UNIX Shell Programming

Reminder that there are some useful tutorials for UNIX Shell use and programming. See the Software Carpentries tutorial and give it a try.

Basic UNIX programming in the BASH shell can help you do make some simple things more possible. More complicated programming is probably better achieved in a scripting language like Python which will be covered in the rest of the course, but BASH can be very powerful and useful to apply these to improve the tools.

See the Software Carpentry tutorial.

Variables

Variables are used to store information in Variables. To access a value of a variable in UNIX you can prefix it with \$.

For example to assign a variable a value

```
NAME="GeneA"
NAME2="GeneB"
echo "$NAME $NAME2"

NAME="GeneC"
NAME3=$NAME.$NAME2
echo "$NAME $NAME2 $NAME3"
```

Loops and Logic

if [TEST]; then DOSOMETHING fi can be used to test for a logical statement. This testing structure also allows for other conditions to be met with elif or "else if" and else.

For example:

```
if [ $NAME == "GeneC" ]
then
    echo "Name is C"
fi
if [ $NAME != "GeneA" ]
then
    echo "Name is not GeneA"
fi
NAME="GeneA"
if [ $NAME == "GeneA" ]; then
    echo "A"
```

```
elif [ $NAME == "GeneB" ]; then
  echo "B"
else
  echo "had another class for NAME: $NAME"
NAME="genea"
if [ $NAME == "GeneA" ]; then
  echo "A"
elif [ $NAME == "GeneB" ]; then
  echo "B"
else
  echo "had another class for NAME: $NAME"
fi
The structure requires the [ ] and there is expected to be a space between the
[ or ] and the options; The then is also require but if you want to compact this
slightly differently.
if [ $NAME == "GeneC" ]; then echo "Name is C"; fi
Multiple tests can be applied in same if statement but require double brackets.
NAME=GeneC
if [[ $NAME == "GeneC" || $NAME == "GeneB" ]]; then
  echo "Name is $NAME"
fi
More logical operator such as testing if a number if smaller or greater with -gt
and -lt.
NUM=10
if [ $NUM -gt "0" ]; then
  echo "NUM is greater than 0"
fi
Can test if one file is newer than another with -nt. Also showing how to use
touch fileA.fasta
echo "second file" > fileB.fasta
if [ fileA.fasta -nt fileB.fasta ]; then
  echo "File A is newer"
```

Really useful testing options

echo "File B is newer"

fi

• -f - if the variable is a file and exists

```
-s - if file exists and is not zero
-d - if the variable is a directory
-z - if variable is empty
if [ ! -f $file1 ]; then
echo "file $file1 does NOT exist"
fi
if [ -z $var1 ]; then
echo "variable $var1 is empty"
fi
if [ -s $file2 ]; then
echo "file $file2 exists and is not empty"
fi
```

Loops

Loops are important components for iterating through data. For loops we can specify a list to go through explicitly. for loops are structured with for VARIABLE in LIST; do DOSOMETHING done

```
for n in A B C D
do
echo "$n"
done
```

Can also use the results of a function to loop through a dataset, folder of files, etc. For loops are used when the specific list is available at the start of the loop.

```
for file in $(ls *.fa)
do
   echo "file $file is found"
done
```

Can use the seq function to make a list of numbers. Arguments are either the ending number, or start and end, or start, end, and offset.

```
seq 3 # start at 1 and count to 3
1
2
3
seq 5 7 # start at 5 end at 7
5
6
7
```

```
seq 5 2 10 # start at 5, end at 10, offset by 2
5
7
9
So if you want to iterate through a bunch of numbers.
for m in $(seq 3 15)
do
    echo "m is $m"
done
```

Using UNIX tools with Variables

Capturing output from a program is also a useful. For example if you want to do simple mathematical arithmetic with the UNIX tool expr (or "evaluate expression"). It takes arguments for simple math.

To save the result from a command you can use the \$() structure and also can use the "'" backquote, they both will work for taking the output from an application and saving it in a variable.

```
n=$(echo "ABCDEFG" | wc -c) # this prints out the number of characters
echo "$n characters"
n=`echo "ABCD" | wc -c`
echo "$n characters"
a=1
echo "a is $a"
expr $a + 1
a=$(expr $a + 1)
echo "a is now $a"
```

Loops again

While loops can be used which can run

```
N=1
while [ $N -lt 10 ]
do
   echo "N is $N"
   N=$(expr $N + 1)
done
```

Can also use while to read data from a file using the read directive.

```
echo "wolf tooth animal" > data.txt
echo "snake fang animal" >> data.txt
```

```
echo "mantis mandible insect" >> data.txt
while read COL1 COL2 COL3
do
 echo "COL1 is $COL1; COL3 is $COL3"
done < data.txt</pre>
How these columns are delimited are dependent on an environment variable
defined $IFS. For example to separate columns based on comma:
echo "wolf,tooth,animal" > data.csv
echo "snake,fang,animal" >> data.csv
echo "mantis,mandible,insect" >> data.csv
IFS=,
while read COL1 COL2 COL3
 echo "COL1 is $COL1; COL3 is $COL3"
done < data.csv
Can also pass data INTO the while loop with pipes. This is a really useful way
to parse out columns of data.
echo "Hop, Skip, Jump" | while read COL1 COL2 COL3;
   echo "COL1=$COL1 ... COL2 is $COL2"
done
Data Processing
https://www.safaribooksonline.com/library/view/bioinformatics-data-
skills/9781449367480/ch07.html#chapter-07
https://github.com/biodataprog/GEN220_data/tree/main/data
sort Sort data and files.
sort file.txt > file.sorted.txt
Type of sorting: * -d/-dictionary order: consider only blanks & alphanumeric
characters * -n/-numeric-sort : compare according to string numerical value *
-f/-ignore-case: upper/lower doesn't matter * -r/-reverse: reverse the order *
```

for n in \$(seq 100); do echo \$((\$RANDOM%100)); done > numbers.txt

-k: specify the key positions to sort by

sort numbers.txt | head -n 10

10

#generate some random numbers between 0 and 100

```
10
12
25
30
34
39
42
49
```

But if sort by numeric - you see there are some numbers < 10 which weren't shown.

```
sort -n numbers.txt | head -n 10
0
1
2
3
4
6
6
8
8
13
```

 ${\bf uniq}$ - Collapse runs of words/numbers into unique list. This only works if the data are sorted.

```
sort -n numbers.txt | uniq | head -n 10
0
1
2
3
4
6
8
13
15
16
```

To see the numbers (or words) uniquely with counts of the occurrences use '-c'.

```
sort -n numbers.txt | uniq -c | head -n 10
    1 0
    1 1
    1 2
    1 3
    1 4
    2 6
```

```
2 8
   3 13
   2 15
   2 16
Hey let's sort this list so we know the numbers that show up most frequently
$ sort -n numbers.txt | uniq -c | sort -r -n | head -n 8
   4 91
   4 54
   4 32
   3 57
   3 22
   3 17
   3 13
   2 95
Sort Multicolumn data - you can sort by the 2nd or 3rd column.
head -n 10 data/rice_random_exons.bed
Chr7
        21408673
                     21408826
Chr9
        16031526
                     16031938
Chr11
         4762531
                     4762595
Chr8
        54040
                  54193
Chr10
         19815475
                      19815747
Chr3
        16171331
                     16172869
Chr10
         2077882
                     2077938
Chr3
        20517604
                     20517936
         9777446
Chr10
                     9777527
        4967096
                    4967246
Chr2
$ sort -k1,1 -k2,2n data/rice_random_exons.bed | head -n 5
Chr1
        12152
                  12435
Chr1
        98088
                  98558
Chr1
        216884
                   217664
Chr1
        291398
                   291534
Chr1
        338180
                   338310
$ sort -k1,1 -k2,2n data/rice_random_exons.bed | tail -n 5
        22369724
Chr9
                     22369776
Chr9
        22508926
                     22509014
Chr9
        22753347
                     22753458
Chr9
        22924316
                     22924424
ChrSy
         136034
                    136323
```

3

2

336

330

110

1

1

14

336

6

217148

198385

211012

14799

217483

198695

211119

15132

8.6e-83

4.1e-18

2.1e-10

1.3e-77

298.8

84.0

58.4

281.6

cut Cut - subselect and print certain columns from a file

108 25

YAR060C Chr I 100.00 336 0

YAR060C Chr_I 97.02 336 8

YAR060C Chr I 74.07

YAR060C Chr_I 64.00 325 95 22

```
YAR060C Chr_I 72.48 109 25
                                   6
                                         110
                                                20974
                                                         21081
                                                                   2.3e-10
YAR061W Chr_I 100.00 204 0
                                         204
                                                                    3.4e-54
                                    1
                                               218131
                                                         218334
YAR061W Chr_I 70.62
                      194 57
                                         194
                                               203400
                                                         203593
                                                                    6.5e-23
                                   1
YAR061W Chr_I 94.61
                      204 7
                                   204
                                           1
                                                13951
                                                          14150
                                                                   5e-48
YAR061W Chr_I 67.88 193 62
                              0
                                    194
                                           2
                                                27770
                                                         27962
                                                                   3.9e-20
YAL030W Chr_I 100.00 252 0
                                    103
                                        354
                                                87502
                                                         87753
                                                                   2.5e-55
Just print out the first column of sequence names.
cut -f1 data/yeast_orfs-to-chr1.FASTA.tab | head -n 7
YAR060C
YAR060C
YAR060C
YAR060C
YAR060C
YARO61W
Print out Column 2
cut -f2 data/yeast_orfs-to-chr1.FASTA.tab | head -n 5
Chr_I
Chr_I
Chr_I
Chr_I
Chr_I
Get the Query name and Percent Identity
cut -f1,3 data/yeast_orfs-to-chr1.FASTA.tab | head -n 5
YAR060C
           100.00
YAR060C
           64.00
YAR060C
           74.07
YAR060C
           97.02
YAR060C
           72.48
YARO61W
           100.00
YAR061W
           70.62
YARO61W
           94.61
YARO61W
           67.88
YAL030W
           100.00
YAL030W
           98.15
Cut two columns out, and run sort to sort on the column
sort -k3,1nr data/yeast_orfs-to-chr1.FASTA.tab | cut -f1,3 | head -n 5
        100.00
HRA1
YALO01C
           100.00
YALO02W
           100.00
           100.00
YAL003W
```

58.2

90.0

207.7

182.6

203.1

99.2

YAL003W

100.00

```
$ sort -k9,1n data/yeast_orfs-to-chr1.FASTA.tab | head -n 5
YAL069W
           100.00
                     335
                             649
YAL068W-A
             100.00
                       538
                               792
                              2169
YAL068C
           100.00
                     1807
YARO20C
           79.76
                    2008
                             2169
YALO67W-A
             100.00
                                2707
                       2480
```

Made up example, but you can cut two columns out. And also PASTE things back together.

```
cut -f1,3,4 data/yeast_orfs-to-chr1.FASTA.tab > first_cols.tab
cut -f1,7 data/yeast_orfs-to-chr1.FASTA.tab > second_cols.tab
paste first_cols.tab second_cols.tab | head -n 5
YAL027W 100.00 786
                       YALO27W 1
tL(CAA)A
               100.00 44
                               tL(CAA)A
                                               39
tL(CAA)A
               100.00 38
                               tL(CAA)A
                                               1
YAL028W 100.00 1587
                       YAL028W 1
YAL029C 100.00 4416
                       YAL029C 4416
```

AWK

Can also use awk to process column data.

```
awk '{print $1}' yeast_orfs-to-chr1.FASTA.tab # print out the first column of a file
# specificy a different delimiter (,)
head -n 3 data/random_exons.csv
Chr5,27781790,27781800
Chr11,14656670,14656870

$ awk -F, '{print $1,$2}' data/random_exons.csv | head -n 3
Chr5 27781790
Chr11 14656670
Chr3 14560358

Here get the length of an alignment (column 6 is the START and column 7 is the end)
```

Advanced Variable usage

BASH also supports the concepts of Arrays. This tutorial provides useful summary of how to use arrays.

awk '{print \$7-\$6}' data/yeast_orfs-to-chr1.FASTA.tab

A simple example is like this

```
animals=(dog cat mouse)
for name in ${animals[@]};
do
  echo "name is $name"
done
# add to the array
animals+=(snake)
for name in ${animals[@]};
do
  echo "name is $name"
done
```