FINDING GENES AND EXPLORING THE GENE PAGE AND RUNNING A BLAST (Exercise 1)

1.1 Finding a gene using text search. Note: For this exercise use <u>http://www.plasmodb.org</u>

a. Find all possible kinases in Plasmodium.

Hint: use the keyword "kinase" (without quotations) in the "Gene Text Search" box.



- How many genes did you get?
- How many of those are in *P. falciparum*? How did you find this out?
- What happens if you search using the word "kinases"? How many results did you return?

b. How can you increase the number of possible kinases in your results?

Hint: the search you did in 'a' will miss things like "6-phosphofructokinase" or "kinases" so you need to use a wild card in your search – try "kinase*", "*kinase" and "*kinase*" (without quotations).

- Did you get more results?
- Which one of the above wild card combinations gave you the largest number of kinases?
- How can you quickly examine the genes that were identified using the key word "*kinase*" but not with the word "kinase"? Hint: You can easily do this by combining search strategies. Click on "Add Step" then select "existing strategy":

				Add Step		
→	Run a new Search for Transform by Orthology Add contents of Basket Add existing Strategy Filter by assigned Weight	 Gene	,	Opened	,	Text* Text(2)*

- Select the right strategy from your list of Gene Strategies and combine the strategies with the correct operation:





- Do the results make sense? Do all the product names contain the word kinase?

c. Find only the kinases that specifically have the word "kinase" in the gene product name.

Hint: Use the text search page, the specific page where you can define the fields to be. There are many ways to navigate to the Text Search page.

- How did you get there?
- How many kinases have the word kinase in their product names?
- Did you remember to use the wild card?



1.2 Combing text search results with results from other searches

a. In exercise 1.1 you identified genes that have the word "kinase" somewhere in their product name. Can you now find out how many of these kinases are likely secreted?

Hint: grow your search strategy by adding a step. Choose a search that identifies genes with likely secretory signal peptides. How did you combine the search results?

My Strategies:	New Opened (1)	All (3) 🙆 Basket Example	s Help
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b. Now that you have a list of possible secreted kinases, how would you expand this strategy even further?

Hint: there is no wrong answer here....

- From a biological standpoint what else would be interesting to know about these kinases? Add more searches to grow this strategy.

- For example, how many of these secreted kinases also have transmembrane domains?

c. In the above example, how can you define kinases that have either a secretory signal peptide <u>AND/OR</u> a transmembrane domain(s)?

Hint: to do this properly you will have to employ the "Nested Strategy" feature. Why?



Notice the different results obtained in figures A (with nesting) and B (without nesting) below:





- 1.3 Visiting a specific gene page. Note: For this exercise use <u>http://www.plasmodb.org</u>
- a. Find the bifunctional dihydrofolate reductase-thymidylate synthase (DHFR-TS) gene (or, if you prefer, the apical membrane antigen 1 gene; AMA1) in *P. falciparum*.
 - How did you navigate to this gene? What other ways could you get there? (hint: what about using the gene ID? PF3D7_0417200 for DHFR-TS or PF3D7_1133400 for AMA1)

- What chromosome is this gene on?
- How many exons does this gene have?
- How many nucleotides of coding sequence?
- b. What genes are located upstream & downstream of DHFR-TS (AMA1) in *P. falciparum*?
 - Is synteny (chromosome organization) in this region maintained in other species? (hint: look in the genomic context section of the gene page).
 - How complete is the genome assembly for other species? (hint: in the genomic context section, each genome is displayed as genes and the underlying contig. Are all the contigs complete?)
- c. Does the *P. falciparum* DHFR-TS (or AMA1) gene contain any single Nucleotide Polymorphisms (SNPs)? (hint: SNPs are represented graphically in the genomic context section and also in a table called "SNP Overview").
 - What is the total number of SNPs in the gene?
 - What do the different color diamonds represent? (hint: mouse over the diamonds to get more information).
 - How many impact the predicted protein sequence?
 - What is the maximum number of SNPs per strain?
 - Is this likely to define the full spectrum of sequence variation in these particular strains?
 - How do these results compare with SNP distribution in other genes? (hint: look at SNPs in upstream and downstream genes).

d. Is the DHFR-TS (or AMA1) gene expressed?

Hint: look at the gene page sections entitled "Protein" and "Expression" - you may have to click on the show link to reveal the underlying data).

- What kinds of data in PlasmoDB provide evidence for expression?
- At what life cycle stage is DHFR-TS (AMA1) most abundant?
- Does this make sense?
- How do the different life cycle microarray expression profiles compare to each other?
- Are the results similar? What about RNA-sequence data, does it agree with microarray data?
- How abundant is DHFR-TS (AMA1) protein? How confident are you of this analysis?

1.4 Finding a gene by BLAST.

Note: For this exercise use <u>http://www.toxodb.org</u> Also, you need to open another window in your browser to retrieve the sequence we will use in our BLAST search, got to:

http://goo.gl/6BGjZ

- Imagine that you generated an insertion mutant in *Toxoplasma* that is providing you with some of the most interesting results in your career! You sequence the flanking region and you are only able to get sequence from one side of the insertion (the sequence in the link above). You immediately go to ToxoDB to find any information about this sequence. What do you do?
- Try running a BLAST search with this sequence (hint: you can get to the BLAST tool by clicking on the BLAST link under tools on the home page).



- Which blast program should you use? (hint: try different combinations, just keep in mind that you have a nucleotide sequence so you have to use an appropriate BLAST program).

Note on BLAST programs:

- blastp compares an amino acid sequence against a protein sequence database;
- blastn compares a nucleotide sequence against a nucleotide sequence database;
- blastx compares the six-frame conceptual translation products of a nucleotide sequence (both strands) against a protein sequence database;

- tblastn compares a protein sequence against a nucleotide sequence database dynamically translated in all six reading frames (both strands);
- tblastx compares the six-frame translations of a nucleotide sequence against the six-frame translations of a nucleotide sequence database.



- Are you getting any results from blastx? tblastn? What about blastn?
- What is your gene? (hint: after running a blastn against *Toxoplasma* genomic sequence, click on the "link to the genome browser". In the genome browser zoom out to see what gene is in the area).

1.5 More BLASTing in EuPathDB (optional). Note: for this exercise use <u>http://www.eupathdb.org</u>

- a. The first thing we will need to do is get a sequence to use for BLAST. Search for the keyword "dihydrofolate" (without quotations). (Hint: use the Gene Text Search on the upper right hand side of the EuPathDB home page).
- b. You should get multiple hits. Find the first one that is annotated as "dihydrofolate reductase-thymidylate synthase" (Look in the product description column).
- c. Once you find one, click on the gene and go to the gene page. It might be helpful to open the gene page in a new window or tab.
- d. Scroll down to the bottom of the page to the "Sequences" section.
- e. Copy the amino acid sequence and go back to EuPathDB (if you have not done so already, it might be helpful to open EuPathDB in a new window or tab).
- f. Go to the BLAST page from the EuPathDB home page. (hint: under Tools on the EuPathDB home page).
- g. Paste the amino acid sequence into the input window.
- h. Select target data type (start with "Proteins").
- i. Select BLAST Program. (Hint: BLASTP).
- j. Select the target organism (let's start by selecting all, but you may need to exclude Amoebazoa). Click on "Get Answer".

Based on the results you should have identified excellent hits in almost all pathogens in EuPathDB but can you find good hits in *Giardia*, *Entamoeba* or *Trichomonas*?

Let's try a different BLAST method:

- k. Go back to the BLAST window. Change the target data type to Genome.
- I. Select the BLAST Program. Notice you cannot select BLASTP anymore. Try the other options. Notice how your input sequence type has to change when you select a different program. (Hint: TBLASTN is the one you need).
- m. Select all target organisms. Click on "Get Answer".

Note that the results are still missing a dihydrofolate from *Giardia* and *Trichomonas and Entamoeba*. *Let's try a different BLAST method.*

n. Go to your gene page window (in CryptoDB) and copy the nucleotide coding sequence.

- o. Go to the BLAST window and paste the nucleotide sequence into the input window.
- p. Select the target data type (try different ones) and the BLAST program. Notice you can only select TBLASTX or BLASTN when your input sequence is nucleotide. (Hint: select TBLASTX).
- q. Select the target organisms. This time let's specifically only select *Giardia*, *Entamoeba* and *Trichomonas*. Click on "Get Answer".

Getting frustrated?

Not getting a hit for *Giardia*, *Entamoeba* and *Trichomonas* in this case is actually the correct answer! These organisms do not have dihydrofolate reductase or thymidylate synthetase activity.