## CS217: Computer Programming Language: File I/O lab

Instructor: Jakramate Bootkrajang

### Outlines

- Uploading file to colab
- DNA sequence file
- Reading from file
- Exercises

# **Colab file uploading**

- In order to practice opening file we need a file to begin with
- You can upload files to Colab using the following code snippet

from google.colab import files
uploaded = files.upload()

• The uploaded file is only available for the current login session

### **DNA sequence file**

- Download the file from
- http://www.dalkescientific.com/writings/NBN/ data/10\_sequences.seq
- Save the file in your PC
- Upload the saved file to Colab using the code snippet in the previous slide

#### **Processing sequences stored in a file**

- We've previously worked with DNA sequence and did some interesting processing
- We will now generalise the process to cover multiple sequences.
- The sequences are stored in 10\_sequences.seq file

## **Reading lines from a file**

- Using open() function and loop over each line
- Document is a variable that stores input file handler
- Recall that file handler can be iterated

```
document = open("10_sequences.seq")
for line in document:
    print(line)
```

## More complex example

List the sequences starting with a thymine

```
document = open("10_sequences.seq")
for line in document:
   if line.startswith("T"):
      print(line)
```

- Each line contains additional '\n', what is its effect ?
- How to remove the extra '\n' ?

# rstrip() and lstrip()

- To remove extra white space at the end use rstrip() method
- To remove extra white space at the beginning of a string use lstrip() method

### **Exercise 1: Number lines in a file**

- Read the file 10\_sequences.seq. Print out the line number (starting with 1) then the line.
   Remember to use rstrip() to remove the extra newline.
- The output should look like this

1 CCTGTATTAGCAGCAGATTCGATTAGCTTTACAACAATTCAATAAAATAGCTTCGCGCTAA

- 2 ATTTTTAACTTTTCTCTGTCGTCGCACAATCGACTTTCTCTGTTTTCTTGGGTTTACCGGAA
- 3 TTGTTTCTGCTGCGATGAGGTATTGCTCGTCAGCCTGAGGCTGAAAATAAAATCCGTGGT
- 4 CACACCCAATAAGTTAGAGAGAGAGTACTTTGACTTGGAGCTGGAGGAATTTGACATAGTCGAT
- 5 TCTTCTCCAAGACGCATCCACGTGAACCGTTGTAACTATGTTCTGTGC
- 6 CCACACCAAAAAAACTTTCCACGTGAACCGAAAACGAAAGTCTTTGGTTTTAATCAATAA
- 7 GTGCTCTCTTCTCGGAGAGAGAGAGGTGGGCTGCTTGTCTGCCGATGTACTTTATTAAATCCAATAA
- 8 CCACACCAAAAAAACTTTCCACGTGTGAACTATACTCCAAAAACGAAGTATTGGTTTATCATAA
- 9 TCTGAAAAGTGCAAAGAACGATGATGATGATGATGAGAGGAACCTGAGCAGCCATGTCTGAACCTATAGC
- $10\ {\tt GTATTGGTCGTCGTGCGACTAAATTAGGTAAAAAAGTAGTTCTAAGAGATTTTGATGATTCAATGCAAAGTTCTATTAATCGTTCAATTG$

### **Exercise 2**

• List the sequences in 10\_sequences.seq which have the pattern CTATA

### **Exercise 3**

• Modify the previous program to print the index of the first time CTATA pattern is found

#### **Exercise 4**

- Based on sequences.seq write a program to answer the following questions
- 1) How many sequences are in that file ?
- 2) How many have the pattern CTATA ?
- 3) How many have more than 500 bases ?
- 4) How many have over 50% GC composition ?

GC composition is the ratio of the number of G and C over the length of the sequence

### References

http://www.dalkescientific.com/writings/NBN/