# Intro to the Command Line

J Fass | 19 June 2017

### The Unix Philosophy

Small, sharp tools ... each of which does a well defined job very well.

```
jfass@nickel:~$ ssh cabernet
 HPC-14 currently consists of 66 nodes \
 with 2248 CPUs and 11.03TB RAM. Type
 sinfo for more info.
Ubuntu 14.04.2 LTS 4.4.0-67-generic x86 64
***** 2016-05-11 *****
Memory has been enabled as a consumable resource. If your job will be
using more than 2G/CPU, you will need to use --mem-per-cpu= in your job
scripts or your job will be automatically killed by the scheduler.
***** 2017-03-03 *****
As of March 13 2017 we will be requiring the --time option for all jobs.
Shortly thereafter we will be reducing the grace period from 8 days to
1 hour. Contact sysadmin@genomecenter.ucdavis.edu with any questions.
Last login: Wed Jun 14 09:35:04 2017 from nickel.genomecenter.ucdavis.edu
groups: cannot find name for group ID 1103844286
jfass@cabernet:~$
```

```
jfass@nickel:~$ ssh cabernet
                    .genomecenter.ucdavis.edu
[\ldots]
jfass@cabernet:~$
```

```
jfass@nickel:~$ ssh cabernet
                    .genomecenter.ucdavis.edu
[ \dots ]
jfass@cabernet:~$ <ctrl>-1
```

```
jfass@cabernet:~$
```

<ctrl>-l or -k ...
Clears terminal, start at top

See also: reset

```
jfass@cabernet:~$
```

```
<-- prompt (includes $ and one
space after)

Huge # of possible
configurations; in this case:

<uname>@<hostname>:<pwd>$<space>

(pwd = present working directory)
```

#### Command Line Basics - Don't work on head node!

```
jfass@cabernet:~$ srun -t 1440 -n 1 --mem 8000
--reservation workshop --pty /bin/bash
srun: job 5461103 queued and waiting for resources
srun: error: Lookup failed: Unknown host
srun: job 5461103 has been allocated resources
jfass@c4-1:~$
```

This logs you into an interactive session on one of the cluster nodes, so you don't all play on the head node.

More on this later. Just log into cabernet, then enter the above and press <enter>.

From now on, I'll omit "jfass@hostname:" for brevity!

#### **Command Line Basics**

```
~$ <type command here>
~$ pwd<enter>
/home/jfass
```

Follow command with <enter>

E.g. "pwd" ... lists your present working directory

```
~$ command fubar <control-c>
~$ # looks like this:
~$ command fubar ^C
~$ sleep 1000
^C
```

```
<control-c>: escape from
entering a command ...
```

```
(notice: "#" prevent
interpretation of text that
follows)
```

... kill a running command
("sleep" actively counts off
the specified number of
seconds before letting you do
anything else)

```
~$ R
[\ldots]
> <control-d>
Save workspace image? [y/n/c]: n
```

```
<control-d> ... escape from
some interactive sessions (R,
python, ...)
```

(R is a powerful
data-centered, statistical
computing language)

```
~$ yes
          more
[ \dots ]
<q>
```

```
q = quit:
Escape from paginators!
(less, man, etc.)
("yes" says "y" until killed
... it's a dinosaur)
("|" is the pipe character ...
we'll explore it more soon)
("more" shows you a page of
text, then waits for you to
hit <space> to show another,
or exit with "q")
```

~\$ exit

"exit" kills the current shell: the program that's interpreting your commands for the operating system.

#### Command Line Basics - Where Am I?

```
~$ 1s
[\ldots]
~$ pwd
[\ldots]
```

list file in the pwd

present working directory

### **Command Line Basics - Options!**

```
~$ 1s -R
[\ldots]
[\ldots]
<control-c>
```

```
list recursively
```

What did I just do???

## Command Line Basics - Read The Manual (RTM)!

```
\sim$ man 1s
[\ldots]
<up, down arrows>
[\ldots]
<q>
```

man <command> consults the manual that exists for basic, OS commands. Any software author can write a "man page" for their software, but most scientific software authors don't.

## Command Line Basics - Options, options, options!

```
~$ 1s -1
~$ ls -a
~$ ls -1 -a
~$ ls -la
~$ ls -ltrha
```

man 1s ...

Can combine single letter options ...

list all files (in pwd), in
long format, in reverse time
order with human readable
file sizes

### Command Line Basics - Options, options, options!

```
~$ alias
[\ldots]
alias ll='ls -alFtrh'
[\ldots]
~$ 11
[\ldots]
```

"alias" allows you to define your own commands. "alias" by itself only *lists* the aliases defined in your bash configuration files. Try one on for size ...

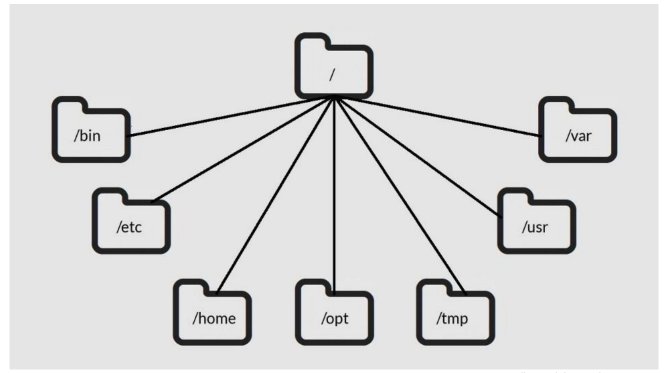
### Command Line Basics - Directory Structure

```
~$ 1s -R
[\ldots]
./R/x86 64-pc-linux-gnu-library/3.3/BH/include
[\ldots]
<control-c>
```

'/' separates directories

Names can include many characters, but avoid spaces and other weird stuff.

### Command Line Basics - Directory Structure



www.linuxtrainingacademy.com

#### Command Line Basics - '.' and '..'

```
~$ ls -a
.bash history
[\ldots]
```

```
"." = pwd
```

".." = up one level

Don't be confused between use of "." and filenames that start with "." ... the latter are valid filenames, that are just "hidden" unless you use the "ls" command's "-a" option.

#### Command Line Basics - Absolute/Relative Address

```
~$ ls /home/jfass/
[\ldots]
~$ 1s ./
[same ...]
~$ ls ../jfass/
[same ...]
~$ ls ../jfass/../jfass/../jfass/
[yup, same ...]
```

```
"." = pwd
".." = up one level
```

### Command Line Basics - <Tab> Completion

```
~$ ls /home/jfas<tab>
~$ ls /home/jfass/
```

<tab> will literally save
your life. Hours of it.

A single <tab> auto-completes when it's possible (when only a single possible completion exists).

### Command Line Basics - <Tab> Completion

```
~$ ls /home/j<tab>
~$ ls /home/j
~$ ls /home/j<tab>
jacob/ [...]
jagadish/
jagomez/
                                             jwbucha/
[\ldots]
```

<tab> will literally save
your life. Hours of it.

Two <tab>s in a row will show you all the possible completions, when there wasn't a *single* one for the single <tab> to use.

# Command Line Basics - <Tab> Completion

```
~$ ls /h<tab>
~$ ls /home/<tab>
~$ ls /home/<tab>
~$ ls /home/j<tab>
~$ ls /home/j<tab>
~$ ls /home/jf<tab>
~$ ls /home/jfass/
```

Use it!

Watch out for RSI ...

## Command Line Basics - Create and Destroy

```
~$ mkdir temp
~$ cd temp/
~/temp$ echo "Hello, world!" > first.txt
~/temp$ cat first.txt
~/temp$ rm first.txt
~/temp$ cd ../
~$ rmdir temp
```

Create a directory

Change directories

Put text into a file

Concatenate file to screen

Remove file

Up and out

Remove (empty) directory

### Command Line Basics - Pipe and Redirect

```
~$ mkdir CLB; cd CLB/
~/CLB$ echo "first" > test.txt
~/CLB$ echo "second" > test.txt
~/CLB$ cat test.txt
~/CLB$ echo "third" >> test.txt
~/CLB$ cat test.txt
```

```
">" redirects the output from
one command to a file,
instead of the screen.
```

">" replaces

">>" appends

#### Command Line Basics - Pipe and Redirect

```
~/CLB$ cut -c 1-3 test.txt
~/CLB$ cat test.txt | cut -c 1-3
~/CLB$ cat test.txt > cut -c 1-3
```

"cut" cuts lines of text.

"|" pipes output from one command to be the input of another command.

">" is wrong here ... what does this command do?

### Command Line Basics - Pipe and pipe and pipe ...

```
~/CLB$ cat test.txt
~/CLB$ cat test.txt | cut -c1-3
~/CLB$ cat test.txt | cut -c1-3 | grep s
```

Pipes allow us to build up compound operations, filtering and changing data as we go.

("grep" searches for matches to regular expressions ... patterns)

#### Command Line Basics - History

```
~/CLB$ history
~/CLB$ history | head
~/CLB$ history | tail
~/CLB$ history | tail -n 15
~/CLB$ history | less
```

Since we often develop long commands through trial and error, it helps to see and access what we've done.

In "less," up and down arrows, pgUp, pgDn, and "q" to exit. Also, "/pattern" searches for pattern each <enter>. "n" and "N" for next and previous match, "g" and "G" for beginning and end of file / stream.

#### **Command Line Basics - History**

```
cat test.txt | cut -c1-3 | grep s
 558
      history
 559
      history
 560
~/CLB$ !560
```

"!#" repeats command # from your history.

#### Command Line Basics - History Search

```
~/CLB$ <control-r>
(reverse-i-search)`first': echo "first" > test.txt
```

<control-r>text triggers a
recursive search for "text"
from your history. After
finding the most recent match
that you like, use
<control-r> again to get to
an earlier match (and again,
and ...).

<enter> executes the command;
left or right arrow fills the
command line with the command
but allows you to edit it
before running it.

### Command Line Basics - History Search

```
~/CLB$ <up> <dn> ...
~/CLB$ cat text.txt | grep s
 <control-a>
                        <control-e>
```

And, by the way, the up and down arrows take you backwards and forwards through your history of commands. Reach one you like, and start editing.

Also, by the way, <control-a> and <control-e> bring you to the beginning and end of your command.

## Command Line Basics - Editing Commands

```
~/CLB$ blah blah blah
~/CLB$ blah blah <control-k>blah
~/CLB$ blah blah
~/CLB$ blah blah <control-w>
~/CLB$ blah
```

Left arrow to before the last "blah," then <control-k> ... kills text from here 'til the end of the line.

Now, <control-w> ... kills preceding word.

### Command Line Basics - Compression

```
~/CLB$ gzip test.txt
~/CLB$ file text.txt.gz
~/CLB$ gunzip test.txt.gz
~/CLB$ bzip2 test.txt; bunzip2 test.txt.bz2
```

Compress big files using "gzip," "bzip2." Bzip2 compresses smaller, but takes longer.

("file" gives you info about the *type* of file you're looking at)

#### **Command Line Basics - Archives**

```
~/CLB$ wget
ftp://igenome:G3nom3s4u@ussd-ftp.illumina.com/PhiX/I
llumina/RTA/PhiX Illumina RTA.tar.gz
~/CLB$ tar -xzvf PhiX Illumina RTA.tar.gz
```

Large directory trees may be compressed as "tarballs" ... see "tar."

Let's grab one and expand it.

### Command Line Basics - Forced Removal



This is a dangerous one.
Remove a file / directory, do
it recursively to all
sub-directories, and force
removal (by-pass confirmation
questions).

Caution is warranted. There's no Trash Bin, and no gauranteed way to recover deleted files.

#### Command Line Basics - Wildcard Characters

```
~/CLB$ tar -xzvf PhiX Illumina RTA.tar.gz
~/CLB$ ls PhiX/Illumina/RTA/Sequence/*/*.fa
[\ldots]
```

Let's re-unarchive that tarball, to have something to look at.

List all files a few directories down that end in ".fa" ...

### Command Line Basics - Wildcards and Find

```
~/CLB$ find . -name "*.fa"
[\ldots]
~/CLB$ find . -name "*.f?"
```

"\*" can fill in for anything in a filename, except "/" ... there's a more appropriate command to use when you don't know which directory level the files you're looking for are at: "find"

"?" is like "\*," except only fills in for a single character.

### Command Line Basics - Fasta example

```
~/CLB$ curl -k
https://bioshare.bioinformatics.ucdavis.edu/bioshare
/download/pdhqicmfgw2bra8/variant.neighborhoods.fa >
regions.fa
~/CLB$ wc -1 *
~/CLB$ grep -c ">" regions.fa
```

```
Grab a sequence file from the web ("curl" copies from a url) ... see also "wget"
```

```
"wc" counts words, etc.; "-1" only gives line count
```

```
"-c" causes grep to count matches instead of printing them
```

### Command Line Basics - Fasta example ... Pipes!

```
~/CLB$ grep ">" regions.fa | cut -c 2-
~/CLB$ grep ">" regions.fa | cut -c 2- | cut -f1 -d:
~/CLB$ grep ">" regions.fa | cut -c 2- | cut -f1 -d:
 sort
~/CLB$ grep ">" regions.fa | cut -c 2- | cut -f1 -d:
 sort | uniq -c
~/CLB$ grep ">" regions.fa | cut -c 2- | cut -f1 -d:
 sort | uniq -c | sort -rn -k1,1
```

Follow each step of these commands ... the entire final command counts how many sequences come from each contig.

"sort," "uniq" are self-explanatory; "uniq -c" adds the counts (must be sorted first), and "sort -rn -k1,1" sorts lines in reverse numerical order based only on the first column (the counts).

## Command Line Basics - Symbolic Links

```
\sim/CLB$ ln -s
PhiX/Illumina/RTA/Sequence/WholeGenomeFasta/genome.f
a .
~/CLB$ ls -ltrha
[...] genome.fa ->
PhiX/Illumina/RTA/Sequence/WholeGenomeFasta/genome.f
а
~/CLB$ grep -c ">" genome.fa
```

"ln -s [target] [link location/name]" creates a "shortcut" to the file.

The target file can be deleted, which leaves a broken link. The link can be deleted without affecting the file.

Otherwise, addressing the link addresses the file.

#### Command Line Basics - Bioinformatics At Last?

```
~/CLB$ bwa mem genome.fa regions.fa > aln.sam
[E::bwa idx load] fail to locate the index files
~/CLB$ cat aln.sam
~/CLB$
```

Let's align sequences to a sequence!

Woops ... what went wrong?

I redirected output to a file, why didn't it go there? (Even if I wouldn't want that kind of stuff to end up in an alignment file!)

### Command Line Basics - STDOUT & STDERR

```
~/CLB$ bwa mem genome.fa regions.fa 1> aln.sam 2>
aln.err
~/CLB$ cat aln.sam
~/CLB$ cat aln.err
[E::bwa idx load] fail to locate the index files
```

Programs can write to two separate output streams: standard out, and standard error. Former mostly used for output, latter mostly used for error messages.

("1>" is equivalent to ">")

Now we can store error messages from many jobs run at once, to separate files.

## Command Line Basics - Loops

```
\sim/CLB$ for i in {1..21}; do echo $i >> a; done
for name in {list}; do
    commands
done
~/CLB$ cat a
[...]
```

For loop, for a defined number of steps.

# Command Line Basics - Loops

```
~/CLB$ grep ">" regions.fa | cut -c2- | while read
header; do echo "contig $header" >> b; done
while {condition}; do
    commands
done
~/CLB$ cat b
[\ldots]
```

While loop, for stopping on a condition.

### Command Line Basics - Paste

```
~/CLB$ paste a b
[\ldots]
```

"paste" jams columns of equal size together. See also "join" ...

# Command Line Basics - Running in the Background

```
~/CLB$ sleep 100000
[\ldots]
<control-z>
^Z
[1]+
      Stopped
                                sleep 100000
bg
[1]+ sleep 100000 &
```

When a long job is running, use "<control-z>", then "bg" to suspend it, then run it in the background. If you want to kill it before it finisheds on its own, either "kill %1" (or %2, etc., if multiple jobs), or "fg," then <control-c>.

# Command Line Basics - Running in the Background

```
~/CLB$ sleep 100000 &
[1] 2166
~/CLB$ kill %1
~/CLB$
     Terminated
[1]+
                              sleep 100000
```

To start a job already running in the background, append "%" to the command.

# Command Line Basics - Running in the Background

```
~/CLB$ nohup sleep 100000 &
[1] 2178
~/CLB$ nohup: ignoring input and appending output to
'nohup.out'
~/CLB$ jobs
[1]+
     Running
                              nohup sleep 100000 &
```

To make a background job resistant to a lost connection or terminal problems, use "nohup."

"jobs" will list jobs running in the background.

With nohup, you should be able to exit from the shell, and the job keeps running.

(Not as useful in a cluster environment?)

#### Command Line Basics - Table of Processes

```
~/CLB$ top
                                                             "Top" prints a self-updating
                                                             table of running processes
                                                             and system stats. Try "M,"
                                                             "P," and "u" then uname ...
        top - 14:15:13 up 94 days, 4:52, 35 users, load average: 3.00, 3.00, 3.00
               557
                                      554
                                           62.5
                                   0.0
                                                    37.5
                                                                       0.0
                                                                               0.0
                  12293548
                                 11816988
                                                  476560
                                                                 260124
                   1952764
                                   358824
                                                 1593940
                                                                9564044
          PID USER
                                 VIRT
```

### Command Line Basics - Shell Scripts

```
#!/bin/bash
grep -o . $1 | \
    sort | \
   uniq -c \
    sort -rn -k1,1
<control-o><control-x>
```

"Nano" is a simple text editor; open it with "nano test.sh" and type / copy in this ... your first shell script! Save and exit as shown at the bottom, via <control-o><control-x>.

#### Command Line Basics - Permission Needed

```
~/CLB$ 11 test.sh
-rw-rw-r-- 1 jfass biocore 79 Jun 18 19:00 test.sh
~/CLB$ chmod u+x test.sh
~/CLB$ 11 test.sh
-rwxrw-r-- 1 jfass biocore 79 Jun 18 19:00 test.sh*
```

Though you can run the commands in test.sh in several ways, to really make it a script you need to give yourself permission to execute it.

Permissions now are to read and write for user and group, and only read for the world (ignore first dash, then rwfor user, rw- for group, and r-- for world). Add execute for user as shown.

#### Command Line Basics - Permission Needed

```
~/CLB$ ./test.sh genome.fa
  1686 T
   1292 A
   1253 G
   1155 C
     1 x
     1 p
     1 i
     1 h
      1 >
```

Execute by addressing the shell script with its path, and feed it an argument as required in the script.

Congrats! You've now counted all characters in the phiX genomic fasta file.

Questions ... comments ... confusion?