

Genome-scale technologies 2 /
Algorithmic and statistical aspects of DNA sequencing
Assignment 2

winter semester 2021/2022

Task

Design and implement an assembly algorithm, destined to work on single-end reads originating from the same strand of a single chromosome.

In your program you:

- can use the codes from the classes,
- can not use programs and libraries to read assembly, mapping, alignment, etc.
- can not use multiprocessing commands.

The solution should include:

- program file `assembly`, executable using syntax:

```
./assembly input_reads.fasta output_contigs.fasta
```

- `readme` file with short description of your approach,
- program code (if executable is binary).

Input and minimum performance requirements

Typical parameters of input dataset:

- number of reads: 1000,
- read length: 80bp,
- average percentage of mismatches: $\leq 5\%$,
- average coverage: $\geq 5\times$.

Typical input dataset should be processed in time less then 1h on a common laptop, using up to 0.5GB memory.

Output and evaluation

The solutions will be evaluated on simulated data (i.e. artificial reads generated from a reference sequence). Output contigs will be locally aligned to the reference sequence and resulting alignments will be processed in the following way:

- ambiguous alignment fragments (sharing a reference sequence interval) will be trimmed away,
- alignments of length $< 300bp$ will be excluded.

Alignments passing filtering criteria will be scored according to the following formula:

$$S = \frac{ref_cov \cdot cont_cov \cdot \max(0.5, 10 \cdot (ident - 0.9))}{\log_5(4 + n_alments)}$$

where:

- *ref_cov* is the proportion of the reference sequence covered by the alignments,
- *cont_cov* is the proportion of the contigs' sequence covered by the alignments,
- *ident* is the identity proportion in the alignments,
- *n_alments* is the number of alignments.

Training data

You can download from moodle a training data package consisting of:

- directory **reference/** with a reference sequence file **reference.fasta** and **bowtie2** index files for this sequence,
- directory **reads/** with simulated read files:
 - **reads1.fasta** – 1000 reads containing ~1% mismatches,
 - **reads2.fasta** – 1000 reads containing 1-3% mismatches,
 - **reads3.fasta** – 1000 reads containing 3-5% mismatches,
- scripts to evaluate your assembly.

Scripts require **bowtie2** program and Python module **pysam**. Usage:

```
./evaluate.sh contigs.fasta
```

Two reference algorithms have been tested on the training datasets, the table below presents intended algorithm assessment and obtained *S* scores (*m* is the maximum assessment – see next section for details):

	algorithm1	algorithm2
assessment	<i>m</i> – 4	<i>m</i> – 2
score for dataset		
- reads1.fasta	0.06	0.59
- reads2.fasta	0.03	0.21
- reads3.fasta	0.03	0.08

Assessment and deadlines

- The assignment can be completed individually or in 2- or 3-person teams.
- Submit your team by email to dojer@mimuw.edu.pl till December 21.
- Submit your solution to moodle till January 18.
- Maximum assessment for the solution is
 - 12 points for assignments completed individually,
 - 11 points for assignments completed in 2-person teams,
 - 10 points for assignments completed in 3-person teams.
- If deadlines are not met, the maximum assessment is reduced by 2 points.