomic DNA and to fragment that isolate in such a way it is possible to determine whether it includes the unfortunate mutation or not from the pattern of fragments generated.

For a variety of reasons, including the ready availability and ever decreasing cost of sequencing, this is typically not the preferred way to proceed. It is normally preferable to use Polymerase Chain Reaction (PCR) to isolate the region around the mutation and then to sequence samples from all individuals under examination. To do this, the first step would be to design suitable PCR primers. One program, in many different forms, is almost exclusively used for this purpose. The program is **primer3**. It is free and can be downloaded and run under linux and windows (at least). It is available as part of the **EMBOSS** package (**eprimer3**) and from a number of websites, including at the **Massachusetts Institute** of Technology (**MIT**)<sup>1</sup>:

```
http://bioinfo.ut.ee/primer3/
```

This site is popular with many users wanting the very latest version of the software, complete control over the various options offered by **primer3** and are not too concerned with using a database search to check the uniqueness of the products they will produce.

Another excellent **primer3** web interface developed in the Netherlands is available at:

```
http://www.bioinformatics.nl/cgi-bin/primer3plus/primer3plus.cgi
```

The site incorporates access to a **blast** search to check the uniqueness of the selected primers (important if unwanted **PCR** products are to be avoided).

Mostly because of its completely seamless inclusion of a **blast** search to compare potential primers with appropriate sequence collections, I suggest we here use **primer3** as implemented at the **NCBI**, even though it offers less than complete control over the execution of **primer3** itself. Go to:

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

Click on the BLAST option. Select



from the Specialized BLAST section.

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Upload your genomic PAX6 sequence using the Browse (or Choose File) button for the PCR Template.

You have established that the mutation of greatest interest is the G/C substitution at position 15714 of the genomic sequence copied from Ensembl. It is logical therefore to specify that this feature be included in the PCR production.

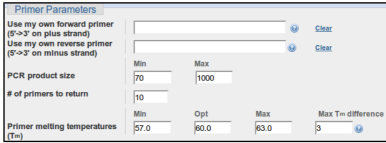


specify that this feature be included in the PCR product not too near either end. Accordingly, request the Forward primer to be chosen From the region starting at base pair 15000 and continuing To base pair 15700. Set the range for the Reverse primer to be From 15800 and To 16500.

The default PCR product size is specified in the Primer Parameters section as between 70 and 1000 base pairs. This seems fine.

Primer Parameters
Use my own forward prints on plus strand)
Use my own reverse prints.

I would not presume to advise you on the melting temperatures that were most suitable<sup>2</sup>. For this exercise, the defaults work splendidly.



By default, primer-BLAST will report the (Tm) Primer melting temperatures best 10 primer pairs it can find (# of primers to return). This is r

best 10 primer pairs it can find (# of primers to return). This is plenty for the exercise.

Do you think 10 primer pair suggestions is sufficient? If not, what number would you choose?

In addition to running **primer3** to suggest primers, **Primer-BLAST** checks against the possibility of unwanted **PCR** products by comparing potential primers against an appropriate sequence database with **blast**.

In the Primer Pair Specificity
Checking Parameters section, set the
Database selection to RefSeq
representative genomes. Leave the
Organism set as Homo sapiens.

You thus request each potential pair of **PCR** primers to be compared to the entire human genome. Thus unintended products of similar size to the intended product, can be identified.

The ideal conclusion is "just one product will be produced, on Splice variant har chromosome 11, in the region of the PAX6 gene".

Primer Pair Specificity Ch	necking Parameters						
Specificity check	☑ Enable search for primer pairs specific to the intended PCR template  ☑						
Search mode	Automatic 💲						
Database	Refseq representative genomes 💲 📦						
Exclusion	Exclude predicted Refseq transcripts (accession with XM, XR prefix)						
	sample sequences (a)						
Organism	Homo sapiens						
	Enter an organism name (or organism group name such as enterobacteriaceae, rodents), taxonomy id or select						
	from the suggestion list as you type.						
	Add more organisms						
Entrez query (optional)	•						
Primer specificity stringency	Primer must have at least 2 📫 total mismatches to unintended targets, including						
	at least 2 ‡ mismatches within the last 5 ‡ bps at the 3' end 🕖						
	Ignore targets that have 6 🗘 or more mismatches to the primer. 🔞						
Max target size	4000						
Splice variant handling	☐ Allow primer to amplify mRNA splice variants (requires refseq mRNA sequence as PCR template input) ⓐ						
V/C 22							

Use the appropriate button to discover the purpose of the Max target size parameter.

This is a relatively new parameter replacing a

Max target size

4000

This specific are latively new parameter replacing a

which was somewhat less obvious. The reason for the **Max target size** parameter is surely pretty transparent, so maybe there is now less requirement to wake up its button? For the present, the maximum size of any proposed **PCR** product, in this instance, is **1,000** base pairs (the form default). So the greatest size of an unwanted product that might be a problem (the **Max target size**) must be small enough to potentially be mistaken for a real product of **1,000** base pairs. **4,000** base pairs seems a bit cautious to me? However, unless you feel strongly about the matter, accept the default value of **4000**.

What value would you choose here if you were looking for uncluttered results?

2 My policy has been to not discuss parameters that pertain to the experimental conditions. In future versions of these notes, I will include discussion of some of these parameters. In the mean time, the buttons are very helpful. I would also suggest the MIT site (or the Wageningen site) for very readable explanations linked from every parameter. The full **primer3** manual can be found here.

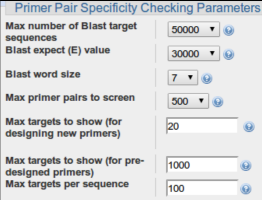
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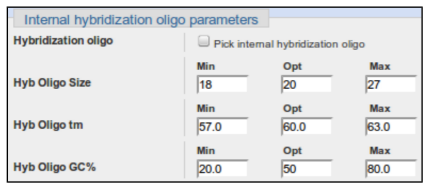
Before setting primer-BLAST going, click on the Advanced parameters button. Not really so Advanced? More Avoidable by those in a hurry. At the top are the Primer Pair Specificity Checking Parameters that control the

way that **blast** is run. Note the buttons offering explanation.

Note the very high default **Blast expect (E) value**, suggesting you will be interested in matches with your primers that might occur up too 30000 times by chance! This does make sense as the primers will be very short and so many good, even exact, "chance" matches might be expected against a large database.

Comment upon the small default value for the **Blast word size**?

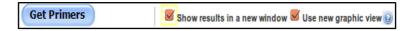




Note that you could get primer-BLAST to suggest an Internal hybridisation oligo, but decline the invitation this time.

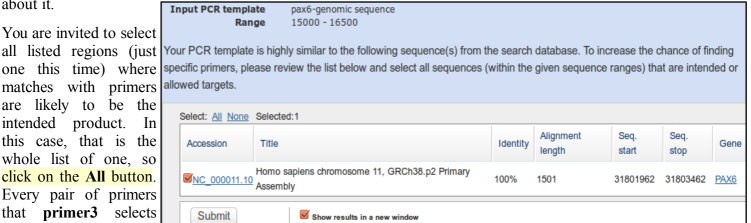
Accept all the Advanced parameters as they are. Ask primer-BLAST to Show results in a new window.

Click on the Get Primers button.



After a few moments of deep thought, **primer-BLAST** will notice that the template sequence you are using is **highly** similar (identical in fact) to part of an entry in the database being searched. Hardly surprising if one was to think about it

You are invited to select matches with primers allowed targets. are likely to be the intended product. this case, that is the whole list of one, so click on the All button. Every pair of primers that **primer3** selects must match this region



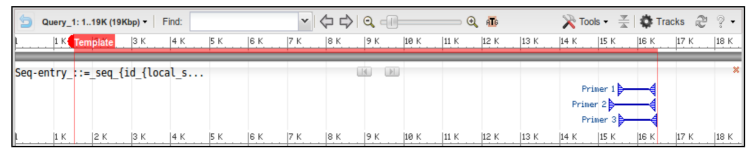
of Chromosome 11 as it is precisely the region investigated by primer3 in the first place. This process avoids blast reporting intended products as unintended products.

Finally, all is ready, so ask to **Show results in a new window** and then click on the **Submit** button.

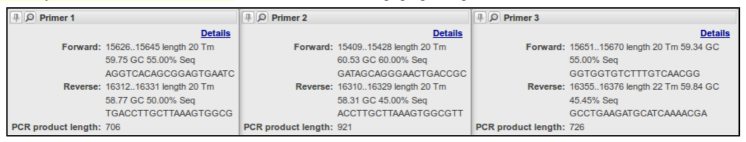
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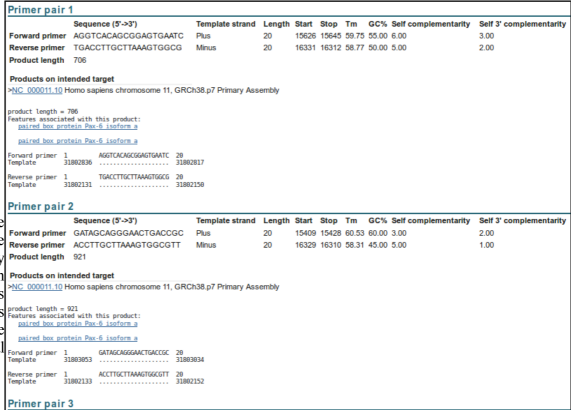
Once you have revelled in the opportunity to twiddle the fingers and scratch the ear(s) whilst **primers3** and **blast** go merrily about their appointed tasks, you will receive your results. These should look disarmingly like mine if all has gone well.

The summary **Graphic view** suggest just 3 solutions met the default criteria for success used by **primer3**. Up to 10 were permitted<sup>3</sup>.



Hover your mouse over one or more and further details will pop up in separate windows.





20

Template strand Length Start Stop Tm GC% Self complementarity Self 3' complementarity

15651 15670 59.34 55.00 5.00

16376 16355 59.84 45.45 8.00

None of your three suggested primer pairs are reported with any unintended products, even given the very generous suggestion that products 4000 bases long should be considered a potential problem<sup>4</sup>.

Which rather makes mock of all the deep thought employed deciding upon the most sensible maximum number of predictions to be reported.

Sequence (5'->3')

Reverse primer GCCTGAAGATGCATCAAAACGA Minus

NC 000011.10 Homo sapiens chromosome 11, GRCh38.p7 Primary Assembly

GGTGGTGTCTTTGTCAACGG 20 .......31892792 GCCTGAAGATGCATCAAAACGA 22 .......31892197

Forward primer GGTGGTGTCTTTGTCAACGG

Product length 726

Products on intended target

Reverse primer 1 Template 31802086 3.00

2.00

<sup>4</sup> This was not true until very recently. **Primer-BLAST** reported many more primer pair suggestions and quite a few unintended products for each. The previous parameter restriction the length of unintended products was somewhat more generous.

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As well as suggesting primers for PCR (or other purposes) and (optionally) suggesting hybridisation oligos, **primer-BLAST** can be used to evaluate user-selected primers. Earlier, you saved a pair of primer sequences associated with **PAX6** when searching the nucleotide databases at the **NCBI**. It would be interesting to discover the product these might produce. To do this you need an unsullied **Primer-BLAST** page. Go again to:

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

#### Click on the BLAST option. Select



from the **Specialized BLAST** section.

Upload your genomic PAX6 sequence using the Browse (or Choose File) button for the PCR Template.



Open up the file you made containing the primers from **GenBank** (pax6\_primers.fasta) in a text editor.

Copy and Paste the two primer sequences into the Use my own forward primer and Use my own reverse primer boxes as appropriate.

In the Primer Pair Specificity Checking Parameters section, set the Database selection to RefSeq representative genomes.

**Leave the Organism** as **Homo sapiens**.

Length Start Stop Tm GC%

12036 12060 69.45 60.00 6.00

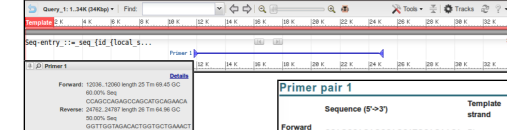
24787 24762 64.96 50.00 4.00

Raise the Max target size parameter from 4000 to 20000. You should check for enormous unintended products with this run of Primer-BLAST. The reasons for this will soon become apparent.

### Ask primer-BLAST to Show results in a new window. Click on the Get Primers button.







primer Reverse

primer

Product

After a short thrill filled pause, you will receive a result that should again looks more that a trifle like mine.

Self

complementarity

Self 3

0.00

1.00

complementarity

Seemingly a fine match. Even the single **potentially unintended product** reported is actually the **intended product**. For some reason, **Primer-BLAST** does not eliminate intended products when searching for unintended ones when investigating user specified primers<sup>5</sup>?

Products on potentially unintended templates

NC 000011.10 Homo sapiens chromosome 11, GRCh38.p7 Primary Assembly

product length = 12752

Features associated with this product:
 paired box protein Pax-6 isoform a

CCAGCCAGAGCCAGCATGCAGAACA Plus

**GGTTGGTAGACACTGGTGCTGAAACT Minus** 

paired box protein Pax-6 isoform a

paired box protein Pax-6 isoform a

Forward primer 1 CCAGCCAGAGCCAGCATGCAGAACA 25

- I have asked the guys at NCBI to explain. No full answer as yet, further prodding required. Prodded last 2016.04.02. Maybe I give up?
- Apparently, such a PCR product is possible! However, above 5,000 base pairs would be slow, require very close attention and be prone to errors.

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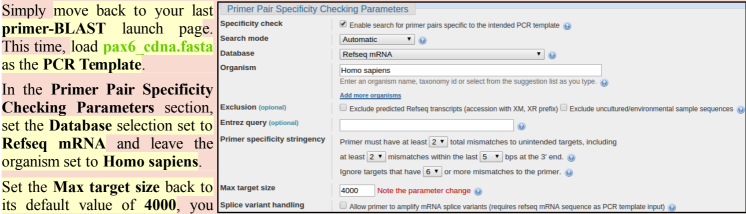
Unfortunately, the only paper referenced does not explain what might be going on particularly clearly. However, there is a hint that the primers you saved were designed for use with mRNA/cDNA data. Therefore it might be interesting to run primer-BLAST one last time with pax6 cdna.fasta as the PCR Template.

primer-BLAST launch page. This time, load pax6 cdna.fasta as the PCR Template.

In the Primer Pair Specificity Checking Parameters section, Exclusion (opion set the Database selection set to Entrez query (optional) Refseq mRNA and leave the organism set to **Homo sapiens**.

Set the Max target size back to Max target size its default value of 4000, you Splice variant handling

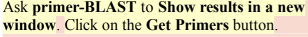
length

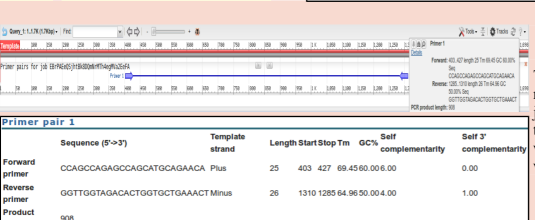


should expect much smaller mRNA products this time, so no need for extending this maximum beyond 4000.

These selections suppose that the design of PCR product was for selection from a library of all human cDNAs.

**Get Primers** 





The result is a much more reasonable **Product length** of just 908 base pairs, reinforcing the theory that these primers were indeed designed for use with a cDNA library.

Show results in a new window W Use new graphic view (

Before moving on, afford a quick glance at the report offered concerning possible unintended products. Here primer-BLAST warns against human mRNAs that might be cloned along with the intended target.

The first thing to note is that the intended target is not generated from a RefSeq mRNA. It comes from an mRNA taken from an aniridia patient directly. Therefore, there is no unintended product that we can ignore because it is really the intended product discovered by a different route, even though no filtering of the **RefSeq** database was undertaken.

All the unintended products could/would potentially be generated by the primers under investigation and have the potential to cause confusion. If you look down the list, you should conclude that the 16 unintended products come from 16 of the 24 RefSeq PAX6 transcripts first noted by GeneCards and then confirmed later by blast.

```
Products on potentially unintended templates
NM 001310159.1 Homo sapiens paired box 6 (PAX6), transcript variant 9, mRNA
product length = 908
Reverse primer 1
                GGTTGGTAGACACTGGTGCTGAAACT 26
             1021 ...... 996
Template
NM_001310158.1 Homo sapiens paired box 6 (PAX6), transcript variant 8, mRNA
product length = 950
                 CCAGCCAGAGCCAGCATGCAGAACA 25
             Template
Reverse primer 1
                  GGTTGGTAGACACTGGTGCTGAAACT 26
Template
            1445 ..... 1420
XM 006718246.2 PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X11, mRNA
product length = 707
.
Forward primer 1
                 CCAGCCAGAGCCAGCATGCAGAACA 25
Template 457 .....
                  GGTTGGTAGACACTGGTGCTGAAACT 26
             1163 ...... 1138
XM_011520152.1 PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X10, mRNA
product length = 749
orward primer 1
             Template
Reverse primer 1
                  GGTTGGTAGACACTGGTGCTGAAACT 26
             1205 .....
Template
XM 005252956.3 PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X9, mRNA
Forward primer 1
                  CCAGCCAGAGCCAGCATGCAGAACA 25
Reverse primer 1
                  GGTTGGTAGACACTGGTGCTGAAACT 26
```

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9 of the 11 NM\_ good quality transcripts are detected. 7 of the 13 poorer quality XM\_ "PREDICTED" transcripts are also present. So 16 of the 24 PAX6 transcript sequences in RefSeq were detected.

Why do you suppose **blast** did not pick up all the transcripts? XM 005252955.3 PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X8, mRN. Note that the intended product is 908 base pairs long. Note that all the unintended products except two, near the top of the list are either 908 long or 950 long. A difference of 42. >XM\_011520150.1 PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X6, mRNA How would you tell quickly which isoform was represented by each mRNA listed here? >XM\_011520149.1 PREDICTED; Homo sapiens paired box 6 (PAX6), transcript variant X5, mRNA Some fairly redundant questions to finish this section. I think I have already answered them all. But maybe you might wish to differ? XM\_005252954.3 PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X4, mRNA Is the number of "potentially unintended products" as you would you expect, given the evidence from GeneCards, Ensembl and blast?\_ NM 001258465.1 Homo sapiens paired box 6 (PAX6), transcript variant 7, mRNA NM\_001258464.1 Homo sapiens paired box 6 (PAX6), transcript variant 6, mRNA For all the "potentially unintended products", the selected primers match exactly. Can you explain this? >NM\_001258463.1 Homo sapiens paired box 6 (PAX6), transcript variant 5, mRNA product length = 950 >NM 001258462.1 Homo sapiens paired box 6 (PAX6), transcript variant 4, mRNA The "potentially unintended products" are of different sizes. Can >NM\_001604.5 Homo sapiens paired box 6 (PAX6), transcript variant 2, mRNA you explain the difference between the possible product lengths? 1 GGTTGGTAGACACTGGTGCTGAAACT 26 1392 ......1367 >NM\_000280.4 Homo sapiens paired box 6 (PAX6), transcript variant 1, mRNA NM\_001127612.1 Homo sapiens paired box 6 (PAX6), transcript variant 3, mRNA Are the numbers of "potentially unintended products" of each possible length consistent with your **blast** results? 1 GGTTGGTAGACACTGGTGCTGAAACT 26 1362 ......13

#### **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.

#### **Model Answers**

From your investigations of Primer Design

#### Do you think 10 primer pair suggestions is sufficient? If not, what number would you choose?

Until very recently, the default here was 5. That seemed rather low to me. I included this question to solicit opinion rather than to impart knowledge. A default of 10 seems more in line with my instincts, but people who use this program seriously mostly tell me that they can select suitable primers from the first 2 or 3 suggestions of the program. So, 5 would seem a good choice and 10 would be moving towards cautiously overdoing things.

On the whole, informed opinion suggests that 10 suggestions will be more than enough in most circumstances.

#### What value would you choose here if you were looking for uncluttered results?

#### Summary:

Clearly, the smaller the number chosen, the shorter will be the list of spurious products. However, pick something too small and you risk including unintended product(s) that could cause confusion. The size selected must be sufficient that larger unwanted PCR product(s) could easily be spotted by other means (simply by size?).

#### Full Answer:

Well, mostly for me, and just in case you were curious, when I first wrote the question, the parameter was very different and not so easy to understand. Pure self indulgence, I know, but here is the history. The parameter explained itself, via the button, thus:



I interpreted this to mean that only **blast** predicted products of up to **X**+**4**,000 base pairs, where **X** base pairs is the length of the intended target, will be given any regard. It is thus assumed that a difference of **4**,000 base pairs between an intended PCR product (predicted by **primer3**) and a spurious product (detected by **blast**) can easily be detected simply by size difference.

Of course this parameter also will reject unwanted **blast** predicted products that are less than X-4,000 base pairs will be given any regard. Given the largest possible **primer3** suggestion will be 1,000 base pairs (the form setting for the exercise specifies products of between  $100^{7}$  and 1,000 base pairs), this is hardly an issue here.

#### Comment upon the small default value for the **Blast word size**?

By default, **blast** will be looking for aligned exactly matching blocks of 7 nucleotides when identifying where a primer might match a database entry. The entire primer match with the template sequence does not have to be exact for the primer to be acceptable. The entire primer is typically only around **20** bases long. And word size much more that 7 would clearly miss too much to be effective.

<sup>7</sup> The form explicitly declares a minimum of 70, but the ranges from which the **forward** & **reverse** primers must come (15000-15700 & 15800-16500) make the smallest possible **primer3** prediction 100 base pairs long.

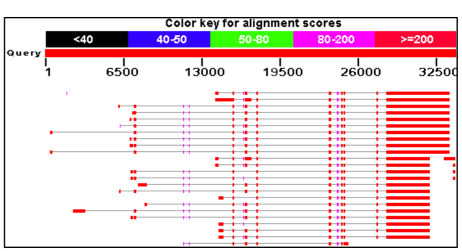
Why do you suppose **blast** did not pick up all the transcripts?

#### Summary:

Well, the simple answer is that the transcripts that were not detected as unwanted products cannot include either the forward primer, or the reverse primer, or both. This is, almost, the only possible explanation.

#### **Full Answer:**

Of course, for this run, you did specify that you were not interested in products longer than **4,000** base pairs, so it could be that one or more products were possible but longer than that? I suspect this would only be feasible if there were retained introns involved, but previous **blast** results do not suggest this to be the case. I would say the only possible candidate for an over-length product might be the second hit down in the graphical representation generated



previously by **blast**. The first and third exons from the left look a bit bloated, but not really sufficiently to cause a problem.

It might also be that unwanted PCR products are eliminated/introduced due to variations in the predicted transcripts. However, this can be ruled out as previous experiments, **blast** assures us that all **24** potential transcripts match the genomic sequence exactly.

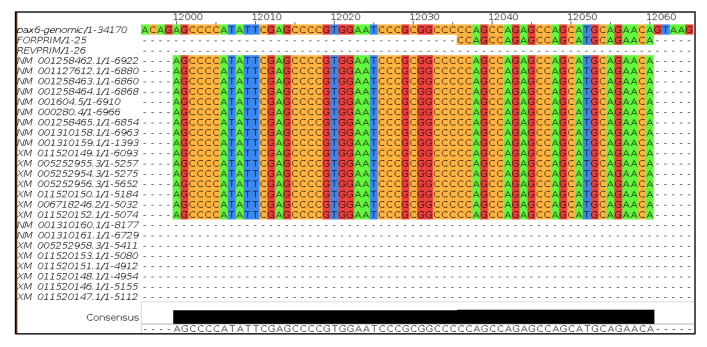
**Enough!** Only because I want to, I will compute the alignments to prove the missing primer matches. Read no further unless you are truly in the mood. Much of the reason for recording the rest of this answer is that, apart from enjoying the pursuit of irrelevant detail, I also wanted to remember how I made the alignments and certainly feel I could have made both these, and my point much more simply? Suggestions welcome.

Home sapiens paired box 6 (PAX6), transcript variant 1, mRNA   9659   15161   24%   0.0   100%   NM 0/4	01310161.1 01310160.1 01310158.1 01258465.1 01258464.1 01258463.1 01258462.1
Homo sapiens paired box 6 (PAX6), transcript variant 3, mRNA   9659   12929   20%   0.0   100%   NM   00	01310158.1 01258465.1 01258464.1 01258463.1
Homo sapiens paired box 6 (PAX6), transcript variant 7, mRNA   9659   12729   20%   0.0   100%   NM   00	01258465.1 01258464.1 01258463.1
Homo sapiens paired box 6 (PAX6), transcript variant 1, mRNA   9659   12761   20%   0.0   100%   NM   00	01258464.1 01258463.1
Home sapiens paired box 6 (PAX6), transcript variant 5, mRNA   9659   12737   20%   0.0   100%   NM   01	01258463.1
Homo sapiens paired box 6 (PAX6), transcript variant 4, mRNA   9659   12862   20%   0.0   100%   NM   01	
Homo sapiens paired box 6 (PAX6), transcript variant 2, mRNA   9659   12833   20%   0.0   100%   NM   01	01258462.1
Homo sapiens paired box 6 (PAX6), transcript variant 1, mRNA   9659   12942   20%   0.0   100%   NM   01	
Homo sapiens paired box 6 (PAX6), transcript variant 3, mRNA   9659   12791   20%   0.0   100%   NM   00	01604.5
PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X13, mRNA 6613 10063 15% 0.0 100% XM 00	00280.4
	01127612.1
DREDICTED: Home seniors paired box 6 (DAY6) transcript variant V12 mRNA 6613 9439 14% 0.0 100% VM 0.0	05252958.3
TITLE DIGITED. FROM COMPANDE PARTEU DOX O (FAXO), BAILSCHIPE VARIABLE X12, BILLINA	11520153.1
PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X11, mRNA 6613 9329 14% 0.0 100% XM 00	06718246.2
PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X10, mRNA 6613 9410 14% 0.0 100% XM 0	11520152.1
PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X9, mRNA 6613 10507 16% 0.0 100% XM 00	05252956.3
PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X8, mRNA 6613 9783 15% 0.0 100% XM 00	05252955.3
PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X7, mRNA 6613 9091 14% 0.0 100% XM 0:	11520151.1
PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X6, mRNA 6613 9637 15% 0.0 100% XM 0	11520150.1
PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X5, mRNA 6613 11324 17% 0.0 100% XM 0:	11520149.1
PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X4, mRNA 6613 9814 15% 0.0 100% XM 00	05252954.3
PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X3, mRNA 6613 9172 14% 0.0 100% XM 0	11520148.1
PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X2, mRNA 6613 9502 15% 0.0 100% XM 0:	11520147.1
PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X1, mRNA 6613 9576 15% 0.0 100% XM 0	11520146.1
PREDICTED: Homo sapiens elongator acetyltransferase complex subunit 4 (ELP4), transc 1775 1775 2% 0.0 100% XM 00	05252865.2
Homo sapiens paired box 6 (PAX6), transcript variant 9, mRNA 647 2630 4% 0.0 100% NM 00	

OK, I started by computing an alignment that was a mapping of all **24** transcripts onto the **PAX6** genomic regions as represented in the file **pax6\_genomic.fasta**. I used a program called **gmap**, which like **spline** (used in the exercise) is designed to align cDNA/mRNA sequences with corresponding genomic sequences. The version of **gmap** I used runs under **linux** from the command line. It has the advantage over **spline** that is will align more than one cDNA/mRNA sequence against the genome in one run. Unfortunately, it does not generate an output format that can be easily displayed in the way I required here. I did try to persuade a couple of general multiple alignment programs (**clustalw** & **muscle**) to make me a usable alignment, but ran into the same difficulties we experienced in the exercise. I failed to find gap penalties that would get the programs to gap the larger introns. Even if I had succeeded to get the gaps in the right place, I would not have believed them to be placed with sufficient accuracy for the same reasons this was not possible when we tried the same trick with general alignment software for just one cDNA sequence against the genome in the exercise.

So, I made a rough alignment with **clustalw** and edited it to exactly what was suggested by **gmap** using **jalview**. This took **HOURS**. There has to be a better way!! You have already used all the software mentioned except **gmap** and **clustalw**. You will use **clustalw** and see how **jalview** can be use to edit, as well as just view, alignments a little later.

All that effort to show that the region around the forward primer looks like this:

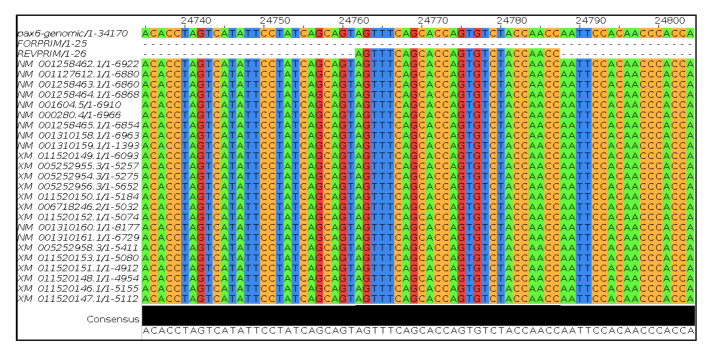


Showing clearly that the 8 transcripts:

NM\_001310160.1 NM\_001310161.1 XM\_005252958.3 XM\_011520153.1 XM\_011520148.1 XM\_011520146.1 XM\_011520147.1

Have the exon that includes the forward primer spliced out and so will not produce any PCR product. Feel free to check this by comparing the textual results of your **blast** of the genomic sequence against the **RefSeq** mRNAs and the results of **PRIMER-BLAST**. I did, it was lots and lots of fun and I ended up content that all was logically consistent.

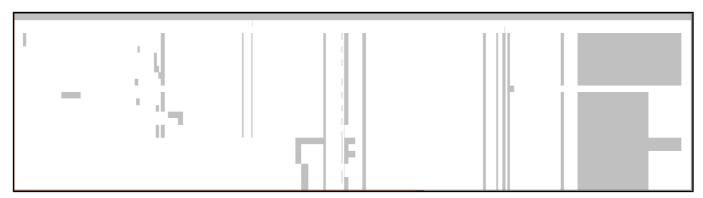
The alignment around the reverse primer looks like this:



All 24 putative transcripts match the reverse primer perfectly. So blast should indeed find 16 of the 24 transcripts sequences in RefSeq, and it does.

#### **Model Answers**

**Jalview** offers an overview of the entire alignment. The top row shows the genomic sequence. The second row shows the position of the forward primer. The third row shows the position of the reverse primer.



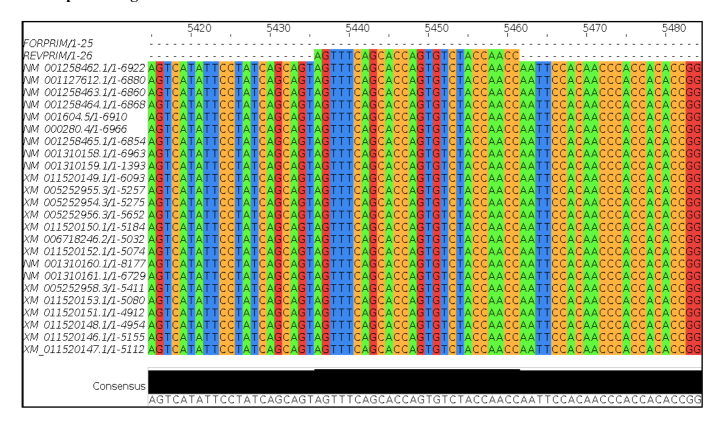
Except for the order of the transcripts, this view is very similar to the overview graphic generated by **blast**. The transcripts missing the forward primer, which isoform each transcripts represents and the fact that all transcripts match the reverse primer should be very clear.

Finished Dave? Well no, not quite. I wondered why I had included the genomic sequence in my alignment. Finding no answer to that question, I tried to make an alignment of just the primer sequences and the mRNAs. I thought this would be easy. I was wrong. The general programs are still going to get the gaps wrong whatever penalties are used. Some transcripts have exons entirely missing in all other transcripts leaving no clues as to which way round they should be aligned. The scaffold provided by the genomic sequence was essential. So, I made an mRNA only alignment by editing the alignment discussed above with **jalview**. This was easy (although you would not think so given the time it took me to work out how to do it!). I loaded the alignment into **jalview**, deleted the genomic sequence and then removed all empty columns (that is, all columns with no bases in them due the the removal of the genomic sequence). Clever eh? Just because it is there, here are the pictures.

#### **Forward primer region** (the primer is right at the end of an exon):

	2760	<del></del>	277	70 .	27	780	27	90		2800		2810		2820	
FORPRIM/1-25					- CCA	GCCAG	AGCCÁ	GCAT(	ĠCAG/	AACA					
REVPRIM/1-26															
NM 001258462.1/1-6922	CCG	GGAA	TCCCG	CGGCC	CCCA	GCCAG	AGCCA	GCAT(	GCAG	AACA					
NM 001127612.1/1-6880	CCG	GGAA	TCCCG	CGGCC	CCCA	GCCAG	AGCCA	GCAT(	GCAG	AACA					
NM 001258463.1/1-6860	CCG	GGAA	TCCCG	CGGCC	CCCA	GCCAG	AGCCA	3CAT	GCAG,	4ACA					
NM 001258464.1/1-6868	CCG	GGAA	TCCCG	CGGCC	CCCA	GCCAG	AGCCA	GCAT(	GCAG	AACA					
NM 000280.4/1-6966	CCG	GGAA	TCCCG	CGGCC	CCCA	GCCAG	AGCCA	GCAT(	GCAG	AACA					
NM 001258465.1/1-6854	CCG	GGAA	TCCCG	CGGCC	CCCA	GCCAG	AGCCA	GCAT(	GCAG	AACA					
NM 001310158.1/1-6963															
NM 001310159.1/1-1393	CCG	GGAA	TCCCG	CGGCC	CCCA	GCCAG	AGCCA	GCAT(	GCAG	AACA					
XM 011520149.1/1-6093	CCG	GGAA	TCCCG	CGGCC	CCCA	GCCAG	AGCCA	GCAT(	GCAG	AACA					
XM 005252955.3/1-5257	CCG	GGAA	TCCCG	CGGCC	CCCA	GCCAG	AGCCA	<b>GCAT</b> (	GCAG	AACA					
XM 005252954.3/1-5275	CCG	GGAA	TCCCG	CGGCC	CCCA	GCCAG	AGCCA	GCAT(	GCAG	AACA					
XM 005252956.3/1-5652	CCG	GGAA	TCCCG	CGGCC	CCCA	GCCAG	AGCCA	3CAT	GCAG	AACA					
XM 011520150.1/1-5184	CCG	GGAA	TCCCG	CGGCC	CCCA	GCCAG	AGCCA	GCAT(	GCAG	AACA					
XM 006718246.2/1-5032															
XM 011520152.1/1-5074															
NM 001310160.1/1-8177											CTTTTC				
NM 001310161.1/1-6729											CTTTTC	AATTA	GCCTT	CCATG	CATG
XM 005252958.3/1-5411											CTTTTC	AATTA	GCCTT	CCATG	CATG
XM 011520153.1/1-5080											CTTTTC	AATTA	GCCTT	CCATG	CATG.
XM 011520151.1/1-4912															
XM 011520148.1/1-4954															
XM_011520147.1 <b>/</b> 1-5112															
Consensus															
	CCG	ГGGAA	TCCCG	CGGCC	CCCA	GCCAG	AGCCA	GCAT	GCAG,	AACA	СТТТТС	AATTA	AGCCTI	CCATG	CATG

#### Reverse primer region:



#### Overview:

Without the evidence of the genomic sequence, the two leftmost exons could logically swap position. There is no transcript that includes both these exons and no overlap between either and any other exon in any transcript (most clearly verified from the previous **Overview** plot). Thus, there is no exon evidence of the order in which the two should appear.



Now I am done! This has to be the most over the top answer yet, but at least it kept me out of trouble for a while.

How would you tell quickly which isoform was represented by each mRNA listed here?

#### Summary:

All the mRNAs reported were of length 908, 950, 707 or 749.

A reasonable guess might be based on the length of the products? All those that are 908 bases might be assume to produce the 422 amino acid canonical isoform. All those that are 950 (i.e. 42 base pairs longer) might be assumed to 436 produce amino acid isoform 5a proteins (i.e. 14 amino acids longer).

Analogous reasoning might be applied to the mRNAs that are either 707 or 749 base pairs in length.

Just a guess of course, but one I would be happy to have faith in. To be certain, one would need to read the annotations of each listed **RefSeq** entry!

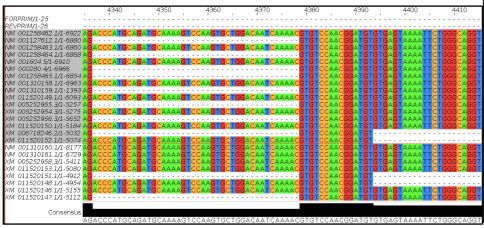
#### Full Answer:

From the illustrations of the last "Full Answer" (in particular, the jalview overviews), it is clear that all the mRNAs that produce a product include the region that determines which isoform is represented. That is, all are one isoform or the other.



The last two of the mRNAs that produce a PCR product, have a bit chewed out just after the isoform defining region (an exon spliced out, if you prefer). It is logical to suppose these would be the mRNAs from which the two shorter products were generated.

Indeed, looking at the relevant part of the mRNA only alignment shows them to be XM\_006718246 (product length 707, excluding the isoform 5a exon that suggest it codes for a canonical protein) and XM\_011520152 (product length 749, including the extra 42 base pairs suggesting it codes for an isoform 5a protein).



	NM 001310159.1 H	omo sapi	ens paired box 6 (PAX6), transcript variant 9, mRNA						
	product length Forward primer Template	= 908 1 114	CCAGCCAGAGCCAGCATGCAGAACA 25						
	Reverse primer Template	1 1021	GGTTGGTAGACACTGGTGCTGAAACT 26 996						
ŀ	> <u>NM 001310158.1</u> H	omo sapi	ens paired box 6 (PAX6), transcript variant 8, mRNA						
ŀ	product length Forward primer Template	= 950 1 496	CCAGCCAGAGCCAGCATGCAGAACA 25 520						
	Reverse primer Template	1 1445	GGTTGGTAGACACTGGTGCTGAAACT 26 1420						
Ŀ	>XM 006718246.2 PF	REDICTE	D: Homo sapiens paired box 6 (PAX6), transcript variant X11, mRNA						
	product length Forward primer Template	= 707 1 457	CCAGCCAGAGCCAGCATGCAGAACA 25						
	Reverse primer Template	1 1163	GGTTGGTAGACACTGGTGCTGAAACT 26 1138						
:	>XM 011520152.1 PREDICTED: Homo sapiens paired box 6 (PAX6), transcript variant X10, mRNA								
	product length Forward primer Template	= 749 1 457	CCAGCCAGAGCCAGCATGCAGAACA 25						
!	Reverse primer Template	1 1205	GGTTGGTAGACACTGGTGCTGAAACT 26 1180						
	>XM 005252956.3 PF	REDICTE	D: Homo sapiens paired box 6 (PAX6), transcript variant X9, mRNA						
ш	product length Forward primer Template	= 908 1 876	CCAGCCAGAGCCAGCATGCAGAACA 25						
!	Reverse primer Template	1 1783	GGTTGGTAGACACTGGTGCTGAAACT 26 1758						
	>XM 005252955.3 PF	REDICTE	D: Homo sapiens paired box 6 (PAX6), transcript variant X8, mRNA						
	product length Forward primer Template		CCAGCCAGAGCCAGCATGCAGAACA 25						
	Reverse primer Template	1 1388	GGTTGGTAGACACTGGTGCTGAAACT 26 1363						
ŀ	>XM 011520150.1 PF	REDICTE	D: Homo sapiens paired box 6 (PAX6), transcript variant X6, mRNA						
П	product length Forward primer Template		CCAGCCAGAGCCAGCATGCAGAACA 25 390						
	Reverse primer Template	1 1315	GGTTGGTAGACACTGGTGCTGAAACT 26 1290						
ŀ	>XM 011520149.1 PF	REDICTE	D: Homo sapiens paired box 6 (PAX6), transcript variant X5, mRNA						
П	product length Forward primer Template		CCAGCCAGAGCCAGCATGCAGAACA 25 1299						
	Reverse primer Template	1 2224	GGTTGGTAGACACTGGTGCTGAAACT 26 2199						
ŀ	>XM 005252954.3 PF	REDICTE	D: Homo sapiens paired box 6 (PAX6), transcript variant X4, mRNA						
1	product length Forward primer Template	= 950 1 457	CCAGCCAGAGCCAGCATGCAGAACA 25						
	Reverse primer	1	GGTTGGTAGACACTGGTGCTGAAACT 26						

All other transcripts that generate PCR products generate products of length either 908 or 950. Given the difference (42 base pairs) is exactly the size of the **isoform 5a** exon, it is reasonable to assume the transcripts generating PCR products of length 908 represent **canonical** proteins, whereas the transcripts generating PCR products of length 950 represent **isoform 5a** proteins.

Prettier? True, and I submit including sufficient evidence to be more that just a guess now.

#### **Model Answers**

Is the number of "potentially unintended products" as you would you expect, given the evidence from GeneCards, Ensembl and blast?

Yes, I think so, given you accept my investigation (see above) as to why there were only 16 "potentially unintended products" when you might have expected 24, given your blast results. GeneCards now encourages an initial expectation of 24 "potentially unintended products". Ensembl only uses the higher quality RefSeq mRNAs. Currently, Ensembl uses 10 of the 11 good quality RefSeq mRNAs to make its transcripts predictions. Close enough?

For all the "potentially unintended products", the selected primers match exactly. Can you explain this?

Well, of course they do??? All the transcripts found are generated from the same region of genomic DNA and therefore will be identical in all shared regions, including the primer regions. I suppose, in other instances, it would be possible to have transcripts with variation in the regions matching the primers insufficient to stop the primers working? But not in this case.

One might conclude there are no genuinely "unintended" products? All are real **PAX6** transcripts of varying certainty. A genuine unintended product would come from an entirely different part of the genome and would not necessarily match exactly with respect to the primers. They would just need to be "good enough to work".

The "potentially unintended products" are of different sizes. Can you explain the difference between the possible product lengths?

Are the numbers of "potentially unintended products" of each possible length consistent with your blast results?

Yes yes yes! I think both these questions made a bit more sense a few generations of these notes ago. We have already answered them sufficiently I suggest. I refer you to the answers above.

DPJ - 2016.12.06