Bi 621 - Problem Set 6

Our goal with this assignment is to generate some measures of accuracy for whole genome assemblies and then use these measures to assay our success in several Velvet assemblies.

Part 0 – Write unit tests

- 1. Read through Part 1 to be sure you understand the details of what your algorithm should do.
- 2. Create a file called Unit_test.fa that contains a text file that you know will yield a specific answer when analyzed by your code from Part 1.
- 3. Create a second file called expected_results.txt that contains the expected results from your test.
- 4. After finishing your code in Part 1, run your unit test to ensure your code behaves as expected.
- 5. OPTIONAL trade unit tests and results with 1 or more people in the class. See if you can pass their unit test.

Part 1 – Contig length distributions

- 1. Parse the contigs fa file that is output by velvetg. Extract the FASTA ID lines as you parse the file (remember: these strings will begin with the ">" character).
- 2. You can use the sample data contigs.fa from Talapas to test your code.
- 3. Using Python regular expressions, extract k-mer length of each contig (in red below). In addition, extract the k-mer coverage for the contig (in blue). Assume a k-mer length of 49.

```
>NODE 11 length 3717 cov 19.315845
```

- 4. Adjust the k-mer length to represent the physical length. Calculate the number of contigs, the maximum contig length, the mean contig length, and the total length of the genome across the contigs. Calculate the mean depth of coverage for the contigs.
- 5. Calculate the N50 value of your assembly.
- 6. Calculate the distribution of contig lengths and bucket the contig lengths into groups of 100bp. So, all contigs with lengths between 0 and 99 would be in the 0 bucket, those with lengths between 100 and 199 would be in the 100 bucket, etc.
- 7. Print out the distribution.

```
# Contig length Number of contigs in this category
0    0
100  5324
200  3345
300  1130
```

Part 2 – Velvet

You will need to install Velvet in your home directory on Talapas to use it.

- On Talapas, in your home directory:
 wget https://www.ebi.ac.uk/~zerbino/velvet/velvet 1.2.10.tgz
- In your home directory, make a directory called bin (if you don't already have one)

- Move velvet 1.2.10.tgz into ~/bin
- Untar the file
- Move into the velvet 1.2.10 directory
- Issue the command make 'MAXKMERLENGTH=50' 'OPENMP=1'
- Move velveth and velvetg up one directory (into ~/bin)
- 1. All your work in this section should be completed using the queuing system on HPC. (See Nick's lecture notes on HPC and https://hpcrcf.atlassian.net/wiki/display/TCP/How-to+Submit+a+Job to remind yourself how the queuing system works)
- 2. You can find the data here:

```
/projects/bgmp/Bi621/rs_female_1983.13.1.fq.gz
/projects/bgmp/Bi621/rs_female_1983.13.2.fq.gz
```

Please do not copy the data, but rather refer to its location in your script. Remember to NOT write to the group project directory.

- 3. Run Velvet on the dataset. You can find the Velvet manual here: https://www.ebi.ac.uk/~zerbino/velvet/Manual.pdf
 - a. Calculate the k-mer coverage for the dataset assuming a stickleback genome size is 460 Mb
 - b. Run velveth/velvetg with k-mer sizes of 31, 41, and 49
 - c. Use your code from Part 1 to collect assembly statistics on each result
- 4. With a k-mer size of 49, adjust the coverage cutoff to 4x, 8x, and 'auto'. Again, assay your results with your code.
- 5. Finally, adjust the minimum contig length to 500bp and again, assay your results.

Part 3 – Questions

- 1. Describe how the assembly changes with different k-mer values using the assembly statistics you have collected. How does the contig length distribution change?
- 2. How does an increased coverage cutoff affect the assembly? What is happening to the de Bruijin graph when you change the value of this parameter? How does velvet calculate its value for 'auto'?
- 3. How does increasing minimum contig length affect your contig length distribution?

To turn in your work for this assignment, do the following:

Be sure to turn in your unit tests and expected results, your code, your output (mean, max, N50, etc.) and plots, as well as the answers to the questions above to GitHub.