Unix exercises

Find the number of unique users on a shared system

We know that w will tell us the users logged in. Try it now on a system that has many users (i.e., not your laptop) and see the output. Likely there are dozens of users, so we'll connect the output of w to head using a pipe | so that we only see the first five lines:

```
[hpc:login20~]$ w | head -5
09:39:27 up 65 days, 20:05, 10 users, load average: 0.72, 0.75, 0.78
USER
       TTY
               FROM
                                LOGIN@
                                        IDLE
                                              JCPU
                                                   PCPU WHAT
                                       0.00s 0.05s 0.02s w
kyclark pts/2
                gatekeeper.hpc.a 09:38
emsenhub pts/0
               gatekeeper.hpc.a 04:05
                                      14.00s 0.87s 0.87s -bash
joneska pts/3
               gatekeeper.hpc.a 08:25
```

Really we want to see the first five *users*, not the first five *lines* of output. To skip the first two lines of headers from w, we can pipe w into awk and tell it we only want to see output when the Number of Records (NR) is greater than 2:

```
[hpc:login20~]$ w | awk 'NR>2' | head -5
                                           0.00s 0.07s 0.03s w
kyclark pts/2
                 gatekeeper.hpc.a 09:38
emsenhub pts/0
                 gatekeeper.hpc.a 04:05
                                          26.00s 0.87s 0.87s -bash
joneska pts/3
                 gatekeeper.hpc.a 08:25
                                           1:13m 0.16s 0.12s vim results x2r
shawtaro pts/4
                 gatekeeper.hpc.a 08:06
                                          58:34
                                                  0.17s 0.17s -bash
darrenc pts/5
                 gatekeeper.hpc.a 07:58
                                          51:07
                                                  0.14s 0.07s qsub -I -N pipe
```

awk takes a PREDICATE and a CODE BLOCK (contained within curly brackets {}). Without a PREDICATE, awk prints the whole line. I only want to see the first column, so I can tell awk to print just column \$1:

```
[hpc:login2@~]$ w | awk 'NR>2 {print $1}' | head -5
kyclark
emsenhub
joneska
shawtaro
darrenc
```

We can see that the some users like "joneska" are logged in multiple times:

```
[hpc:login2@~]$ w | awk 'NR>2 {print $1}'
kyclark
emsenhub
joneska
shawtaro
darrenc
guven
guven
```

```
guven
joneska
dmarrone

Let's uniq that output:
[hpc:login20~]$ w | awk 'NR>2 {print $1}' | uniq
kyclark
emsenhub
joneska
shawtaro
darrenc
guven
joneska
dmarrone
```

Hmm, that's not right – "joneska" is listed twice, and that is not unique. Remember that uniq only works on sorted input? So let's sort those names first:

```
[hpc:login2@~]$ w | awk 'NR>2 {print $1}' | sort | uniq
darrenc
dmarrone
emsenhub
guven
joneska
kyclark
shawtaro
```

To count how many unique users are logged in, we can use the wc (word count) program with the -1 (lines) flag to count just the lines from the previous command

```
[hpc:login2@~]$ w | awk 'NR>2 {print $1}' | sort | uniq | wc -1
```

So what you see is that we're connecting small, well-defined programs together using pipes to connect the "standard input" (STDIN) and "standard output (STDOUT) streams. There's a third basic file handle in Unix called standard error" (STDERR) that we'll come across later. It's a way for programs to report problems without simply dying. You can redirect errors into a file like so:

```
$ program 2>err
$ program 1>out 2>err
```

The first example puts STDERR into a file called "err" and lets STDOUT print to the terminal. The second example captures STDOUT into a file called "out" while STDERR goes to "err."

NB: Sometimes a program will complain about things that you cannot fix, e.g., find may complain about file permissions that you don't care about. In those

cases, you can redirect STDERR to a special filehandle called <code>/dev/null</code> where they are forgotten forever – kind of like the "memory hole" in 1984.

```
find / -name my-file.txt 2>/dev/null
```

Count "oo" words

Do those numbers add up?

On almost every Unix system, you can find /usr/share/dict/words. Let's use grep to find how many have the "oo" vowel combination. It's a long list, so I'll pipe it into "head" to see just the first five:

```
$ grep 'oo' /usr/share/dict/words | head -5
abloom
aboon
aboveproof
abrood
abrook
```

Yes, that works, so redirect those words into a file and count them. Notice the use of !\$ (bang-dollar) to reference the last argument of the previous line so that I don't have to type it again (really useful if it's a long path):

```
$ grep 'oo' /usr/share/dict/words > oo-words
$ wc -1 !$
10460 oo-words
Let's count them directly out of grep:
$ grep 'oo' /usr/share/dict/words | wc -l
10460
Do any of those words additionally contain the "ow" sequence?
$ grep 'oo' /usr/share/dict/words | grep 'ow' | head -5
arrowroot
arrowwood
balloonflower
bloodflower
blowproof
How many are there?
$ grep 'oo' /usr/share/dict/words | grep 'ow' | wc -l
158
How many do not contain the "ow" sequence?
$ grep 'oo' /usr/share/dict/words | grep -v 'ow' | wc -l
10302
```

```
$ bc <<< 158+10302
10460
```

Excellent. Smithers, massage my brain.

Something with sequences

Now we will get some sequence data from the iMicrobe FTP site. Both wget and ncftpget will do the trick:

```
$ mkdir -p ~/contigs
$ cd !$
$ wget ftp://ftp.imicrobe.us/biosys-analytics/contigs/contigs.zip
     NB: How do we know we got the correct data? Go back and look at
     that FTP site, and you will see that there is a "contigs.zip.md5" file
     that we can less on the server to view the contents:
$ ncftp ftp://ftp.imicrobe.us/biosys-analytics/contigs
NcFTP 3.2.6 (Dec 04, 2016) by Mike Gleason (http://www.NcFTP.com/contact/).
Connecting to 150.135.44.10...
Welcome to the imicrobe.us repository
Logging in...
Login successful.
Logged in to ftp.imicrobe.us.
Current remote directory is /biosys-analytics/contigs.
ncftp /biosys-analytics/contigs > ls
                   contigs.zip.md5
contigs.zip
ncftp /biosys-analytics/contigs > cat contigs.zip.md5
1b7e58177edea28e6441843ddc3a68ab contigs.zip
ncftp /biosys-analytics/contigs > exit
You can read up on MD5 (https://en.wikipedia.org/wiki/Md5sum) to understand
that this is a signature of the file. If we calculate the MD5 of the file we dowloaded
and it matches what we see on the server, then we can be sure that we have the
```

exact file that is on the FTP site:

```
$ md5 contigs.zip
MD5 (contigs.zip) = 1b7e58177edea28e6441843ddc3a68ab
```

Yes, those two sums match. Note that sometimes the program is also named md5sum.

So, back to the exercise. Let's unpack the contigs:

```
$ unzip contigs.zip
Archive: contigs.zip
  inflating: group12_contigs.fasta
  inflating: group20 contigs.fasta
```

```
inflating: group24_contigs.fasta
$ rm contigs.zip
```

These files are in FASTA format (https://en.wikipedia.org/wiki/FASTA_format), which basically looks like this:

>MCHU - Calmodulin - Human, rabbit, bovine, rat, and chicken ADQLTEEQIAEFKEAFSLFDKDGDGTITTKELGTVMRSLGQNPTEAELQDMINEVDADGNGTID FPEFLTMMARKMKDTDSEEEIREAFRVFDKDGNGYISAAELRHVMTNLGEKLTDEEVDEMIREA DIDGDGQVNYEEFVQMMTAK*

>gi|5524211|gb|AAD44166.1| cytochrome b [Elephas maximus maximus]
LCLYTHIGRNIYYGSYLYSETWNTGIMLLLITMATAFMGYVLPWGQMSFWGATVITNLFSAIPYIGTNLV
EWIWGGFSVDKATLNRFFAFHFILPFTMVALAGVHLTFLHETGSNNPLGLTSDSDKIPFHPYYTIKDFLG
LLILILLLLLALLSPDMLGDPDNHMPADPLNTPLHIKPEWYFLFAYAILRSVPNKLGGVLALFLSIVIL
GLMPFLHTSKHRSMMLRPLSQALFWTLTMDLLTLTWIGSQPVEYPYTIIGQMASILYFSIILAFLPIAGX
IENY

Header lines start with ">", then the sequence follows. Sequences may be broken up over several lines of 50 or 80 characters, but it's just as common to see the sequences take only one (sometimes very long) line. Sequences may be nucleotides, proteins, very short DNA/RNA, longer contigs (shorter strands assembled into contiguous regions), or entire chromosomes or even genomes.

So, how many sequences are in "group12_contigs.fasta"? To answer, we just need to count how many times we see ">". We can do that with "grep":

```
$ grep > group12_contigs.fasta
Usage: grep [OPTION]... PATTERN [FILE]...
Try 'grep --help' for more information.
```

What is going on? Remember when we captured the "oo" words that we used the ">" symbol to tell Unix to *redirect* the output of grep into a file. We need to tell Unix that we mean a literal greater-than sign by placing it in single or double quotes or putting a backslash in front of it:

```
$ grep '>' group12_contigs.fasta
$ grep \> group12 contigs.fasta
```

You should actually see nothing because something quite insidious happened with that first "grep" statement – it overwrote our original "group12_contigs.fasta" with the result of "grep"ing for nothing, which is nothing:

```
$ ls -l group12_contigs.fasta
-rw-rw---- 1 kyclark staff 0 Aug 10 15:08 group12_contigs.fasta
```

Ugh, OK, I have to go back and wget the "contigs.zip" file to restore it. That's OK. Things like this happen all the time.

```
$ ls -lh group12_contigs.fasta
-rw-rw---- 1 kyclark staff 2.9M Aug 10 14:38 group12_contigs.fasta
```

Now that I have restored my data, I want to count how many greater-than signs in the file:

```
$ grep '>' group12_contigs.fasta | wc -1
132
```

Hey, I could see doing that often. Maybe we should make this into an "alias" (see above). The problem is that the "argument" to the function (the filename) is stuck in the middle of the chain of commands, so it would make it tricky to use an alias for this. We can create a bash function that we add to our \$HOME/.bashrc:

```
function countseqs() {
  grep '>' $1 | wc -1
}
```

After you add this, remember to source this file to make it available:

```
$ source ~/.bashrc
$ countseqs group12_contigs.fasta
132
```

Same answer. Good. However, someone beat us to the punch. There is a powerful tool called "seqmagick" (https://github.com/fhcrc/seqmagick) that will do this (and much, much more). It's installed into the "hurwitzlab/bin" directory, or you can install it locally:

Run "seqmagick -h" to see everything it can do.

Moving on, let's find how many contig IDs in "group 12_contigs.fasta" contain the number "47":

```
$ grep 47 group12_contigs.fasta > group12_ids_with_47
[login3@~/work/sequences]$ cat !$
cat group12_ids_with_47
>Contig_247
>Contig_447
>Contig_476
>Contig_4764
>Contig_4767
>Contig_13471
```

Here we did a little "useless use of cat," but it's OK. We also could have used "less" to view the file. Here's another useless use of cat to copy a file:

```
$ cat group12_ids_with_47 > temp1_ids
```

Additionally, we want to copy the file again to make duplicates:

```
$ cp group12_ids_with_47 temp2_ids
```

How can we be sure these files are the same? Let's use "diff":

```
$ diff temp1_ids temp2_ids
```

You should see nothing, which is a case of "no news is good news." They don't differ in any way. We can verify this with "md5sum":

```
$ md5sum temp*
957390ab4c31db9500d148854f542eee temp1_ids
957390ab4c31db9500d148854f542eee temp2 ids
```

They are the same file. If there were even one character difference, they would generate different hashes.

Now we will create a file with duplicate IDs:

```
$ cat temp1_ids temp2_ids > duplicate_ids
```

Check contents of "duplicate_ids" using "less" or "cat." Now grab all of the contigs IDs from "group20_contigs.fasta" that contain the number "51." Concatenate the new IDs to the duplicate_ids file in a file called "multiple_ids":

```
$ cp duplicate_ids multiple_ids
$ grep 51 group20_contigs.fasta >> !$
grep 51 group20_contigs.fasta >> multiple_ids
```

Notice the ">>" arrows to indicate that we are appending to the existing "multiple ids" file.

Remove the existing "temp" files using a "*" wildcard:

\$ rm temp*

Now let's explore more of what "sort" and "uniq" can do for us. We want to find which IDs are unique and which are duplicated. If we read the manpage ("man uniq"), we see that there are "-d" and "-u" flags for doing just that. However, we've already seen that input to "uniq" needs to be sorted, so we need to remember to do that:

```
$ sort multiple_ids | uniq -d > temp1_ids
$ sort multiple_ids | uniq -u > temp2_ids
$ diff temp*
1,7c1,11
< >Contig_13471
< >Contig_1947
< >Contig_247
< >Contig_447
< >Contig_476
< >Contig_4764
```

```
< >Contig_4767
---
> >Contig_10051
> >Contig_1651
> >Contig_4851
> >Contig_5141
> >Contig_5143
> >Contig_5164
> >Contig_5170
> >Contig_5188
> >Contig_6351
> >Contig_9651
> >Contig_9851
```

Let's remove our temp files again and make a "clean_ids" file:

```
$ rm temp*
$ sort multiple_ids | uniq > clean_ids
$ wc -l multiple_ids clean_ids
25 multiple_ids
18 clean_ids
43 total
```

We can use "sed" to alter the IDs. The "s//" command say to "substitute" the first thing with the second thing, e.g., to replace all occurences of "foo" with "bar", use "s/foo/bar" (http://stackoverflow.com/questions/4868904/what-is-the-origin-of-foo-and-bar).

```
$ sed 's/C/c/' clean_ids
$ sed 's/_/./' clean_ids
$ sed 's/>//' clean_ids > newclean_ids
```

That last one removes the FASTA file artifact that identifies the beginning of an ID but is not part of the ID. We can use this with "seqmagick" now to extract those sequences and find out how many were found:

30751 16768.14

7

5587

We can get stats on all our files:

```
$ seqmagick info *fasta > fasta-info
$ cat !$
```

newgroup12_contigs.fasta FALSE

name	alignment	min_len	max_len	avg_len	num_seqs
<pre>group12_contigs.fasta</pre>	FALSE	5136	116409	22974.30	132
<pre>group20_contigs.fasta</pre>	FALSE	5029	22601	7624.38	203
group24 contigs.fasta	FALSE	5024	81329	12115.70	139

We can use "cut" to view various columns:

```
$ cut -f 2 fasta-info
$ cut -f 2,4 fasta-info
$ cut -f 2-4 fasta-info
```

But it does not line up very nicely. We can use "column" to fix this:

```
$ cut -f 2-4 fasta-info | column -t
alignment min_len max_len
FALSE 5136 116409
FALSE 5029 22601
FALSE 5024 81329
FALSE 5587 30751
```

Gapminder

Do the following:

```
$ git clone https://github.com/kyclark/metagenomics-book
$ cd metagenomics-book/problems/gapminder/data
```

How many files are in the "data" directory?

```
$ ls | wc -1
```

How many lines are in each/all of the files?

```
$ wc -1 *
```

You can use cat to spew at the entire contents of a file into your shell, but if you'd just like to see the top of a file, you can use:

```
$ head Trinidad_and_Tobago.cc.txt
```

If you only want to see 5 lines, use -n 5 or -5.

For our exercise, we'd like to combine all the files into one file we can analyze. That's easy enough with:

```
$ cat * > all.txt
```

Let's use head to look at the top of file:

\$ head -5 all.txt Afghanistan 22227415 41.763 1997 Asia 635.341351 Afghanistan 2002 25268405 Asia 42.129 726.7340548 Afghanistan 2007 31889923 43.828 974.5803384 Asia Afghanistan 1952 8425333 Asia 28.801 779.4453145 Afghanistan 1957 9240934 30.332 820.8530296 Asia

Hmm, there are no column headers. Let's fix that. There's one file that's pretty different in content (it has only one line) and name ("country.cc.txt"):

```
$ cat country.cc.txt
country year pop continent lifeExp gdpPercap
```

Those are the headers that you can combine to all the other files to get named columns, something very important if you want to look at the data in Excel and R/Python data frames.

```
$ rm all.txt
$ mv country.cc.txt headers
$ cat headers *.txt > all.txt
$ head -5 all.txt | column -t
country
            year pop
                           continent lifeExp
                                               gdpPercap
Afghanistan 1997 22227415 Asia
                                      41.763
                                               635.341351
                                      42.129
Afghanistan 2002 25268405 Asia
                                               726.7340548
Afghanistan 2007
                                      43.828
                                               974.5803384
                 31889923 Asia
Afghanistan
           1952 8425333
                                      28.801
                                               779.4453145
                           Asia
```

Yes, that looks much better. Double-check that the number of lines in the all.txt match the number of lines of input:

```
$ wc -l *.cc.txt headers
$ wc -l all.txt
```

How many observations do we have for 1952?

```
$ grep 1952 all.txt | wc -l
$ cut -f 2 *.cc.txt | grep 1952 | wc -l
```

Those numbers aren't the same! Why is that?

```
$ grep 1952 all.txt | cut -f 2 | sort | uniq -c
142 1952
    1 1982
    1 1987
$ grep 1952 all.txt | grep 198[27]
```

Lebanon 1982 3086876 Asia 66.983 7640.519521 Mozambique 1987 12891952 Africa 42.861 389.8761846

How many observations for every year?

How many observations are present for Africa?

```
$ grep Africa all.txt | wc -1
```

How many for each continent?

```
$ cut -f 4 *.cc.txt | sort | uniq -c
```

What was the world population in 1952? As we've seen, just using grep 1952 is not sufficient. We want to take the 3rd column if the 2nd column is equal to "1952." awk will let us do just that. Normally awk will split on whitespace, so we need to use -F"\t" to tell it to split on the tab (\t) character. Use man awk to learn more.

```
$ awk -F"\t" '$2 == "1952" { print $3 }' *.cc.txt
```

I'll bet you didn't notice that one of those numbers was in scientific notation. That's going to cause a problem. Here it is:

```
$ awk -F"\t" '$2 == "1952" { print $3 }' *.cc.txt | grep [a-z]
3.72e+08
```

We have to throw in a grep -v to get rid of that (the -v reverses the match), then use the paste command is used to put a "+" in between all the numbers:

It bothers me that it's not an integer, so I'm going to use printf in the awk command to trim that:

How did population change over the years? Let's put a list of the unique years into a file called "years" and then cat over that to run the above for each year:

```
$ cut -f 2 *.txt | sort | uniq > years

$ for year in `cat years`; do echo -n $year ": " && awk -F"\t" "\$2 == $year { printf \"%d\n

1952 : 2406957150

1957 : 2664404580

1962 : 2899782974

1967 : 3217478384

1972 : 3576977158

1977 : 3930045807
```

1982 : 4289436840 1987 : 4691477418 1992 : 5110710260 1997 : 5515204472 2002 : 5886977579

2007 : 6251013179

That's kind of useful! Here's how I might put that into a script:

```
$ cat pop-years.sh
#!/bin/bash
```

```
set -u
YEARS="years"
cut -f 2 ./*.cc.txt | sort | uniq > "$YEARS"
NUM=$(wc -l $YEARS | awk '{print $1}')
if [[ "$NUM" -lt 1 ]]; then
      echo "No years ($NUM)!"
      exit 1
fi
while read -r YEAR; do
            echo -n "$YEAR: "
            awk -F"\t" "\s2 == \$YEAR \{ printf \"\d\n\", \s3 \}" ./*.cc.txt | grep -v "[a-z]" | pasterner | paster
done < "$YEARS"
$ ./pop-years.sh
1952: 2406957150
1957: 2664404580
1962: 2899782974
1967: 3217478384
1972: 3576977158
1977: 3930045807
1982: 4289436840
1987: 4691477418
1992: 5110710260
1997: 5515204472
2002: 5886977579
2007: 6251013179
How has life expectancy changed over the years? For this we'll need to write a
little Python program. I'll cat the program so you can see it. You can type this
in with nano and then do chmod +x avg.py to make it executable (or use the
one I added):
$ cat avg.py
#!/usr/bin/env python3
import sys
args = list(map(float, sys.argv[1:]))
print(str(sum(args) // len(args)))
$ for year in `cat years`; do echo -n "$year: " && grep $year *.txt | cut -f 5 | xargs ./av
1952: 49.0
1957: 51.0
1962: 53.0
```

```
1967: 55.0
1972: 57.0
1977: 59.0
1982: 61.0
1987: 63.0
1992: 64.0
1997: 65.0
2002: 65.0
2007: 66.0
```

How many observations where the life expectancy ("lifeExp," field #5) is greater than 40? For this, let's use the awk tool.

```
 * awk -F"\t" '$5 > 40' all.txt | wc -1
```

How many of those are from Africa? We can either use cut to get the 4th field or ask awk to print the 4th field for us:

How many countries had a life expectancy greater than 70, grouped by year?

```
$ awk -F"\t" '$5 > 70 { print $2 }' *.cc.txt | sort | uniq -c
5 1952
9 1957
16 1962
25 1967
30 1972
38 1977
44 1982
49 1987
54 1992
65 1997
75 2002
83 2007
```

How could we add continent to this?

As you look at the data and want to ask more complicated questions like how does gdpPercap affect lifeExp, you'll find you need more advanced tools like Python or R. Now that the data has been collated and the columns named, that will be much easier.

What if we want to add headers to each of the files?

```
$ mkdir wheaders
$ for file in *.txt; do cat headers $file > wheaders/$file; done
$ wc -l wheaders/* | head -5
```

- 13 wheaders/Afghanistan.cc.txt
- 13 wheaders/Albania.cc.txt
- 13 wheaders/Algeria.cc.txt
- 13 wheaders/Angola.cc.txt
- 13 wheaders/Argentina.cc.txt

\$ head wheaders/Vietnam.cc.txt

country	year	pop	contin	ent 1	LifeExp	${\tt gdpPercap}$
Vietnam	1952	262468	39 A	sia 4	10.412	605.0664917
Vietnam	1957	289985	543 A	sia 4	12.887	676.2854478
Vietnam	1962	337961	.40 A	sia 4	15.363	772.0491602
Vietnam	1967	394639	010 A	sia 4	17.838	637.1232887
Vietnam	1972	446550)14 A	sia 5	50.254	699.5016441
Vietnam	1977	505335	506 A	sia 5	55.764	713.5371196
Vietnam	1982	561421	.81 A	sia 5	58.816	707.2357863
Vietnam	1987	628264	91 A	sia 6	32.82	820.7994449
Vietnam	1992	699407	'28 A	sia 6	67.662	989.0231487