Lab 1: Strings and Loops

IBCS SL and HL

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1 DNA Background

In this lab you will practice handling Strings in Java. To do so, you will read in a DNA sequence that is mean to create a pump for a cell membrane. DNA provides a blueprint to create proteins which in turn create all the necessary parts for us to stay alive. This process involves a few steps:

- 1. DNA starts with one strand and that strand's compliment, forming a two strand double helix.
- 2. When ready, the DNA will split apart and messenger RNA, or mRNA, will create a copy.
- 3. The mRNA will then travel into a ribosome, The ribosome will read in the mRNA and create amino acids by reading a section at a time.
- 4. A chain of amino acids will then form the proteins necessary.
- 5. Watch this video and this one.

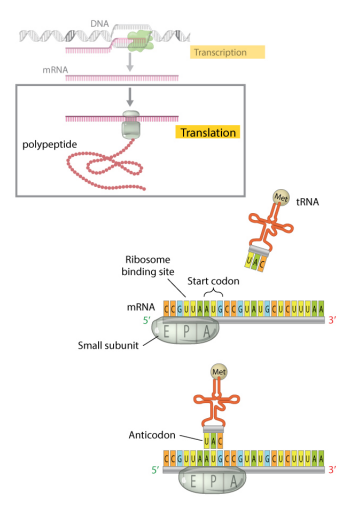


Figure 1: Protein Creation

2 Assignment

- 1. Create a DNAStrand class.
 - (a) Create a method, readDNA, that reads in the dna sequence file by <u>looping</u> through each line. <u>Store</u> the sequence in an Arraylist.
 - (b) Create another method, *createCompliment*, which uses the read-in sequence and takes in no parameters, <u>iterates</u> through it, and creates its compliment by <u>checking</u> each letter. <u>Store</u> the compliment in an Arraylist. Remember that the compliment to "a" is "t" and "c" is "g". **The output should keep all strings as LOWERCASE.** HINT: Use the replace method for strings.
 - (c) Create **getCompliment** and **getDnaSequence** methods that return the corresponding Arraylist. So **getCompliment** should return the Arraylist created in createCompliment.
 - (d) Create a *mutateDna* method that will <u>loop</u> through the DNA sequence and change it in any way that you want. You can get <u>creative</u> on how this changes the DNA.
 - (e) Create a *printPeptide* method that <u>loops</u> through an Arraylist of amino acids and prints the sequence inside of it.
- 2. Create Mrna class.
 - (a) Create a *createCopy* method that reads in a DNA arraylist and <u>stores</u> the translated copy in an Arraylist. Remember that mRNA doesn't use "T", but instead the translation of "a" is "u". The output should keep all strings as LOWERCASE.
 - (b) Create a getMessengerDnaCopy that returns the arraylist with the copy of the DNA.
- 3. Create a Ribosome class.
 - (a) Create a *createProtein* static method that returns an ArrayList of amino acids. This should take in an Arraylist as a parameter.

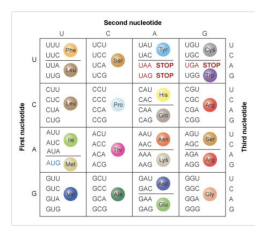


Figure 2: Proteing List For Ribosome

3 HL Extension

- 1. Create a Medicine class.
 - (a) Create a *cancerCure* method that checks if a sequence is the same as the original sequence from the txt file and then returns a fixed Arraylist with the correct sequence.