Dear ODTÜClass Users,

There will be maintenance work at Turnitin on January 27, 2024 between 19:30 - 23:30.

Therefore, we recommend that you do not add assignments with a deadline of January 27, 2024.

Best regards, **ODTÜClass Support Team** 

# [CENG 315 ALL Sections] Algorithms

<u>Dashboard</u> / My courses / <u>571 - Computer Engineering</u> / <u>CENG 315 ALL Sections</u> / <u>December 3 - December 10</u> / <u>THE6</u>

Description

Submission view

# THF6

Available from: Saturday, December 16, 2023, 12:00 PM

Due date: Sunday, December 17, 2023, 11:59 PM

**▼ Requested files**: the6.cpp, test.cpp, the6.h (**L** <u>Download</u>)

Type of work: 4 Individual work

#### EDIT: If you are planning to work on your local, you can find the sample io here.

Input files are formatted as follows:

<node\_count> <edge\_count> followed by the <vertex1> <vertex2> <weight> lines for the bond\_energies graph

<node\_count> <edge\_count> followed by the <vertex1> <vertex2> <weight (always1)> lines for the molecule\_structure graph

**Output** files are formatted as follows:

<total\_lowest\_energy>

<longest\_chain>

Only numerical values are compared as fixed outputs. Since the structures may have possible correct answers, they are evaluated by the solver in the VPL.

You, an Al hater, recently learned that an Al model discovered new molecule structures equivalent to 800 years' worth of knowledge[1][2], and you are extremely annoyed by this news. While on an anger rant, you claim that you can write a basic program to find new optimal structures by yourself. Your chemist friends dare you to do so. They also offer to check the validity of your program, if you also write them another program that helps with their project.

You come up with an idea to find new molecule structures: If we have a graph where each vertex is an atom and each edge is the energy of the bond between the two atoms, maybe a new possible molecule structure can be found by selecting the edges with lowest energy bonds, without creating a cycle. Your chemist friends tell you this is most definitely a wrong approach, but you insist on trying.

Additionally, your chemist friends need a program to find the longest chain in a given molecule, so you agree to also write that program for them.

#### **Problem**

This exam consists of two parts, graded independently.

#### > PART1

You need to complete the function *find\_stucture()* which returns the total bond energy of the found molecule:

```
int find_structure(std::vector< std::vector<std::pair<int,int> > > &bond_energies,
std::vector< std::vector<std::pair<int,int> > &lowest_energy_structure);
```

- **bond\_energies**: a bidirectional graph represented by an adjacency list, where the vertices **(v)** are atoms, edges are bonds, and weights **(w)** of edges are the energies of the bonds. In other words, an entry in the adjacency list for vertex v0 is <v1, w>, which represents an edge between v0-v1 with a weight of w.
- **lowest\_energy\_structure**: the found molecule structure represented as an adjacency list, with the same format as **bond\_energies**. You are expected to find the structure and assign it to this argument.
- There is at most one bond with two atoms with only one weight value, i.e. there is at most one undirected edge between any two vertices in the graph, meaning at most one weight value for each pair. Undirectionality is shown in the adjacency list by adding two mirror entries for each edge for ease of implementation.
- Vertices are represented as integers starting from 0, and the maximum number of vertices in the graph is 1000.
- Weights are represented as integers starting from 1, and the maximum weight value is 100.
- · return value is the total bond energy of the lowest\_energy\_structure.

#### > PART2

You need to complete the function *find\_longest\_chain()* which **returns the number of the atoms in the longest chain** of the given molecule

int find\_longest\_chain(std::vector < std::vector < std::pair < int,int > > & molecule\_structure, std::vector < int > chain);

- molecule\_structure: a graph with the same representation and limits as the bond\_energies and lowest\_energy\_structure arguments of PART 1. Additionally, it is guaranteed that the structure is a connected, acyclic, undirected graph, with all edges having the same weight (i.e. weights are not important). Undirectionality is shown in the adjacency list by adding two mirror entries for each edge for ease of implementation.
- **chain:** a vector of integers, where each integer maps to a vertex ID in the found longest chain. The vector should follow the order of the chain. The validity of the chain will be checked by the tester to see if consecutive vertices in the vector have edges between them or not.
- return value is the total number of atoms in the longest chain of the molecule. The longest chain is defined as the count of vertices on the path between the two farthest vertices in the graph, including the start and end vertices.

#### **Constraints and Hints:**

- When deciding between edges with the same weights connecting v to two different vertices u\_1 and u\_2, always favor the u\_i with a smaller index.
- Similarly, if you need to decide between two edges (u1,u3,w1) and (u2,u4,w2) where w1=w2, you should again **favor the edge with the** smaller u i, meaning (u1,u3,w1).
- There are **no self-loops**, meaning there is no edge such that  $(u_i, u_j, w)$  where i=j.

#### **Evaluation:**

• After your exam, black-box evaluation will be carried out. You will get full points if you return the correct order or strongly connected components. The grade you see in the VPL contains 50% of your final grade. We will evaluate your grades with different inputs after the end of the exam.

#### **Grade distribution** is as follows:

- 30% part1 return value lowest\_total\_energy
- 30% part1 lowest\_energy\_structure
- 10% part2 return value *longest\_chain\_size*
- 30% part2 *chain*

# **Example IO:**

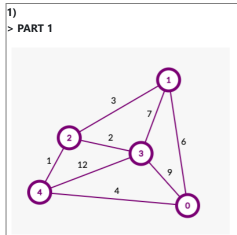


fig 1: bond\_energies graph

# bond\_energies:

- { (1,6) (3,9) (4,4) }
- { (0,6) (2,3) (3,7) }
- { (1,3) (3,2) (4,1) }
- 2 { (0,9) (1,7) (2,2) (4,12) }
- { (0,4) (2,1) (3,12) }

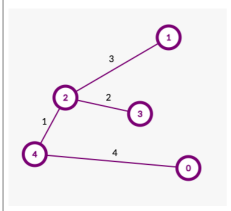


fig 2: lowest\_energy\_structure

# lowest\_energy\_structure found:

- { (4,4) }
- { (2,3) }
- 2 { (1,3) (4,1) (3,2) }
- 3 { (2,2) }
- { (2,1) (0,4) }

# return value *lowest\_total\_energy*: 10

#### > PART 2

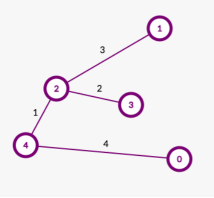


fig3: molecule\_structure

**note:** for the first i/o, input of part2 is the same as the output of part1. this is not guaranteed to be the case for all outputs. please check other i/o.

### molecule\_structure:

{ (4,4) }

```
1 { (2,3) }
2 { (1,3) (3,2) (4,1) }
3 { (2,2) }
4 { (0,4) (2,1) }

chain:
(3)-(2)-(4)-(0) and alternatively, (1)-(2)-(4)-(0)

return value longest_chain_size: 4
```

#### **Specifications:**

- There are 2 tasks to be solved in **36 hours** in this take-home exam.
- You will implement your solutions in *the6.cpp* file.
- You are free to add other functions to the6.cpp
- Do not change the first line of the6.cpp, which is #include "the6.h"
- Some libraries are included in "the6.h" for your convenience, you can use them freely.
- Do not change the arguments and the return value of the functions find\_stucture() and find\_longest\_chain() in the file the6.cpp
- Do not include any other library or write include anywhere in your the6.cpp file (not even in comments).
- You are given **test.cpp** file to test your work on **ODTUClass** or your **locale**. You can, and you are, encouraged to modify this file to add different test cases.
- If you want to test your work and see your outputs you can compile your work on your locale as:

```
>g++ test.cpp the6.cpp -Wall -std=c++11 -o test
> ./test
```

- You can test your the6.cpp on the virtual lab environment. If you click **run**, your function will be compiled and **executed with test.cpp**. If you click **evaluate**, you will get **feedback** for your current work, and your work will be **temporarily graded** for a limited number of inputs.
- The grade you see in lab is not your final grade, your code will be reevaluated with different inputs after the exam.

The system has the following limits:

- a maximum execution time of 8 seconds (your program needs to return in less than a second per test case on average each test case has 2 parts)
- a 1 GB maximum memory limit,
- an execution file size of 4M.
- Solutions with longer running times will not be graded.
- If you are sure that your solution works in the expected complexity, but your evaluation fails due to limits in the lab environment, the constant factors may be the problem.

\*not relevant to the question:

- [1] https://deepmind.google/discover/blog/millions-of-new-materials-discovered-with-deep-learning/
- [2] https://www.nature.com/articles/s41586-023-06735-9

# Requested files

## the6.cpp

```
#include "the6.h"

// do not add extra libraries here

int find_structure(std::vector< std::vector<std::pair<int,int> > % bond_energies, std::vector< std::vector<std::pair<int,int> > % int lowest_total_energy = 0;

return lowest_total_energy;
}

int find_longest_chain(std::vector< std::vector<std::pair<int,int> > % molecule_structure, std::vector<int>% chain){
    int longest_chain_size = 0;
    return longest_chain_size;
}
```

test.cpp

```
#include <iostream>
      #include <fstream>
      #include "the6.h"
      void print_adj_list(std::vector< std::vector< std::pair<int,int> > >& adj_list) {
          int N = adj_list.size();
if (N == 0) {
 8
               std::cout << "list is empty!" << std::endl;
10
11
          }
12
          for (int v=0;v<N;v++) {
   std::cout << v << "\t{";
   for (auto p : adj_list[v]) {
      std::cout << " (" << p.first << "," << p.second << ")";</pre>
14
15
16
17
               std::cout << " }\n";
18
19
20
          return;
21
     }
22
23
      // you can use this if you want to print the adj list as a matrix
24
      void print_adj_list_as_matrix(std::vector< std::vector< std::pair<int,int> > >& adj_list) {
          int N = adj_list.size();
if (N == 0) {
25
26
               std::cout << "list is empty!" << std::endl;
28
29
          int** matrix;
 30
          matrix = new int*[N];
31
          for(int temp=0; temp < N; temp++) matrix[temp] = new int[N];
for (int i=0; i<N; i++){
    for (int j=0; j<N; j++){
        matrix[i][j] = -1; // no edge</pre>
32
33
35
36
               }
37
          }
39
          for (int i=0; i<N; i++){
  for (std::pair<int,int> x: adj_list[i]) {
40
                   matrix[i][x.first] = x.second;
42
43
          }
          for (int i=0; i<N; i++){
   for (int j=0; j<N; j++){
      if (matrix[i][j] == -1) std::cout << "- ";</pre>
45
46
                    else std::cout << matrix[i][j] << " ";
49
               std::cout << std::endl:
50
51
          for(int i=0; i<N; i++) delete[] matrix[i];</pre>
53
54
          delete[] matrix;
56
     }
57
58
      void read_from_file(std::vector< std::vector<std::pair<int,int> > >& bond_energies, std::vector< std::vector<std::pair<int,int> >
59
           char addr[]= "inp01.txt"; // 01-05 are available
          std::ifstream infile (addr);
60
          61
63
64
65
67
          int V_p1, E_p1, V_p2, E_p2;
68
          infile >> V_p1 >> E_p1;
          bond_energies.resize(V_p1);
for (int l=0; l<E_p1; l++) {
70
71
 72
               int v1, v2, w;
               infile >> v1 >> v2 >> w;
bond_energies[v1].push_back(std::make_pair(v2,w));
73
74
               bond_energies[v2].push_back(std::make_pair(v1,w));
 75
76
77
          infile >> V_p2 >> E_p2;
78
          molecule_structure.resize(V_p2);
80
          for (int l=0; l<E_p2; l++) {
               int v1, v2, w;
infile >> v1 >> v2 >> w;
81
82
83
               molecule_structure[v1].push_back(std::make_pair(v2,w));
84
               molecule_structure[v2].push_back(std::make_pair(v1,w));
85
87
          infile.close();
88
     }
90
91
      int main(){
          std::vector< std::pair<int,int> > bond_energies;
92
93
          std::vector< std::pair<int,int> >> molecule_structure;
94
           std::vector< std::pair<int,int> > lowest_energy_structure;
           std::vector<int> chain;
95
          int longest_chain_size, lowest_total_energy;
97
98
          read_from_file(bond_energies, molecule_structure);
99
          lowest_energy_structure.resize(bond_energies.size());
100
          lowest_total_energy = find_structure(bond_energies, lowest_energy_structure);
std::cout << "PART 1: " << std::end1 << "Bond energy graph:" << std::end1;</pre>
101
102
          print_adj_list(bond_energies);
```

```
104
         std::cout << "Graph of the lowest energy structure found:" << std::endl;</pre>
         print_adj_list(lowest_energy_structure);
std::cout << "Total energy of the lowest energy structure: " << lowest_total_energy << std::endl;</pre>
105
106
        108
109
110
111
112
113
114
115
            std::cout << " - (" << chain[i] << ")";
116
117
118
         return 0;
119
```

### the6.h

```
#ifndef THE6_THE6_H
#include <vector>
#include <vector>
#include <queue>
#include <climits>
#include <climits>
#include <algorithm>

//updating this file will not change the execution in the VPL

int find_structure(std::vector< std::vector<std::pair<int,int> >> &bond_energies, std::vector< std::vector<std::pair<int,int> >> &molecule_structure, std::vector<int>& chain);

#endif //THE6 THE6 H
#endif //THE6 THE6 H
```

**VPL** 

You are logged in as aytac sekmen (Log out)

CENG 315 ALL Sections

**ODTÜClass Archive** 

2022-2023 Summer

2022-2023 Spring

2022-2023 Fall

2021-2022 Summer

2021-2022 Spring

2021-2022 Fall

2020-2021 Summer

2020-2021 Spring

2020-2021 Fall

Class Archive

Get the mobile app







