# COM.e PROJECT: Investigating DNA in Python

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TARGET ASSESSMENT LEVEL: 3

# 1 Preface

[Jump ahead to the specification if this is boring.]

In Marjo Petäjäaho's BIO4 course, we are studying DNA and protein synthesis. Interestingly, DNA is the body's way of conveying information in code. After realizing that this project's purpose was text processing, I decided that processing theoretical DNA strands might be interesting. Of course, the project is highly theoretical, not taking mutations and specific complex processes into account; hence, it cannot be used in real-world applications. Nevertheless, it offers quite an interesting insight into how the body might theoretically process DNA.

# 2 Background Theory

DNA is a very complex molecule; however, the way it conveys information is quite understandable. DNA strands contain the organic bases adenine, cytosine, guanine, and thymine, commonly abbreviated as "A," "C," "G," and "T." These organic bases exhibit a characteristic: complementary pairing. A always bonds with T, and G always bonds with C.

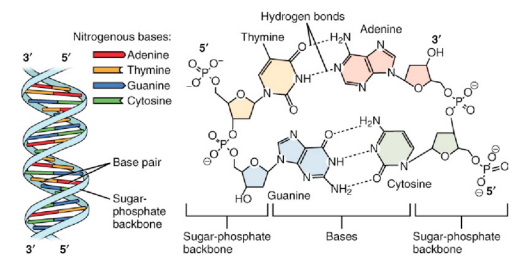


Figure 1: DNA Molecule [LibreTexts]

Complementary pairing can be used to our advantage. When DNA replicates, its two strands unwind. When one strand becomes split, transcription can take place. The singular strand pairs with its complementary counterpart. However, thymine is replaced by uracil (T becomes U). The newly formed strand is called mRNA, which is composed of many different triplets called codons. These codons are either code for a unique amino acid or punctuation. "Codon tables" organize this information (Figure 3).

Punctuation means that codons can indicate "START" or "STOP," signaling when to start or stop the amino acid chain. Translation is the process where amino acids connect to their codon counterparts and form amino acid chains. When these amino acids link up, they form proteins, which are fundamental to all bodily processes. This whole process is simply an algorithm where characters and strings are processed, making it relevant for a CS project [Britannica].

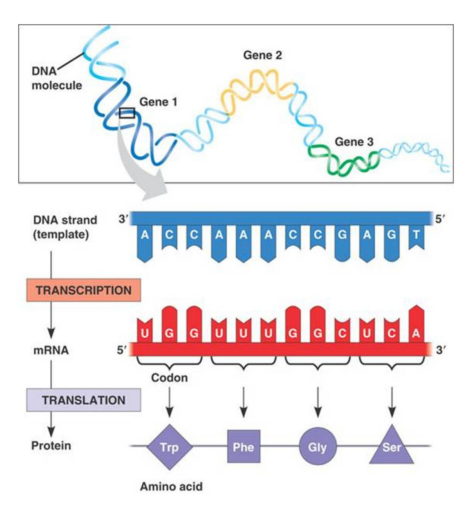


Figure 2: Transcription and Translation [Owlcation]

#### second letter U C G UCU UAU ) UGU Phe UUC UCC UAC J UGC C Ser **UUA UAA** stop **UCA UGA** stop Α uug j UCG , **UAG** stop UGG G Trp U CUU CCU CAU CGU His CAC CUC CCC CGC С Arg C Pro Leu Α CUA CCA CAA CGA first letter Gln G CAG CUG CCG CGG AUU ACU AAU AGU Asn Ser AGC С AAC **AUC** lle ACC Thr AUA . ACA AAA AGA Α AAG AGG G **AUG** ACG , Met GUU GCU GAU GGU U **GUC** GAC GGC C GCC G Val Gly Ala **GUA GCA** GAA **GGA** Α GAG GUG . GCG GGG. G

Figure 3: Codon Table [Vedantu]

# 3 Specification

### 3.1 What does the program do?

The program takes in different pieces of data. First, it takes in a filename where output will be stored (one for text in a .txt file). Then, the program asks the user whether they want a random DNA sequence or a custom one. If the user wants a random sequence, they will be asked for a length. A random sequence will then be generated. If the user chooses custom, they will be prompted to enter a custom sequence. In both cases, the user must enter their input in a specific format, following certain criteria. The random or custom sequence will then be transcribed (see Figure 2). During transcription, start codons are added, and stop codons are deleted. After transcription, the mRNA will be translated to its amino acid counterparts (see Figure 3). It does this with a dictionary in the file Codon\_Table.py. Finally, the amino acids will be strung together. All of these conversions are documented and written into a .txt file with the name that the user inputted.

#### 3.2 Data Format

The user can select any file name for their report (except "" or the absence of value). File extensions do not matter.

The user must enter "custom" or "random", where capitalization does not matter.

If "random" is chosen, a length  $l_1$  must be selected such that:

$$9 \le l_1 \le 30$$
 and  $l_1 \equiv 0 \pmod{3}$ .

If "custom" is chosen, the string must be composed only of the letters "A," "C," "G," and "T," and a length  $l_2$  must be selected such that:

$$9 \le l_2 \le 30$$
 and  $l_2 \equiv 0 \pmod{3}$ .

Capitalization does not matter in any case.

#### 4 Correctness

#### 4.1 Ideal Test Case

In this project's GitHub repository, I have linked some files documenting input and output cases. However, it must be noted that input occurs through the Python console, not through file uploads. Obviously, the date will vary, as it changes.

#### 4.1.1 Example Console Input

#### 4.1.2 Example Text File Output

```
1 REPORT

2
3 DATE: 2025-01-31

4
5 RANDOM BASE ARRANGEMENT: AGTCCCTCT

6
7 RNA sequence: UCAGGGAGA

8
9 Codons: ['AUG', 'UCA', 'GGG', 'AGA']

10
11 Amino Acids: ['Met', 'Ser', 'Gly', 'Arg']

12
13 FINAL CHAIN: Met-Ser-Gly-Arg

14
15 NOTE: NOT A FULLY ACCURATE REPRESENTATION. NOT APPLICABLE TO REAL LIFE CONTEXTS.

16
17 UTSAV CHOUDHURY 2025
```

#### 4.2 Resource Management

The input file is opened using a with-statement, and will therefore be closed automatically. This will conserve system resources

# References

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- [3] LibreTexts. 9.1: The Structure of DNA. 2017. https://bio.libretexts.org/Bookshelves/Introductory\_and\_General\_Biology/Conolecular\_Biology/9.01:  $_T$  he\_Structure\_of\_DNA
- [4] Rhys Baker. Protein Production: A Simple Summary of Transcription and Translation. 2012. https://owlcation.com/stem/protein-production-a-step-by-step-illustrated-guide
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