## Unix, Perl and BioPerl

Session 1: Introduction to Unix for Bioinformatics

## Exercise 1: BLASTing ESTs against a RefSeq database

Goal: Learn the most common Unix commands while manipulating sequence files and "identifying" some rat ESTs by searching RefSeq, an annotated database, with BLAST.

Note: Each command written on multiple lines should be entered as a one-line command, except for actual multiple-line commands, which are delimited with semicolons.

See http://jura.wi.mit.edu/bio/education/bioinfo2005/unix-perl/for course page

#	To do / To answer	Command	Comments
0	Mac OS X: Open the terminal on your computer Mac OS 9: open MacSSH Windows: Open SSHSecureShellClient (See http://jura.wi.mit.edu/bio/education/docs/ssh-sftp.html		If you're running Mac OS X, you're in Unix now.
	for more information about		
1	Open your home account on hebrides.	ssh -1 username hebrides.wi.mit.edu  1st time users: passwd	Username is replaced by your's. You will be prompted for your password.  If it's the first time for connecting, change your password.
2	What is the full path to your home directory?	pwd	"print working directory"
3	What files are in your home directory?	ls	"list" – won't show hidden files
4	What files (including hidden files) are in your home directory? How big are they?	ls -a; ls -al	-a option will also show files starting with a dot
5	What's in these files?	more myfile	myfile is replaced by a name you got with the "ls" command
6	Create a directory called "unix_class"	mkdir unix_class	"make directory"
7	Go to the "unix_class" directory	cd unix_class	"change directory"

8	Make directories called	mkdir rat-ests;	
	"rat-ests" and "dbs", and	mkdir dbs; cd dbs	
9	go to the "dbs" directory.		Note: Comptimes ETD
9	Access the NCBI by FTP	ftp ftp.ncbi.nih.gov	Note: Sometimes FTP
	to get the rat RefSeq	  [Follow instructions	requires a specific user name
	sequence "database" (which isn't really a	from the FTP site.	and password, and other times FTP access can be
	database but rather a	Note: This version of	anonymous (in which case
	multiple sequence file)	the 'ftp' command is	you use "anonymous" as the
	maniple sequence me)	called 'ncftp'.]	username and your e-mail as
			the password)
10	Go to the "refseq"	cd refseq	
	directory		
11	List the refseq files. Get	ls;	README is often a very
	the README file,	get -z README	helpful file on FTP sites. The
	renaming it as	refseq_README	'-z' is not necessary in some
	refseq_README		versions of FTP.
12	Go to ./R_norvegicus/	cd R_norvegicus;	
	mRNA_Prot	cd mRNA_Prot	
13	Get the file "rat.fna.gz"	get -z rat.rna.fna.gz	The '-z' is not necessary in
	containing the RefSeq set	rat.fna.gz	some versions of FTP. The
1.4	of rat cDNA sequences		opposite of "get" is "put"
14	Disconnect from the FTP	quit	or "bye"
1.5	Site	1 ~	V1111
15	Check to make sure you	ls	You should have
	downloaded what you		'refseq_README' and 'rat.fna.gz'
16	wanted to get Look at the README	more refseq README	Hit the space bar to advance
10	one screenful at a time		to the next screenful or
	one sereemar at a time		'q' to quit 'more'
17	Unzip the sequence file.	gunzip rat.fna.gz;	It's generally assumed that a
1,	What's the file called	ls	file ending in .gz needs to be
	now?		unzipped; the opposite is gzip
18	How big is the sequence	ls -l	11 / 11
	file?		
19	How would you list files	man ls	Extra credit: How would you
	in order of modification		list in reverse order of
	time? (Consult the man		modification time (from
	pages for ls, using the		oldest to newest)?
	space bar to advance and		
20	"q" to quit)		TA CTA C
20	Look at rat.fna to check	more rat.fna	FASTA format is a one-line
21	that it's in FASTA format	a non	header followed by sequence
21	What are the arguments	grep	"general regular expression
	to use for "grep"?		parser" – very useful!
<u></u>			

22	Use grep to print all the header lines into a file called rat.headers	<pre>grep "&gt;" rat.fna &gt; rat.headers</pre>	">" marks the beginning of a fasta file	
23	Check out the new file to be sure it looks okay at the beginning and the end	head rat.headers; tail rat.headers	Add the option –n to print n lines with "head" or "tail"	
24	How many sequences are in the sequence file?	<pre>grep "&gt;" rat.fna   wc -1</pre>	wc –l ("word count") actually prints the number of lines	
25	Make a BLAST database of the rat.fna sequence file using the "formatdb" command.	formatdb -i rat.fna -p F -o T	"formatdb —" prints all options. The options used here are the minimal/usual ones.	
26	What files have been created? What does the log file say?	ls; more formatdb.log	formatdb.log will show any formatdb errors.	
27	Change to the 'rat-ests' directory	cd/rat-ests	Remember that '' means up one level in the directory tree	
28	Get a file of ESTs from /home/george/rat-ests and place it into the directory "rat-ests"	cp /home/george/rat-es	sts/* .	
29	Extract the first sequence and place it into a file by itself.	<pre>head ests.fa; head -8 ests.fa &gt; est1.fa</pre>	Can you think of another way to do this?	
30	Run BLAST on the single sequence, using an expect cutoff of 0.5, printing text output (only best 5 hits)	blastall -i est1.fa -d/dbs/rat.fna -p blastn -e 0.05 -v 5 -b 5 -o est1 blast.txt	"blastall" command shows and describes all options. What do these options mean?	
31	Run a similar BLAST search but with a default expect value cutoff and generate tab-delimited output	blastall -i est1.fa -d/dbs/rat.fna -p blastn -v 5 -b 5 -o est1_blast_tab.txt -m 8	Similar to the command above, but with "-m 8" added. Did you use the ↑ command to get back to the previous one?	
32	Open est1_blast.txt in pico (a simple text editor), using ^X (control-x) to exit.	pico est1_blast.txt	Try running BLAST on est1.fa with "-m 9" instead of "-m 8" to find what each field is showing.	
33	Extract a sequence (ex: NM_199463) from the BLAST database	fastacmd -s NM_199463 -d/dbs/rat.fna	This only works if you had used the "-o T" option with formatdb	

34	Make a file with the five sequence IDs from est1_blast.txt, and extract these sequences from rat.fna	<pre>fastacmd -i list.txt -d/dbs/rat.fna &gt; myseqs1.fa</pre>	Use pico to make the list. Accessions (ex: NM_133594) or GIs (ex: 19424297) can be used.
35	BLAST the set of ESTs with standard text output	blastall -i ests.fa -d/dbs/rat.fna -p blastn -e 0.05 -v 5 -b 5 -o est_blast_1.txt	Note that BLAST is very fast when searching a database smaller than the default "nt" database
36	BLAST the set of ESTs with tab-delimited output	blastall -i ests.fa -d/dbs/rat.fna -p blastn -e 0.05 -v 5 -b 5 -T F -m 8 -o est_blast_tab.txt	
37	Any questions?		
38	Logout from hebrides and return to your desktop terminal.	logout	Make sure the "command prompt" no longer shows something like "username@hebrides".
39 a	If you're using Mac OS X, download the tabdelimited output using sftp and view in it Excel.	sftp username@hebrides.wi. mit.edu	Replace "username" with yours. Use cd, ls, and get as before for ftp.
39 b	If you're using Mac OS 9 or Windows, use an application to download the tab-delimited output using sftp and view in it Excel.	[Follow instructions for Fetch or Mac SFTP.]	
40	Delete any of your files from the laptop		Thanks!

## Notes:

1 - In any cases of poor FTP connections to NCBI,
rat.fna.gz and the associated README
can be copied from
/home/george/unix/dbs/

2-If, when trying to use the 'pico' text editor, you get a message that hebrides or barra doesn't know anything about your terminal, use the command setenv TERM vt100 and try again with pico.