Homework 3

Due date: Dec 2, 2020

De Bruijn Graph

• Build condensed De Bruijn Graph.

Write a program that takes fastq-file and k as input and:

- Builds condensed De Bruijn Graph
- Outputs edges in fasta-file
- Counts average kmer coverage for each edge (average kmer coverage is a number of times we see this kmer in reads, so average kmer edge coverage is an average of all its kmer coverages)
- Outputs result in a .dot file. Each edge should have a label with its length and average coverage
- Graph simplification: implement a graph simplification algorithm for basic erroneous edge removal.
 - Tip removal (remove all edges with short length and small coverage with tip topological structure).
 - Remove all low-covered and short edges
 Assess both approaches. Write a reasonable conclusion (you can add figures of your graphs to illustrate).

Hints: use k=55 for illustrations. Add kmer and reverse-complement kmer simultaneously, so your graph will be symmetric.

You can download data from https://drive.google.com/drive/folders/ 1M1XF4zEKwChssqt501XDZZB661PNIslG?usp=sharing. It contains 3 sets of reads and reference genomes for them.