Biological Computation 2022

<u>חישוב ביולוגי</u>

Exercise #1

<u>הגשה בזוגות עד לתאריך 6.6.22</u>

<u>Note</u>: You can discuss the work with other students however you should write all code by yourself / with your exercise partner. Please document your code and also prepare a short (no more than 2 pages) explanation on how the code works and what were the main design and implementation decisions you made. If any code that a student did not write is used, explain where it is taken from and why it's needed.

*In addition to submitting to Moodle prepare a GitHub repository with a readme on how to run the code.

1) a) Write a program (in your favorite programming language) that gets as input a positive integer n and generates all connected sub-graphs of size n.

The output should be a textual file of the following form:

n=2

count=2

#1

12

#2

12

2 1

The first two lines output n and the total number (count) of different sub-graphs of size n. Then the sub-graphs themselves are given each starting with a line labelled #k for motif number followed by all edges, each line i j means an edge from source i to target j.

b) Output the result of your program for n = 1 to 4.

- c) What is the maximal number n for which your program can complete successfully within no more than 1 hour of computing time?
- d) What is the maximal number n for which your program cancomplete successfully within 2,4,8 hours of computing time?
- 2) Write a program that gets as input positive integer n and a graph of the format:
- 12
- 23
- 14

The graph in the example contains 4 vertices 1,2,3,4 and directed edges (1,2) (2,3) (1,4). The program should output all sub-graphs of size n and count how many instances appear of each motif. The format of the output of the identified sub-graphs should be like in question 1, where in the line after #k should appear the count of number of instances, count=m if the motif appears m times. Output count=0 if a motif does not appear in the graph.