Instructor: Boese Assignment #5

DNA Sequences

Objectives

- Read in command-line arguments
- Read in data from files
- Loop through strings

Measuring DNA Similarity

DNA is the hereditary material in human and other species. Almost every cell in a person's body has the same DNA. All information in a DNA is stored as a code in four chemical bases: adenine (A), guanine (G), cytosine (C) and thymine(T). Different order of these bases means different information.

One of the challenges in computational biology is determining what the *codons* in a DNA sequence represent. A *codon* is a sequence of three DNA or RNA nucleotides that corresponds with a specific amino acid or stop signal during protein synthesis. For example, given the sequence GGGA, the codon could be a GGG or a GGA, depending on where the gene begins and the active bases in the gene. Clues about how to interpret a DNA sequence can be found by comparing an unknown DNA sequence to a known sequence and measuring their similarity. If sequences are similar, then it can be hypothesized that they have similar functions and proteins.

Hamming distance and similarity between two strings

Hamming distance is one of the most common ways to measure the similarity between two strings of the same length. Hamming distance is a position-by-position comparison that counts the number of positions in which the corresponding characters in the string are different. Two strings with a small Hamming distance are more similar than two strings with a larger Hamming distance.

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Example:
first string = "AGCCT"
second string = "AACCG"
A G C C T
| * | | *
A A C C G
```

In this example, there are three matching characters and two mismatches, so the Hamming distance is two.

The similarity score is then calculated as follows:

similarity_score = (string length - hamming distance) / string length

So for the example provided,

 $\rightarrow 0.6$

Example 1

mouseDNA1.txt

CATCATCATCATCATCTTTTT

humanDNA1.txt

ATGCATCATCATCATCTTTTT

unknownDNA1.txt

CATCATCTTCATCATCATCTTTTT

Output:

MouseCompare = 0.958333 HumanCompare = 0.833333

mouse



Example 2

(Each file should have the data all on one line in the file)

mouseDNA2.txt

humanDNA2.txt

unknownDNA2.txt

Output:

MouseCompare = 0.655882 HumanCompare = 0.820588 human

Assignment Details:

In this assignment, you will calculate the Hamming distance and similarity scores for sample DNA sequences from a human, a mouse, and an unknown species to determine the identity of the unknown species.

Requirements

- The name of the file must be called **DNA.cpp**
- Comments at the top of your program
 - Your name
 - o Date
 - Homework #
 - o Brief description of the assignment (one or two lines max)
- The program takes three command-line arguments, in the following order:

mouseDNA filename humanDNA filename unknownDNA filename



• Use this algorithm to determine the identity of the unknown DNA (if the difference < 0.0001):

IF unknownDNA is equally similar to both mouse and human,
PRINT "identity cannot be determined"
ELSE IF the unknownDNA is more similar to the humanDNA,
PRINT "human"
ELSE IF the unknownDNA is more similar to the mouseDNA,
PRINT "mouse"

Note: You need to check if they are essentially equal first otherwise this algorithm doesn't work. Do you understand why?

- All data files for a given test case will be the same length.
- You may <u>not</u> create functions, classes, arrays, linked lists, (*e.g., do not use things that we have not covered yet unless you get permission from the instructor first*).
- The output must match exactly to the examples provided (given appropriate inputs).
- If you do not use command-line arguments then you will receive a zero for this assignment.
- Program must be written in C++ and submitted in Moodle.
- Zip the DNA.cpp and submit to Moodle as *Firstname_Lastname_HW5.zip*.

