Sequencing project for Bi1x – class exercise

Overview

The main objective of this exercise is to show you a simplified version of the way microbiologists identify small subunit ribosomal RNA sequences ("16S" sequences) from the environment. You will learn how to (1) convert your sequencing traces to nucleotides (2) align the 16S sequences using the greengenes website (3) scan your sequences for chimeras, which you will eliminate from your database and (4) use MEGA to produce a phylogenetic tree of your sequences and their nearest neighbors (found by greengenes). You will then measure in MEGA the distance between your 16S sequences and the nearest neighbors and determine if they are of the same species, genus or phylum. Finally you will identify the precise taxonomic classification of the nearest neighbor for each of your 16S sequences and explain if given what's known in the literature about the physiology of these isolates it's plausible that you found these microbes in a fresh water pond (or could it be a contamination?). Please follow the instructions below and answer the questions in the bullets as you go along.

Computer requirements: to complete this exercise you will need a (1) PC or a PC emulator on a Mac and (2) internet access.

Part I: Learn a little more about the 16S gene

The *rrsA* gene of *E. coli* codes for a small subunit ribosomal RNA (rRNA) gene. Let's learn a little more about this gene. Go to the BioCyc microbial database (the default organism being *E. coli*)

http://ecocyc.org/

In the search box type *rrsA* and click **rrsA** under **genes**.

- 1. What is the length of this gene?
- 2. Since you used 'all bacterial primers' for your PCR your primers should also match this rRNA gene. Find out the length of the amplicon you generated. The primer sequences you used were

```
357F (16S forward primer) 5' CTCCTACGGGAGGCAGCAG 3' 492RL2D (16S reverse primer) 5' TACGGYTACCTTGTTACGACTT 3'
```

Here is a tool to help you reverse complement a nt sequence http://www.geneinfinity.org/sms reverse complement.html

Hint 1. to get E. coli's 16S sequence click on the rrsA hyperlink under Genes and then press the Nucleotide Sequence button. To convert this sequence into a long string of letters copy this sequence into notepad in order to remove non-text characters. Then copy this sequence into Word and replaces spaces "with an empty string and then replace the newline character "p" with an empty string.

Save E. coli's sequence for later, we will be using it again for our tree.

Hint 2. What is special about the nucleotide sequence of one of the primers above? Look at them carefully...no, it's not a mistake...This website can perhaps help you: http://www.le.ac.uk/bl/phh4/nucredun.htm

- 3. What is the genomic context of this gene? Click on the **Genome Browser** button. You will see that *rrsA* is part of an operon. What other genes are present in this operon? What do they do? Is their function related to *rrsA*?
- 4. Does *E. coli* have more than one copy of this gene? In the EcoCyc website go to **search**|**BLAST** and BLAST the nucleotide sequence of the rrsA gene against *E. coli*'s genome (the default database). How many hits do you get? What is the name of the homologous genes that you found? Why do you think *E. coli* should have multiple copies of this gene?
- 5. How about other organisms? Do other organisms have multiple copies of this gene? What is range of copy numbers in nature? To find out the answer take the forward primer above and BLAST it against other organisms. You can change the organism database in BioCyc by clicking on the **change** hyperlink at the upper right corner. Try this for a few prokaryotes, including the spore forming bacterium *Bacillus subtilis* (a different strain of which causes anthrax), the highly abundant cyanobacterium *Prochlorococcus marinus*, the human pathogen *Streptococcus pyogenes* and finally the extreme thermophile *Thermococcus kodakarensis*. *Hint:* don't forget to set BLAST to compare a nucleotide sequences with a nucleotide database. *Bonus question:* You should have failed to get an answer for one of these prokaryotes. Which was it? Can you give a reason why? Now double check your results using the ribosome RNA copy number database (rrnDB) (also for the prokaryote that failed this test).

http://ribosome.mmg.msu.edu/rrndb/search.php

Just type the name of the prokaryote in the **keyword** box.

6. How do you think the rRNA copy number can affect the PCR reaction you carried out? Do all prokaryotes get equal representation? If all bacteria had the same rRNA copy number per genome, would then all prokaryotes get equal representation?

Part II: Align your SSU rRNA sequences and find nearest neighbor matches

1. Greengenes tutorial. You will use an online tool called greengenes hosted by Lawrence Berkeley National labs to align your 16S sequences and find nearest neighbors. Get acquainted with this tool by reading the online tutorial.

http://greengenes.lbl.gov/cgi-bin/JD Tutorial/nph-Tutorial 2Main2.cgi

2. Convert you abi trace files to nucleotides. Most sequencing reactions produce ~700-900 basepairs of high-quality sequences (called "reads"), which is not enough cover the entire 16S amplicon. To sequence your entire amplicon we would need to run two sequencing reactions in opposite directions using both the forward and reverse primers and then "assemble" the two reads. However the read length that we obtain with just one sequencing reaction is long enough for us to get reasonable classification of the organism. The first step however is to obtain quality nucleotides from your traces. To accomplish this download and install Sequence Scanner to your PC after filling out a quick form

http://marketing.appliedbiosystems.com/mk/get/SSS_login?isource=fr_E_Pg_Prod AB Gbl SeqScan 20050920

We will explain in class how to use this tool. Generate nucleotide sequences for all your 16S amplicons. Try to get the longest sequences that are still of high quality (that is, you can easily identity the base call in the chromatogram).

3. Generate your FASTA files. Save the nucleotide sequence of each trace in a separate FASTA file. A FASTA file is a text file with the following simple format:

```
>SEQ_NAME
SEOUENCE...
```

The format is not case sensitive, but use only alpha-numeric characters and avoid spaces.

4. Chimeras. When running your PCR reaction there is the chance that two different PCR products will combine to form a chimera amplicon, which of course is an unwanted artifact that needs to be identified and discarded. There are several online tools to help you identify these artifactual sequences. We will be using a tool called Bellerophon. Read more about chimeras and how Bellerophon works in this link

http://foo.maths.uq.edu.au/~huber/doc/doc.html

Submit your FASTA file to the Bellerophon server to check for chimeras:

http://foo.maths.uq.edu.au/~huber/bellerophon.pl

Do you have any chimeras? Eliminate any chimeras from analysis.

5. Align sequences. Next you will use greengenes to align your 16S sequences. Alignment of 16S sequences is different than alignment of protein coding nucleotide sequences because the 16S aligners take into account the secondary structure of the 16S gene. For example, sequences on both sides of a stem loop need to match up in order to form the stem loop. Greengenes also attempts to align your sequences against a very large database of pre-aligned 16S sequences and will include in the results the nearest neighbors it could find (i.e. other 16S sequences with the highest similarly to your sequence). You can also specify if you want greengenes to return any nearest neighbors (which include uncultured organisms) or nearest neighbors which are isolates (and thus better characterized). We will choose both. The nearest neighbor sequences will be returned in separate FASTA files (see below). To proceed, load your FASTA files one at a time to the greengenes aligner (Align icon) using the settings below:

http://greengenes.lbl.gov/cgi-bin/nph-NAST align.cgi

- ➤ My Taxonomy (lower right)— activate ncbi taxonomy (choosing all options) → make changes to my interest list → click on the Align icon again
- ➤ Minimum length = 500bp (this should be lower than the length of your submitted sequence)
- ➤ Check options 1,2, 4 and 5 in "Files you desire"
 - o xyz NAST.xls (greengene alignment report)
 - o xyz NAST.fasta (your aligned sequences)
 - o xyz nn NAST.fasta (nearest neighbors)
 - o xyz nni NAST.fasta (nearest isolates)
- > Set the number of sequences to be returned to 5
- Leave the rest of the parameters to their default values
- ➤ Enter your email address and hit **proceed.** Progress is updated on your web browser.
- ➤ Use WinRAR or WinZip to open the files on your PC if they do not open directly. Download WinRaR from here WinRAR

http://download.cnet.com/WinRAR-32-bit/3000-2250 4-10007677.html

- After you obtain the results, based on the tutorial, inspect the excel file to see which sequences were aligned and identified.
- **6.** Create your MEGA FASTA file. Combine your xyz_NAST.fasta file with your xyz_nni_NAST.fasta file and your xyz_nn_NAST.fasta files (do this for each of your 16S pond sequence separately since these sequences can be quite divergent

and the treeing algorithm assumes the sequences are similar). You may want to add a header to the sequence name to distinguish nni from nn sequences. This will be your new FASTA file. We will next use MEGA, which is a little fussy about the input format of the FASTA file. To convert the greengene FASTA file to a format MEGA can read do the following:

- ➤ Replace spaces ("") and any "=" symbols with underscores ".
- Replace '.' symbols with dashes '-'.
- Make sure the extension of the FASTA file is 'fasta'. To changed an extension of text file on a PC use Notepad and in "save as" change the "save as type" from "Text Documents" to "All Files" and use the file extension 'fasta'.

Part III: Phylogenetic analysis

1. Phylogenetic analysis. We would like to generate a Neighbor Joining tree for each of your 16S sequences and its nearest neighbors. This will allow you to visualize the evolutionary distance between each of your sequences and their nearest neighbors. We will also calculate precisely the similarity between these 16S sequences. Based on this percent similarity you can determine the degree of certainty in your taxonomic classification. Since the 16S gene is highly conserved, mutations correlate with time, and so the amount of sequence divergence correlates with evolutionary separation. For example, two 16S sequences having ≥97% identity are generally considered to belong to the same species, ≥95% identity the same genus and ≥80% identity the same phylum. Based on these empirical rules you can determine if your nearest neighbor is the same species, genus or phylum as the 16S sequence you found.

Download and install on your PC a program called MEGA4 from the following link

http://www.megasoftware.net/mega.html

(Mac users – you will need to run a virtual PC to run this program or work on one of the PCs in the lab)

2. Constructing a phylogenetic tree in MEGA.

Open MEGA and drag your corrected FASTA file into MEGA's main window (turquoise window). Now export the file in MEGA format

Data \rightarrow export \rightarrow follow instructions \rightarrow protein coding data? select NO

Close the file and you will be prompted by MEGA to open the MEGA file – click yes. If not, just open the MEGA file you just saved. Next, build a NJ tree with bootstrapping using a Jukes-Cantor nucleotide substitution model. Go to MEGA's main window and then choose:

- **>** Phylogeny→ construct phylogeny → Neighbor Joining (NJ)
- ightharpoonup Test phylogeny tab \rightarrow select bootstrap (1000 replicates)
- ➤ Options summary tab → select Substitution Model→Model→Nucleotide → Jukes-Cantor
- ➤ Hit compute
- ➤ Hit the "caption" button on the tree you just created to see the precise definition of this tree. How many nucleotide positions were used in the tree? You should get at least 500bp, otherwise something maybe wrong ask one of your TAs.

You should obtain a tree that looks something like Fig. 1. Here are some assignments and questions for you regarding your trees (answer for each tree you generate)

- a. Draw your NJ tree using two topologies: rectangular and radiation. What is the difference between these two topologies? (submit only the rectangular tree)
- b. Why is the Jukes-Cantor model more accurate than simply using the pdistance to calculate evolutionary distances? What does the Jukes-Cantor model attempt to correct for?
- c. Do the bootstrap values support your classification?
- d. The tree that you submitted is an unrooted tree. What kind of organism would you need to root this tree? What would you gain by rooting the tree?
- e. Use MEGA to find the percent identity between your sequence and its nearest neighbors. To calculate distances in MEGA, from MEGA's main window choose

Distances \rightarrow compute pairwise \rightarrow Substitution Model \rightarrow Model \rightarrow Nucleotide \rightarrow p-distance

See Fig. 2. for an example how this output looks like.

Hint: you can drag up and down the sequences in the resulting distance matrix window to help you located your nearest neighbors.

- f. Who is the nearest 16S sequence to each of your pond sequences? What is the percent identity to that sequence? Is the nearest neighbor the same species, genus or phylum as your 16S? Can you use the nearest neighbor to classify your organism reliably?
- **3.** Classify your 16S sequences. Obtain further NCBI taxonomic information about the nearest neighbor to your 16S sequence (preferably an isolate) by searching its NCBI ascension number in the NCBI nucleotide database

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

Hint: the NCBI ascension number appears in the organism name given by greengenes and is the second field after the first underscore). See an example in Fig. 3.

- a. What is the precise taxonomic classification of your nearest neighbors?
- b. Read up on the microbes you identified (e.g. by looking up the references in the NCBI record, searching Google scholar for the microbe's name, searching WikiSpecies, etc.). Given this organism's physiology (respiration method, energy and carbon sources, typical osmolarity/pH/temperature ranges) is it plausible you found it in a fresh water pond or could it be a contaminating sequence in your PCR?

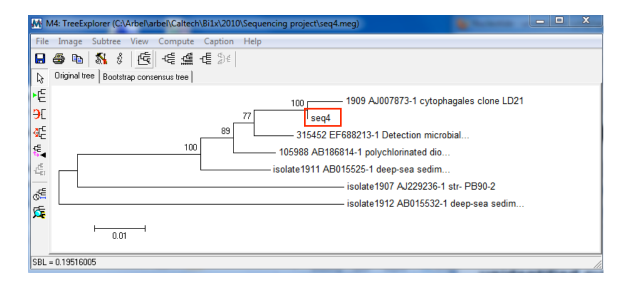


Figure 1. Example of a NJ tree with 1000 bootstrap replicates generated by MEGA to identify seq4. There were a total of 726 positions in the final dataset.

(A,B) (A,B	File Display Average Caption Help								
1. seq4 2. isolate1907 AJ229236-1 str- PB90-2 0.0937 3. isolate1911 AB015525-1 deep-sea sedim 0.0344 0.0854 4. isolate1912 AB015532-1 deep-sea sedim 0.0992 0.1047 0.0923 5. 1909 AJ007873-1 cytophagales clone LD21 0.0063 0.0978 0.0413 0.1061 6. 105988 AB186814-1 polychlorinated dio 0.0220 0.0895 0.0262 0.0937 0.0289	(A,B) 0.00								
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	7. 315452 EF688213-1 Detection microbial	0.0179	0.0868	0.0331	0.0937	0.0248	0.0207		7

Figure 2. p-distance between seq4 and its nearest neighbors. The minimum distance is with respect to the cytophagales clone (99.3% identity).

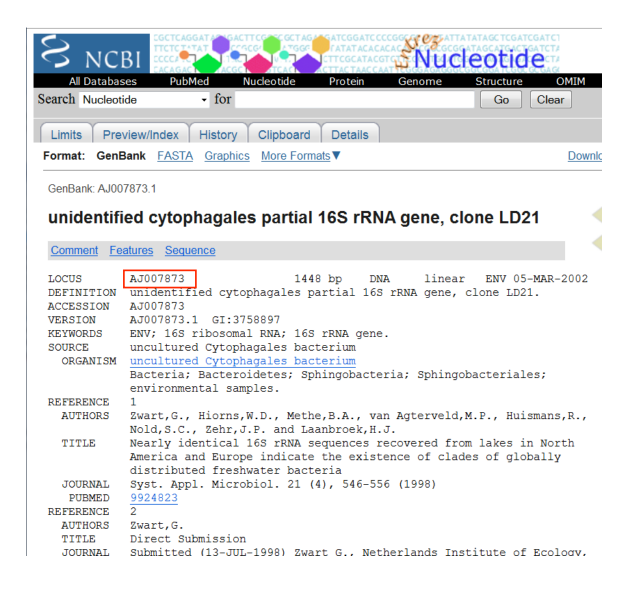


Figure 3. NCBI recored for cytophagales clone (AJ007873-1)