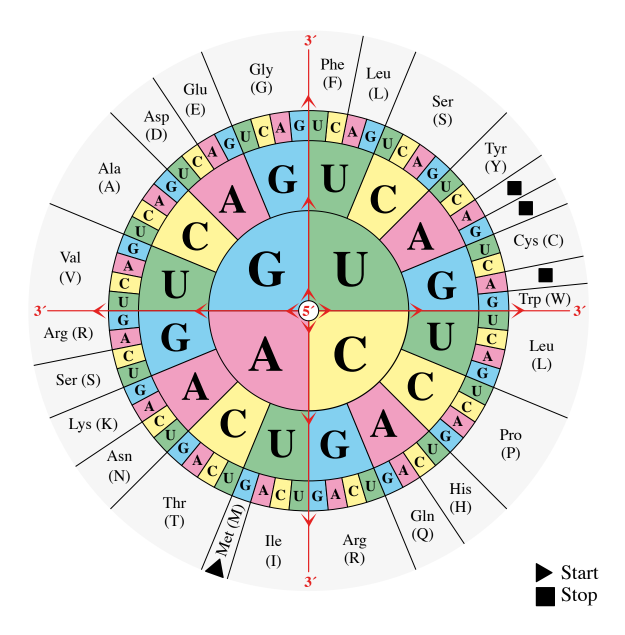
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# Lab 05 - String Operations on DNA / RNA

100pts each part (2 parts), 200 pts total.

Amino Acid Coding Table



Information that you need for this lab is in the lecture notes. https://github.com/Univ-Wyo-Education/F21-1010/blob/main/class/lect/Lect-11/Lect-11.html.pdf

### Part 1

Create a new file conv\_t\_to\_u.py.

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Write a python function that will:

1. Convert the input parameter to all lower case

- 2. Change all the t's to u's
- 3. Return the converted string.

This is very close to the example in class.

Use the example-conv\_t\_to\_u.py as a starting point.

Link on github.com: https://github.com/Univ-Wyo-Education/F21-1010/blob/main/lab/lab-

05/example-conv\_t\_to\_u.py

Input Example: aaTggTaaa

Output Example: aaugguaaa

Implement a test that verifies that this works. See example-conv\_t\_to\_u.py for testing code as well.

#### Part 2

Create a new file rna\_lookup.py.

Implement a function that uses a dictionary to lookup the amino acid that corresponds to a set of 3 RNA letters. The lookup takes in a 3 letter codon string and returns a single letter amino acid. If the three letter is not found then return '!'.

Example Input: "ccu"

Example Output: "P"

For starts codons, "aug", it returns 'M' for stop, "uaa", "uag", "uga" it returns '.'

Call the function rna\_to\_amino\_acid . Passing the 3 letter amino sequence.

Return the single letter coding for the amino acid as a string.

Use the example-rna\_lookkup.py as a starting point. It has the dictionary in it and some of the other code. You will need to implement some test cases. Look at the previous code and copy, then modify the tests to be appropriate for this.

Link on github.com: https://github.com/Univ-Wyo-Education/F21-1010/blob/main/lab/lab-05/example-rna\_lookkup.py

#### Part 3

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Download amino.py from https://github.com/Univ-Wyo-Education/F21-1010/blob/main/lab/lab-05/amino.py.

Run the file amino.py that imports the functions from Part 1 and Part 2 and calls them in order to produce the amino acid sequence.

**Expected Output:** 

RYPQPARPPGSDLRRTVLRLQSTTEICAEDNAAPPSRCSPGLC

Make sure you have all the files in the same folder.

### **Turn** in

Your 2 python files conv\_t\_to\_u.py and rna\_lookup.py.

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