Bioinformatics III Fourth Assignment

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Exercise 4.1: Dijkstra's algorithm for finding shortest paths

(a) Draw a directed or undirected graph with at least one negative edge weight for which Dijkstra's algorithm does not find the shortest path from some node s to another node t. Use your example to explain why Dijkstra's algorithm only works on graphs with non-negative edge weights.

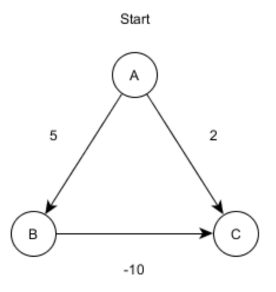


Figure 1: Example directed graph

A-B-C is the shortest path.

We have V = A, B, C, E = (A, C, 2), (A, B, 5), (B, C, -10). So, A-C is found first but not A-B-C. Dijkstra assume that the minimality will never change when "closing" a node. The use of negative numbers change this rule and so is not compatible with Dijkstra.

(b) Dijkstra's algorithm modifications

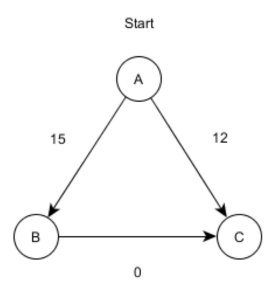


Figure 2: Graph with edge cost ≥ 0 .

We can see that adding 10 to all weights doesn't work. A-C is still the path that Dijkstra find first and in this case, it is the shortest path which was not the case before.

(c) Could BFS be used to find the shortest paths between nodes? If so, what would the edge weights have to look like for BFS to be guaranteed to find the shortest paths between nodes? Why (not)?

With BFS, we can find a path with arbitrary weights but it is not guaranteed to be optimal.

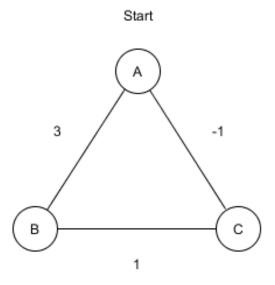


Figure 3: When we want to go from A to B, the shortest path is A - C - B, but with BFS we will first expand A and find the path A-B. When we expand the C node, B is already marked as "duplicate". BFS will ignore the link between B and C in this case.

Exercise 4.2: Force directed layout of networks

(a) Implementation preparation

For the harmonic force we have:

$$F_h(\vec{r}) = -\nabla E_h(\vec{r})$$

According to the definition:

$$\begin{split} &= -\nabla\frac{k}{2} \left\| \vec{r} \right\|^2 = -\frac{k}{2} \nabla \left\| \vec{r} \right\|^2 \\ &= -\frac{k}{2} \nabla (\sqrt{x^2 + y^2 + z^2})^2 \\ &= -\frac{k}{2} \nabla (x^2 + y^2 + z^2) \\ &= -\left(\begin{matrix} kx \\ ky \\ kz \end{matrix} \right) \end{split}$$

For the Coulomb force we have:

$$F_c(\vec{r}) = -\nabla E_c(\vec{r})$$

$$= -\nabla \left(\frac{1}{4\pi\epsilon_0} \frac{q_1 q_2}{\|\vec{r}\|}\right)$$

$$= -\nabla \left(\frac{1}{4\pi\epsilon_0} \frac{q_1 q_2}{\sqrt{x^2 + y^2 + z^2}}\right)$$

$$= -\frac{q_1 q_2}{4\pi\epsilon_0} \nabla \left(\frac{1}{\sqrt{x^2 + y^2 + z^2}}\right)$$

We apply partial derivatives on $\frac{1}{\sqrt{x^2+y^2+z^2}}$ (chain rule):

$$=-\frac{q1q2}{4\pi\epsilon_0}\begin{pmatrix}-\frac{x}{(x^2+y^2+z^2)^{\frac{3}{2}}}\\-\frac{y}{(x^2+y^2+z^2)^{\frac{3}{2}}}\\-\frac{z}{(x^2+y^2+z^2)^{\frac{3}{2}}}\end{pmatrix}=\frac{q1q2}{4\pi\epsilon_0}\begin{pmatrix}\frac{x}{(x^2+y^2+z^2)^{\frac{3}{2}}}\\\frac{y}{(x^2+y^2+z^2)^{\frac{3}{2}}}\\\frac{z}{(x^2+y^2+z^2)^{\frac{3}{2}}}\end{pmatrix}$$

(b) Adapting the energy equations for networks

For the Harmonic force we have:

$$F_h(\vec{r}_{ij}) = -\nabla E_h(\vec{r}_{i,j}) = -\begin{pmatrix} x_i - y_j \\ y_i - y_j \end{pmatrix}$$

For the Coulomb force:

$$F_c(\vec{r_{ij}}) = k_1 k_2 \left(\frac{\frac{x_i - x_j}{((x_i - x_j)^2 + (y_i - y_j)^2)^{\frac{3}{2}}}}{\frac{y_i - y_j}{((x_i - x_j)^2 + (y_i - y_j)^2)^{\frac{3}{2}}}} \right)$$

(c) Understanding the Coulomb and harmonic energy: How does the Coulomb energy and harmonic energy change if the degree of both nodes is increased or decreased? What happens if the distance between two nodes is increased or decreased?

If the degree of both nodes increases, the harmonic force is not affected but Coulomb force will increase with a factor $-k_1k_2$ as seen in (b).

However, if the distance increase, both Coulomb and harmonic will be affected. The Harmonic energy is $\frac{1}{2}||\vec{r^2}||^2$ and the Coulomb $\frac{k_ik_j}{||\vec{r}_{ij}||}$ so if we increase the distance, Coulomb energy is going to decrease and Harmonic energy to increase.

(d) Understanding the forces: Why is the Coulomb force the repulsive force and the harmonic force the attractive force?

In our case, all the node will have a positive charge (the degree) which means that they are all going to repel each other. However, the objective is to keep connected nodes close to each other and at the same time spread the rest of the graph away to give it a nice display. The Coulomb force is the basic force between every node according to their degrees and distance, disregarding their connection. The Harmonic force is applied here only when two nodes are connected to each other and because it has opposite sign, as seen in (b), it will temper the force between the two connected nodes and allow to keep them close ot each other.

(e) Implementing the force directed layout algorithm

```
Listing 1: layout main.py
o from layout import Layout
  from tools import plot_layout, plot_energies
   file_paths = ['star.txt', 'square.txt', 'star++.txt', 'dog.txt']
5 file_paths = ['dog.txt']
   for file_path in file_paths:
       # read the file into your layout class
        layout = Layout(file_path)
10
        \# run the normal layout for 1000 iterations and store the total energies
       \# plot the normal layout
        energies1 = layout.layout(1000)
15
        plot_layout(layout, file_path + "_-_Normal_Layout")
        \# run the simulated annealing layout for 1000 iterations and store the
             total\ energies
        # plot the simulated annealing layout
20
       \begin{array}{ll} energies 2 &= layout.simulated\_annealing\_layout (1000) \\ plot\_layout (layout, file\_path + "\_-\_Simulated\_Annealing\_Layout") \end{array}
        \# plot the total energies of the normal layout and the simulated annealing
              layout
25
         \begin{array}{c} \texttt{plot\_energies} \ ( \, [ \ energies1 \ , \ energies2 \, ] \ , \quad ["\ Normal\_layout" \ , \quad "Simulated\_Annealing\_Layout" \, ] \ , \quad "Layout\_energies" \, ) \end{array} 
                                     Listing 2: layout.py
o from random import gauss
  \mathbf{import} \hspace{0.2cm} \mathrm{math}
   import random
   import itertools
  from generic_network import GenericNetwork
   class Layout:
        \mathbf{def} \ \_\_init\_\_(self , \ file\_path):
             :param file_path: path to a white-space-separated file that contains
10
            node interactions
            # create a network from the given file
             self.network = GenericNetwork()
             self.network.read_from_tsv(file_path)
            \#\ friction\ coefficient
15
             self.alpha = 0.03
            # random force interval
            self.interval = 0.3
            # initial square to distribute nodes
             self.size = 50
20
        def init_positions(self):
             Initialise or reset the node positions, forces and charge.
25
            # Set up the positions and charge
            for key, node in self.network.nodes.items():
```

```
# Pick a coordinate between 1 and 50 (initial square)
30
                                node.pos_x = random.randint(1, self.size + 1)
                                node.pos_y = random.randint(1, self.size + 1)
                                # The charge is the node degree
                                node.charge = node.degree()
35
                       # Calculate the forces
                       self.calculate_forces()
              def calculate_forces (self):
40
                        Calculate the force on each node during the current iteration.
                       # For all pair of nodes...
45
                       for pair in itertools.combinations(self.network.nodes.items(), 2):
                                node1 = pair[0][1]
                                node2 = pair[1][1]
50
                                \begin{array}{lll} coulombx = (node1.charge * node2.charge) * ((node1.pos_x - node2.\\ &pos_x)/((node1.pos_x - node2.pos_x)**2 + (node1.pos_y - node2.\\ \end{array}
                                         pos_y)**2)**(3/2)
                                coulomby = (node1.charge * node2.charge) * ((node1.pos_y - n
                                         pos_y)/((node1.pos_x - node2.pos_x)**2 + (node1.pos_y - node2.pos_x)
                                         pos_y)**2)**(3/2)
                                harmonicx = 0.0
                                harmonicy = 0.0
55
                                # If the nodes are connected, we temper the force with the
                                         harmonic
                                if node1.has_edge_to(node2):
                                         harmonicx = -(node1.pos_x - node2.pos_x)
                                         harmonicy = -(node1.pos_y - node2.pos_y)
60
                                # Add the force to node1 .... opposite to node2
                                fx = coulombx + harmonicx
fy = coulomby + harmonicy
                                node1.force_x += fx
65
                                node1.force_y += fy
                                node2.force_x = fx
                                node2.force_y -= fy
70
              def add_random_force(self, temperature):
                       Add\ a\ random\ force\ within\ [-\ temperature\ *\ interval\ ,\ temperature\ *
                                interval | to each node.
                        (There is nothing to do here for you.)
                        : param\ temperature:\ temperature\ in\ the\ current\ iteration
75
                       for node in self.network.nodes.values():
                                node.force_x += gauss(0.0, self.interval * temperature)
                                node.force_y += gauss(0.0, self.interval * temperature)
80
              def displace_nodes(self):
                       Change the position of each node according to the force applied to it
                       and reset the force on each node.
85
                       for node in self.network.nodes.values():
                                node.pos_x = node.pos_x + node.force_x * self.alpha
                                node.pos_y = node.pos_y + node.force_y * self.alpha
```

```
# Reset the forces to 0
90
               node.force_x = 0
               node.force_y = 0
       def calculate_energy(self):
95
           Calculate the total energy of the network in the current iteration.
           :return: total energy
           energy_total = 0
100
           for pair in itertools.combinations(self.network.nodes.values(), 2):
               node1 = pair[0]
               node2 = pair[1]
105
               # Coulomb energy
               Ec = (node1.degree() * node2.degree()) / (math.sqrt(((node1.pos_x
                    - (node2.pos_x)**2) + ((node1.pos_y - node2.pos_y)**2)))
               #Harmonic energy
               Eh = 0
110
               if node1.has_edge_to(node2):
                   Eh = ((node1.pos_x - node2.pos_x) **2 + (node1.pos_y - node2.
                        pos_y)**2)/2
               energy_total += Ec + Eh
115
           return energy_total
       def layout(self , iterations):
120
           Executes the force directed layout algorithm. (There is nothing to do
               here for you.)
           :param iterations: number of iterations to perform
           :return: list of total energies
           # initialise or reset the positions and forces
125
           self.init_positions()
           energies = []
           for _ in range(iterations):
               self.calculate_forces()
130
               self.displace_nodes()
               energies.append(self.calculate_energy())
           return energies
135
       def simulated_annealing_layout(self, iterations):
           Executes the force directed layout algorithm with simulated annealing.
           :param iterations: number of iterations to perform
           :return: list of total energies
140
           self.init_positions()
           energies = []
           temperature = 100000
           for i in range(iterations):
145
               # TODO: DECREASE THE TEMPERATURE IN EACH ITERATION. YOU CAN BE
                   CREATIVE.
               # Decrease the temperature of 80% at each loop. (It decreases a
                   first\ time\ before\ being\ used...)
               temperature = 0.2 * temperature
               # there is nothing to do here for you
150
               self.calculate_forces()
```

```
self.add_random_force(temperature)
                self.displace_nodes()
                energies.append(self.calculate_energy())
155
           return energies
                                 Listing 3: node.py
 o class Node:
       def __init__(self , identifier):
            self.identifier = identifier
           # contains the identifiers of other nodes connected to this node
           self.neighbour\_nodes = set()
 5
           # fields for the layout algorithm
           self.pos_x = 0.0
           self.pos_y = 0.0
           self.force_x = 0.0
10
           self.force_y = 0.0
           self.charge = 0
       \mathbf{def} = -eq_{--}(self, node):
            :param node: Node-object
15
           :return: True if the other node has the same identifier, False
               otherwise
           return self.identifier == node.identifier
       def __str__(self):
20
            : return: \ string \ representation \ of \ the \ node \ identifier
           return str (self.identifier)
25
       def has_edge_to(self, node):
            : param \ node: \ Node-object
           return: True if this node has an edge to the other node, False
               otherwise
30
           return node.identifier in self.neighbour_nodes
       def add_edge(self, node):
           Adds an edge to the other node by adding it to the neighbour-nodes.
35
           :param node: Node-object
            self.neighbour_nodes.add(node.identifier)
40
       def remove_edge(self, node):
           Removes the edge to the other node, if that edge exists, by removing
               the other node from the neighbour nodes.
            :param node: Node-object
            self.neighbour_nodes.discard(node.identifier)
45
       def degree(self):
           :return: the degree of this node (= number of neighbouring nodes)
           return len(self.neighbour_nodes)
                                 Listing 4: tools.py
 o import matplotlib.pyplot as plt
   from itertools import combinations
```

```
def plot_layout(layout, title):
5
       Plots the layout of a network.
       : param \ layout: \ Layout-object
       :param title: plot title
       for node_1, node_2 in list(combinations(layout.network.nodes.values(), 2))
10
           if layout.network.has_edge(node_1, node_2):
                \# plot the edge between the two nodes, if it exists
                plt.plot\left(\left[\:node\_1.pos\_x\:,\:\:node\_2.pos\_x\:\right]\:,\:\:\left[\:node\_1.pos\_y\:,\:\:node\_2.pos\_y\:\right]
                    ], linestyle='-', color='black')
15
       # plot the nodes
       for node in layout.network.nodes.values():
           plt.plot(node.pos_x, node.pos_y, marker='H', color='red')
       # set the title, clean up the plot layout and show it
       plt.title(title)
20
       plt.tight_layout()
       plt.show()
       plt.clf()
  def plot_energies(energy_lists, legend, title):
       Plots\ list(s)\ of\ total\ energies .
       :param energy_lists: a list that contains a list of total energies
       :param legend: a list of curve labels
30
       : param \quad title: \quad plot \quad title
       # plot each list of total energies
       for energy_list in energy_lists:
           plt.plot(energy_list)
35
       \# set the x-axis and y-axis labels
       plt.xlabel('iteration')
plt.ylabel('Total_Energy')
40
       # set the legend, title, clean up the plot layout and show it
       plt.legend(legend)
       plt.title(title)
       plt.tight_layout()
       plt.show()
45
       plt.clf()
                             Listing 5: generic network.py
o from node import Node
  class GenericNetwork:
       \mathbf{def} __init__(self):
           # key: node identifier, value: Node-object
5
           self.nodes = \{\}
       def read_from_tsv(self, file_path):
           Reads white-space-separated files that contain two or more columns.
10
                The first two columns contain the
            identifiers of two nodes that have an undirected edge. The two nodes
                are added to the network.
            : param \ file\_path: \ path \ to \ the \ file
           \# clear the prior content of the network
           self.nodes = \{\}
```

```
# open the file for reading
           with open(file_path, 'r') as file:
               # iterate over the lines in the file
20
               for line in file:
                    columns = line.split()
                    # skip lines that do not have two node identifiers
                    if len(columns) < 2:
25
                        continue
                    # create the two nodes and remove potential whitespace such as
                         new-line\ from\ their\ identifiers
                    node_1 = Node(columns[0].strip())
                    node_2 = Node(columns[1].strip())
30
                    # add the nodes and the edge between them to the network
                    self.add_node(node_1)
                    self.add_node(node_2)
                    self.add_edge(node_1, node_2)
35
      def add_node(self, node):
           Adds the specified node to the network.
           :param node: Node-object
40
           if node.identifier not in self.nodes.keys():
               self.nodes[node.identifier] = node
      def add_edge(self, node_1, node_2):
45
           Adds an (undirected) edge between the two specified nodes.
           : param \ node_{-1}: \ Node-object
           :param node_2: Node-object
           : raises: \ KeyError \ if \ either \ node \ is \ not \ in \ the \ network
50
           # raise an error if the nodes are not in the network
           if node_1.identifier not in self.nodes.keys():
               raise KeyError('There_is_no_node_in_the_network_with_identifier:',
                    node_1)
           if node_2.identifier not in self.nodes.keys():
               raise KeyError('There_is_no_node_in_the_network_with_identifier:',
55
                    node_2)
           \# add the (undirected) edge
           self.nodes[node_1.identifier].add_edge(node_2)
           self.nodes[node_2.identifier].add_edge(node_1)
60
      \mathbf{def} \ \mathtt{get\_node} \, (\, \mathtt{self} \, \, , \, \, \, \mathtt{identifier} \, ) :
           : param \quad identifier: \quad node \quad identifier
           :return: Node-object corresponding to the given node identifier, if
               the node is in the network
           :raises: KeyError if there is no node with that identifier in the
              network
           if identifier not in self.nodes.keys():
               raise KeyError('There_is_no_node_in_the_network_with_identifier:',
                    identifier)
           return self.nodes[identifier]
70
      def has_edge(self, node_1, node_2):
           : param \ node\_1: \ Node-object
           : param node_2: Node-object
           :return: True if the two nodes have an (undirected) edge, False
75
               otherwise
```

```
:raises: KeyError if either node is not in the network
              # raise an error if the nodes are not in the network
              if node_1.identifier not in self.nodes.keys():
    raise KeyError('There_is_no_node_in_the_network_with_identifier:',
80
                           node_1)
               if node_2.identifier not in self.nodes.keys():
    raise KeyError('There_is_no_node_in_the_network_with_identifier:',
                           node_2)
              \textbf{return} \ \ \texttt{node\_1} \, . \, \\ \texttt{has\_edge\_to} \, (\, \texttt{node\_2} \, ) \ \ \textbf{and} \ \ \texttt{node\_2} \, . \, \\ \texttt{has\_edge\_to} \, (\, \texttt{node\_1} \, ) \\
85
         def size(self):
               :return: number of nodes in the network
              return len(self.nodes.keys())
90
         def max_degree(self):
               : return: \ highest \ node \ degree \ in \ the \ network \, , \ 0 \ if \ there \ are \ no \ nodes
               in the network
95
              return max([node.degree() for node in self.nodes.values()], default=0)
```

(f) Simulated annealing: Explain why simulated annealing is a worthwhile optimisation principle in practice.

Simulated annealing will approximate the global minimum (unlike Gradient descent, which will find a precise local minima). These methods avoid getting stuck in a local minima by making big random jumps in the beginning and reduce their size as we stabilize the layout. If a node is "stuck" in a force-field in which it does not belong,see figure 13, a random force might help it to get out. If a node is already in a good position, this random force might false a little bit the scheme but won't break the overall structure. As the temperature is decreasing, the random force won't have the ability to displace the node far away of their position. The more iterations, the less amount of temperature.

(g) Applying the layout algorithms

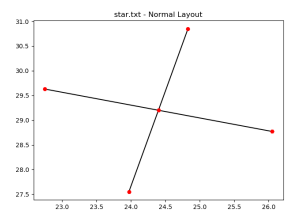


Figure 4:

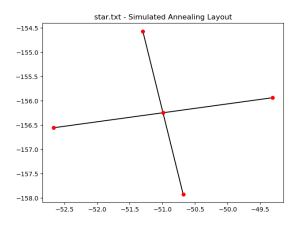


Figure 5:

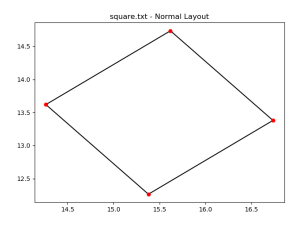


Figure 6:

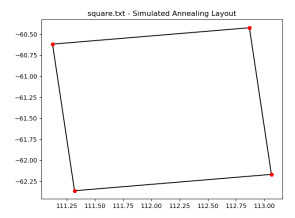


Figure 7:

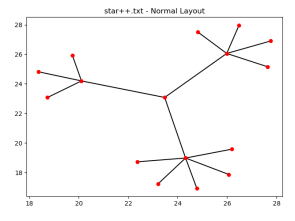


Figure 8:

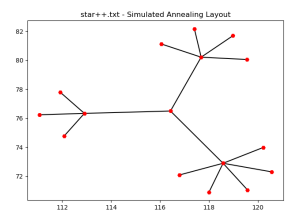


Figure 9:

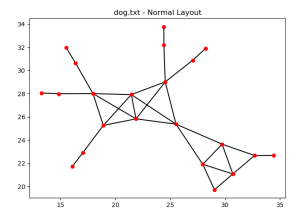


Figure 10:

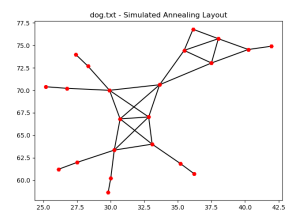


Figure 11:

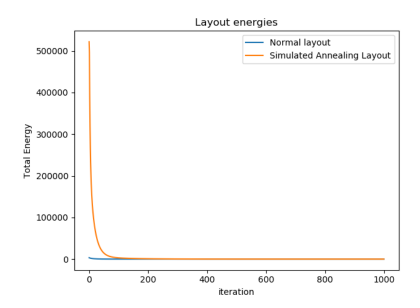


Figure 12: Energies for the star++ graph

We can see in figure 12, the total energy lower quicker with the normal layout than with the Simulated Annealing. Both meet around zero after around 200 iterations.

As said in (f), a node can randomly get stuck somewhere where the forces can't simply make it move and the Simulated Annealing can be useful. In figure 13. After having ran the simulations many times, no such error appeared when using the Simulated Annealing layout.

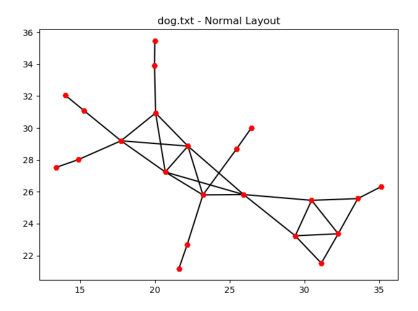


Figure 13: One of the dog's front leg got stuck trying to scratch his back.

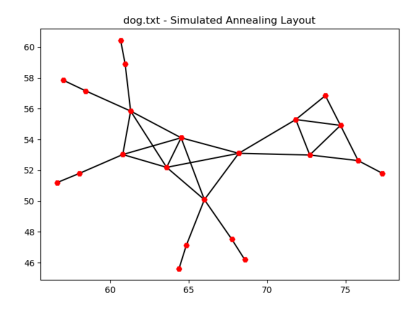


Figure 14: Sometimes, even with annealing layout, the layout is not optimal....

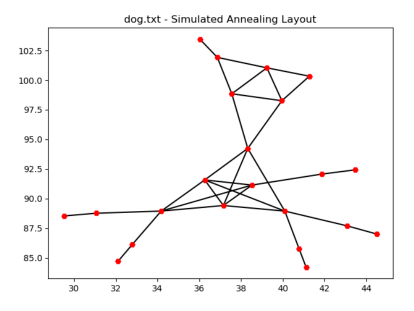


Figure 15: ...or not optimal at all.

Exercise 4.3: Graph Modular Decomposition

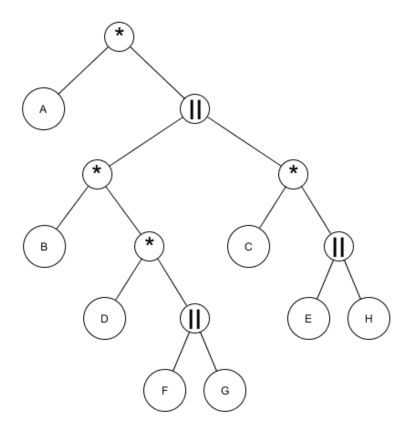


Figure 16: Modular decomposition of the network