#### 1 Unix for Bioinformatics - Solutions

No questions in this section.

### 2 Basic Unix

- 1. ls -al
- 2. There are 4 files in the directory (and 2 subdirectories). You can use 1s -1 to look inside the directory. This will show you which of the contents are files and which are directories. Don't forget to also include the -a option to show any hidden files:
- ls -la Pfalciparum
- **3.** Malaria.fa is the largest file. You can add the -h option to the command above to make the size of the files more readable.
- 4. cd Pfalciparum
- **5.** There is one file in the fasta directory (hint: it's a hidden file!).
- 6. cp Pfalciparum.bed annotation
- 7. mv \*.fa fasta
- 8. 4 files.
- **9.** There are 6 GFFs in the unix directory. To search from the Unix directory, you can either use cd to move up to the directory, or you can specify the path in the find command. This can either be the absolute path, which you can get from pwd, or you can use the relative path, like so:

```
find ../../.. -name *.gff
```

**10.** There are 7 fasta files in the unix directory. Note that fasta files normally end with .fa OR .fasta, so wee need to make sure we look for both of these, by adding a wildcard after fa:

```
find ../../.. -name *.fa*
```

# 3 Looking inside files

- 1. head -n 500 Styphi.gff > Styphi.500.gff
- **2.** There are 6213 lines in the file. Use the -1 option:

```
wc -1 Pfalciparum.bed
```

- 3. sort -k 1 -k 2 -n Pfalciparum.bed
- **4.** Here is one way to do this: First use awk to get the first column of the file. Sort this and then use the -c option for uniq to count how many entries each chromosome has:

```
awk '{ print $1 }' Pfalciparum.bed | sort | uniq -c
```

The expected output would be:

```
190 01
 264 02
 287 03
 292 04
 357 05
 373 06
 395 07
 374 08
 425 09
 452 10
 553 11
 621 12
 773 13
857 14
    Searching inside files with grep
1. grep "^>" exercises.fasta
2. There are 1000 sequences. We can use -c to count the number of matches:
grep -c "^>" exercises.fasta
Or pipe into wc:
grep "^>" exercises.fasta | wc -l
3. Yes, three of them:
>sequence27 spaces in the name
>sequence52 another with spaces
>sequence412 yet another with spaces
One option is two greps piped together:
grep "^>" exercises.fasta | grep " "
Alternatively, in one regular expression
grep "^>.* .*" exercises.fasta
4. grep -v "^>" exercises.fasta
5. Three. First extract the sequences, then search for n:
grep -v "^>" exercises.fasta | grep -c -i n
6. Yes, one sequence. Try:
```

grep -v "^>" exercises.fasta | grep -i -v "^[acgtn].\*\$"

grep -v "^>" exercises.fasta | grep -i "[^ACGTN]"

**7.** 66 sequences. Try:

Alternatively, we can use the ^ to ask for matches NOT in the alphabet [acgtn]

```
grep -v "^>" exercises.fasta | grep -c "GC[AT]GC"
8. We found the total number of sequences earlier:
grep -c "^>" exercises.fasta
... which outputs 1000
This finds the number of unique names:
grep "^>" exercises.fasta | sort | uniq | wc -1
... which outputs 999.
Therefore there is 1000 - 999 = 1 name repeated.
```

### Therefore there is 1000 /// I hame repeated

## 5 File processing with awk

1. Using:

```
awk -F"\t" '{print $1}' exercises.bed | sort -u
Should give you:
contig-1
contig-3
contig-4
contig-5
scaffold-2
```

**2.** There are 5 contigs. Use the command from the previous exercise and count the number of lines with wc:

```
awk -F"\t" '{print $1}' exercises.bed | sort -u | wc -l
```

**3.** There are 164 features on the positive strand. Try:

```
awk -F"\t" '$6=="+"' exercises.bed | wc -1
```

**4.** Ther are 124 features on the negative strand. Try:

```
awk -F"\t" $6=="-"' exercises.bed | wc -1
```

**5.** There are 33 repeats. Try:

```
awk -F"\t" $4 \sim /repeat/' exercises.bed | wc -l
```

#### 6 Advanced Linux

No exercises in this section

# 7 BASH Scripting

1. Here is an example of what this script could look like:

```
#!/usr/bin/env bash
set -e
# check that the correct number of options was given.
# If not, then write a message explaining how to use the
# script, and then exit.
if [ $# -ne 1 ]
then
    echo "usage: example_1.sh filename"
    echo "Prints the number of lines in the file"
    exit
fi
# Use sensibly named variable
filename=$1
# check if the input file exists
if [ ! -f $filename ]
then
    echo "File '$filename' not found! Cannot continue"
    exit
fi
# If still here, we can count the number of lines
number_of_lines=$(wc -l $filename | awk '{print $1}')
echo "There are $number_of_lines lines in the file $filename"
2. Here is an example of what this script could look like:
#!/usr/bin/env bash
set -e
for filename in ../scripts/loop_files/*; do ./exercise_1.sh $filename; done
3. Here is an example of what this script could look like:
#!/usr/bin/env bash
set -e
# Check if the right number of options given.
# If not, print the usage
if [ $# -ne 1 ]
    then
        echo "usage: example_3.sh in.gff"
        echo "Gathers some summary information from a gff file"
        exit
fi
```

```
# store the filename in a better named variable
infile=$1
# Stop if the input file does not exist
if [ ! -f $infile ]
then
   echo "File '$infile' not found! Cannot continue"
   exit 1
fi
echo "Gathering data for $infile..."
# Gather various stats on the file...
# Total number of lines/records in file
total_records=$(wc -l $infile | awk '{print $1}')
echo "File has $total_records records in total"
# Get the sources from column 2.
echo "The sources in the file are:"
awk '{print $2}' $infile | sort -u
# Count the sources
echo
echo "Count of sources, sorted by most common"
awk '{print $2}' $infile | sort | uniq -c | sort -n
# Count which features have no score
echo
echo "Count of features that have no score"
awk '$6=="." {print $3}' $infile | sort | uniq -c
# Find how many bad coords there are
echo
bad_coords=$(awk '$5 < $4' $infile | wc -l | awk '{print $1}')
echo "Records with bad coordinates: $bad_coords"
#_____#
```

```
WARNING: the following examples are more advanced!
#
# if there were records with bad coords, find the sources responsible
if [ $bad_coords != 0 ]
then
    echo
   echo "Sources of bad coordinates:"
    # Instead getting one source per line, pipe into awk again to print them
    # on one line with semicolon and space between the names
    awk '$5 < $4 {print $2}' $infile | sort -u | awk '{sources=sources"; "$1} END{print substr
fi
# Count of the features. Instead of using awk .... |sort | uniq -c
# we will just use awk. Compare this with the above method
# used to count the sources. Although it is a longer command, it is more efficient
echo "Count of each feature:"
awk '{counts[$3]++} END{for (feature in counts){print feature"\t"counts[feature]}}' $infile |
k2n
# This example is even more complicated! It uses a loop to
# get the mean score of the genes, broken down by source.
echo
echo "Getting mean scores for each source..."
for source in `awk '{print $2}' $infile | sort | uniq`
   awk -v s=$source '$2==s {total+=$6; count++} END{print "Mean score for", s":\t", total/count++}
done
# We can use awk to split the input into multiple output files.
# Writing print "line" > filename will append the string "line"
# to a file called filename. If a file called filename
# does not exist already, then it will be created.
# Write a new gff for each of the sources in the original input gff file
echo "Writing a file per source of the original gff file $infile to files called split.*"
awk '{filename="split."$2".gff"; print $0 > filename}' $infile
echo " ... done!"
```