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## Homework 8

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This homework is due on April 4, 2017 at 7:00pm. Please submit as a PDF file on Canvas. Before submission, please re-run all cells by clicking "Kernel" and selecting "Restart & Run All."

**Problem 1 (5 points):** In bioinformatics, k-mers refer to all the possible subsequences (of length k) from a read obtained through DNA sequencing. For example, if the DNA sequencing read is "ATCATCATG", then a the 3-mers in that read include "ATC" (which occurs twice), "TCA" (which occurs twice), "CAT" (occurs twice), and "ATG" (occurs once). You can read more about k-mers on Wikipedia (https://en.wikipedia.org/wiki/K-mer).

a) Write a function that takes a string of nucleotides as input and returns a **dictionary** with all 3-mers present in that string, and the number of times that each 3-mer occurs. Then use your function to find the 3-mers in the DNA sequence my seq1 defined below.

The output of your function should be a dictionary that is structured like this (although it will have several more entries):

```
{"ATC": 3, "ATG": 2, "TGA": 3}
```

where each key is a 3-mer itself (e.g., "ATC") and each value is the number of times that 3-mer occurs.

b) For the sequence my\_seq2 defined below, verify manually that your function generates the correct result, and explain your reasoning in 2-3 sentences.

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In [1]: # Find all 3-mers in these two sequences
        my seq1 = "CAGCCCAATCAGGCTCTACTGCCACTAAACTTACGCAGGATATATTTACGCCGACGTACT"
        my_seq2 = "ATCATCATG"
        # Your code goes here
        #define nucleotide function
        def nucleotides(string):
            num=len(string)
            num range= range(len(string))
            nucleotide_list= {}
            for a in num range:
                if a <= (num-3):
                    sequence = string[a: a+3] #"sequence" takes a length of a codon
                    if sequence in nucleotide list:
                        nucleotide_list[sequence] += 1 #a 3-mer that is already in t
                    else:
                        nucleotide list[sequence]=1 # a new 3-mer is added to the di
            return nucleotide list
        print("The 3-mers and the counts of 3-mers for sequence 1 are:", nucleotides
        print("The 3-mers and the counts of 3-mers for sequence 2 are:", nucleotides
```

```
The 3-mers and the counts of 3-mers for sequence 1 are: {'CAG': 3, 'AGC': 1, 'GCC': 3, 'CCC': 1, 'CCA': 2, 'CAA': 1, 'AAT': 1, 'ATC': 1, 'TCA': 1, 'AGG': 2, 'GGC': 1, 'GCT': 1, 'CTC': 1, 'TCT': 1, 'CTA': 2, 'TAC': 4, 'ACT': 4, 'CTG': 1, 'TGC': 1, 'CAC': 1, 'TAA': 1, 'AAA': 1, 'AAC': 1, 'CTT': 1, 'TTA': 2, 'ACG': 3, 'CGC': 2, 'GCA': 1, 'GGA': 1, 'GAT': 1, 'ATA': 2, 'TAT': 2, 'ATT': 1, 'TTT': 1, 'CCG': 1, 'CGA': 1, 'GAC': 1, 'CGT': 1, 'GTA': 1}

The 3-mers and the counts of 3-mers for sequence 2 are: {'ATC': 2, 'TCA': 2, 'CAT': 2, 'ATG': 1}
```

When I put the sequence of my\_seq2 into the function, the function is able to take the sequence, find the length and range of the sequence, and within the sequence find all the 3-mer's and put the sequences in the dictionary and returns the dictionary. Then the second if statement determines if a certain 3-mer has been repeated or not in the list, and collects how many times a certain 3-mer has been counted within myseq2, which is collected under the dictionary "nucleotide\_list" and is returned.

**Problem 2 (5 points):** DNA sequences are typically stored in a format called FASTA (pronounced fast-ay). A single FASTA file may contain many different sequences. For example, you may have a FASTA file for a mouse, and each mouse gene sequence is stored as a separate sequence in that FASTA file. All sequences in a FASTA file begin on a new line with a greater-than symbol ">" (without quotes).

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Write a function that takes the *name* of a FASTA file as input, opens that file, counts the number of sequences in the file (by counting the number of lines in the file that start with a ">" symbol), and returns the count. Download the file "CD4.fasta"

(http://wilkelab.org/classes/SDS348/2017 spring/homeworks/CD4.fasta)" to your computer and use your function to count the number of sequences in the file. The file CD4.fasta contains amino acid sequences of the CD4 membrane protein that is found on the surface of the immune cells.

The number of sequences is 18