Bioinformatics III

Fifth Assignment

Alexander Flohr (2549738) Andrea Kupitz (2550260)

May 25, 2018

Exercise 5.1: Cliques and Network Evolution

(a) Listing 1 shows source code.

```
Listing 1: Listing of source code
```

```
o import random
  from os.path import exists
  {\bf from}\ {\bf AbstractNetwork}\ {\bf import}\ {\bf AbstractNetwork}
  import matplotlib.pyplot as plt
  class CliqueNetwork(AbstractNetwork):
      function that computes the number of cliques of sizes 3, 4 and 5 in a network
      def __createNetwork__(self , filename , dummy):
10
           Create a network from a file
           if exists(filename):
               # "with" closes the file again after reading
15
               with open(filename) as openfile:
                   for line in openfile:
                       # get entries of a line as list
                       content = line [0:(len(line)-1)]. split ("\t")
20
                       # and store them
                        if len(content) == 2:
                            n1 = self.getNode(content[0])
                            n2 = self.getNode(content[1])
                            n1.addLinkTo(n2)
                            n2.addLinkTo(n1)
25
           else:
               print(filename, "does_not_exist")
      def findCliques(self):
30
           function that computes the number of cliques of sizes 3, 4 and 5 in a network.
           :return: list of cliques
           for node1 in self.nodes.values():
35
               for node2 in node1.nodelist:
                   candidates = list()
                   candidates.append(node1)
                   \verb|candidates|.append(\verb|node2|)
                   clique = set(self.extendClique(candidates, 3))
40
                   #only consider cliques of size 3-5
                    if clique not in cliques and len(clique) > 2:
                       cliques.append(clique)
          return cliques
```

```
45
       def extendClique(self, candidates, depth):
           try to extend clique to larger cliques
            : param \ candidates: \ smaller \ clique
           :param depth: clique size; maximal 5
50
           :return: largest clique possible
           if depth <= 5:
                for nextnode in candidates [0]. nodelist:
55
                    if nextnode not in candidates:
                        #check if node is connected to all nodes in clique
                        if all(nextnode.hasLinkTo(x) for x in candidates):
                             candidates.append(nextnode)
                            self.extendClique(candidates, depth+1)
                        else:
60
                            return candidates
                return candidates
           else:
                return candidates
       def evolve (self, t):
           function that takes a parameter t representing the number of time steps, as well as a
           step, randomly insert or delete one edge in the network
            : param \ t: \ number \ of \ time \ steps
70
           cliques = list()
           for c in range (3):
                cliques.append(t * [0])
               i in range(t):
75
                clique = cn. findCliques()
                r = random.random()
                nodelid = random.choice(list(self.nodes))
                node1 = self.nodes[node1id]
               \#decide if edge gets removed or new edge added
80
                if r <= 0.5:
                    #try to find two nodes without edge
                    while True:
                        node2id = random.choice(list(self.nodes))
                        node2 = self.nodes[node2id]
85
                        if not node1.hasLinkTo(node2):
                            node1.addLinkTo(node2)
                            node2.addLinkTo(node1)
                            break
                else:
90
                    #search for edge
                    while len(node1.nodelist) == 0:
                        node1id = random.choice(list(self.nodes))
                        node1 = self.nodes[node1id]
                    node2 = random.choice(node1.nodelist)
                    node1.removeLinkTo(node2)
               #plot number of cliques of size 3, 4 and 5 at the beginning and after each time ste
               \# with t = 100
                cliques = self.plot(clique, i, cliques)
           for index , item in enumerate(cliques):
100
                plt.plot(range(len(item)), item, marker='x')
                plt.xlabel('t')
plt.ylabel('amount_of_cliques')
                plt.title('Evolving_Networks')
                plt.legend('345')
105
                plt.tight_layout()
           plt.show()
       def printClique(self, clique):
110
            print cliques to check result
```

```
:param clique: clique to print
            for line in clique:
                s =
115
                for c in line:
                    s += str(c)
                print(s)
       def plot(self, clique, t, cliques):
120
            count amount of cliques of size i at time step t
            :param\ clique:\ cliques\ in\ current\ time\ step
            :param \ t: \ time \ step
            :param cliques: matrix to store count
125
            :return: matrix with counts
           for c in clique:
                cliques[len(c)-3][t] = cliques[len(c)-3][t] + 1
130
       def numLinks (self):
           summ = 0
           for n in self.nodes:
                summ = summ + len(self.getNode(n).nodelist)
135
           return summ / 2
```

- (b) Listing 1 shows source code.
- (c) Listing 1 shows source code.
- (d) The number of cliques stays approximately the same independent of the time step because approximately the same amount of edges are added which are removed as shown in figure 1, table 4 and table 4. The amount of cliques of size 3 increases rather than the one of larger cliques because a less edges added may result in a small clique.

clique size	number of cliques before evolution	number of cliques after evolution
3	294	300
4	33	33
5	3	3

Table 1: Number of cliques of size 3, 4 and 5 at the beginning and after letting it evolve for 100 time steps

clique size	number of cliques before evolution	number of cliques after evolution
3	294	327
4	33	27
5	3	3

Table 2: Number of cliques of size 3, 4 and 5 at the beginning and after letting it evolve for 1000 time steps

(e) The goal of randomising networks this way is to create random permutations of a network. In this way the behaviour of similar networks may be studied and the networks quality may be rated.

Listing 2 shows source code.

Listing 2: Listing of source code

 $\begin{array}{ll} \textbf{0} & \textbf{from} & AbstractNetwork & \textbf{import} & AbstractNetwork \\ \textbf{import} & random \end{array}$

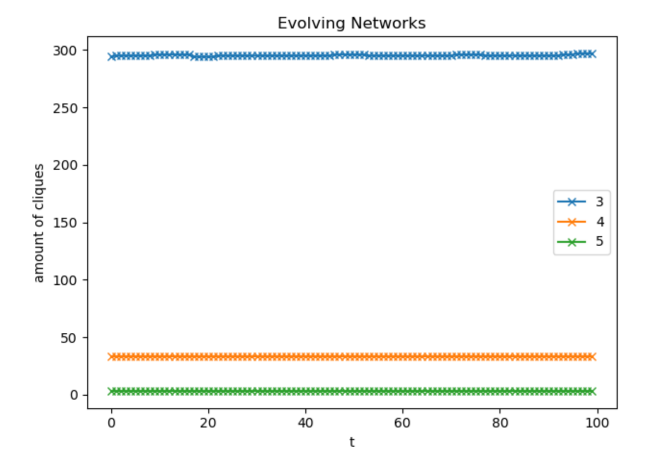


Figure 1: Number of cliques of size 3, 4 and 5 at the beginning and after each time step as a function of time with t=100

from CliqueNetwork import CliqueNetwork

```
class RandomizedNetwork(AbstractNetwork):
       function that takes a network with m edges and returns a randomised version of that network
       def __createNetwork__(self , network , m):
 10
            for 2m iterations, randomly select two edges e1 = (n1; n2) and e2 = (n3; n4) from the
            such\ that\ the\ start\ and\ end\ nodes\ are\ swapped
            :param\ network:\ network
            :param m: number of edges
 15
            self.nodes = network.nodes
            for i in range (2*m):
                node1 = self.chooseNode()
                node2 = random.choice(node1.nodelist)
 20
                node3 = self.chooseNode()
                node4 = random.choice(node3.nodelist)
                node1.addLinkTo(node4)
                node3.addLinkTo(node2)
                node1.removeLinkTo(node2)
 25
                node3.removeLinkTo(node4)
       def chooseNode(self):
            choose random node with edges
 30
            : return: \ random \ node \ with \ at \ least \ one \ edge \\"""
            nodeid = random.choice(list(self.nodes))
            node = self.nodes[nodeid]
            if len(node.nodelist) != 0:
 35
                return node
            else:
                self.chooseNode()
   cn = RandomizedNetwork(CliqueNetwork("test.tsv", "dummy"), 10)
(f) Assuming a p-value of 0.05, none of the clique sizes were significantly enriched, see table 6.
   Listing 3 shows source code.
                            Listing 3: Listing of source code
 o import random
   from CliqueNetwork import CliqueNetwork
   class MotifNetworks:
 5
        class that takes a parameter n and a network and computes if cliques of size 3, 4 and 5 are
        enriched in that network
       network = CliqueNetwork("", "")
       clique = list()
 10
       def __init__(self , network , n):
            self.network = network
            #compute number of cliques in original network
            self.clique = self.network.findCliques()
 15
            #perform n switches
            cliques = self.__createNetwork__(n)
            c_i = dict()
```

#initilize dict
for i in range(3, 6):

 $c_i[i] = 0$

20

```
#count cliques of size i
for i in self.clique:
                c_i[len(i)] = c_i.get(len(i), 0) + 1
           \#\ count\ cliques\ of\ size\ of\ at\ least\ i
25
            for i in range (4, 2, -1):
                    c_i [i] += c_i [i+1]
           \#compute p_{-i}
            extremes = list()
            for key, item in c_i.items():
30
                extremes.append((key, self.count(cliques, key, item)))
            print(extremes)
       def __createNetwork__(self , n):
35
            for 2m iterations, randomly select two edges e1 = (n1; n2) and e2 = (n3; n4) from the
            such that the start and end nodes are swapped
            :param n: number of swaps
40
            cliques = list()
            for i in range(n):
                node1 = self.chooseNode()
                node2 = random.choice(node1.nodelist)
45
                node3 = self.chooseNode()
                node4 = random.choice(node3.nodelist)
                node1.addLinkTo(node4)
                node3.addLinkTo(node2)
                node1.removeLinkTo(node2)
50
                node3.removeLinkTo(node4)
                clique = self.network.findCliques()
                cliques.append(clique)
           return cliques
       \mathbf{def} count(self, cliques, i, c_i):
55
            count \ cliques \ of \ size \ of \ at \ least \ i
            : param \>\>\> cliques: \>\>\> randomized \>\>\>\> cliques
            :param\ i:\ clique\ size
            :param c_i: number of cliques with clique size of at least i in original network
:return: number of networks with at least as many cliques of size of at least i over no
60
                     networks
            extremes = 0
            for network in cliques:
65
                count = 0
                for clique in network:
                     if len(clique) >= i:
                         count += 1
                if count >= c_i:
70
                     extremes += 1
            extremes /= float(len(cliques))
            return extremes
       def chooseNode(self):
            choose\ random\ node\ with\ edges
            :return: random node with at least one edge
            nodeid = random.choice(list(self.network.nodes))
            node = self.network.nodes[nodeid]
            if len(node.nodelist) != 0:
                return node
            else:
                self.chooseNode()
85
  cn = CliqueNetwork("rat_network.tsv", "dummy")
```

MotifNetworks (cn, 100)

clique size i	p_i
3	0.38
4	0.27
5	1.0

Table 3: Number of randomised networks in which the number of cliques is at least as high as in the original network divided by the number of randomised networks for each clique size

Exercise 5.2: Annotations in Protein Protein Interaction Networks

- (a) Implemented in PPIGONetwork.py, see Listing 4
- (b) Required information provided in table 4

Table 4: Reults for exercise 5.2 b

Task	Result
total number of proteins and interactions in the network	17087
total number of unique annotations in the network	275472
total number and percentage of proteins without any annotation	2334~(13.6595%)
smallest number of annotations per protein	0
average number of annotations per protein	7.013
highest number of annotations per protein	184
smallest number of associated proteins per annotatio	1
average number of associated proteins per annotatio	7.013
highest number of associated proteins per annotatio	1519

- (c) In table 5, we see the five most and least annotated GO terms. We see that the most annotated GO terms are all involved in crucial biological processes for cell viability. In this way, we have important transcription factors, which regulate the different expression levels. Additonally, we have one GO term encoding signal transduction, which is also important for cell communication, appoptosis, etc. Further, the GO terms are relatively unspecific, so they do not describe just one specific process, this fewer information content leads to a high number of annotations. Therefore it is no surprise that these GO terms are high annotated. In contrast, the least annotated GO terms describe various, precise biological processes. Therefore they can only be assigned to a specific subset of human proteins. Therefore we see only a few annotations of them. So they are less common but have a higher information content than the most annotated proteins GO terms.
- (d) Not implemented
- (e) Not implemented
- (f) Listing 4 shows source code.

Listing 4: Listing of source code

```
o from CliqueNetwork import CliqueNetwork
from os.path import exists
import sys

class Protein:

Protein class: Storage for protein specific informations
obtained by the network, uniprot and go-term-file
```

Table 5: Reults for exercise 5.2 b

Task	Result			
Most Common				
GO term	Occurrence	Biological Process		
GO:0006351	1519	transcription, DNA-templated		
GO:0045944	1001	positive regulation of transcription by RNA polymerase II		
GO:0007165	980	signal transduction		
GO:0006357	937	regulation of transcription by RNA polymerase II		
GO:0006355	740	regulation of transcription, DNA-templated		
Least Common				

GO term Occurrence Biological Process GO:0000003 1 reproduction GO:0000011 1 vacuole inheritance GO:0000032 1 cell wall mannoprotein biosynthetic process GO:0000053 1 argininosuccinate metabolic process GO:0000097 1 sulfur amino acid biosynthetic process

```
def __init__(self , name):
           # storage for protein name (from PPI)
10
           s\,e\,l\,f\,\,.\,name\,\,=\,\,name
           \# storage for accession number from Go file
           self.accession = ""
           # storage for alternative names (uniprot)
15
           self.alternatives = set ([name])
           # storage for go terms from go file
           self.GOs = set()
       # Getter
       def getName(self):
           return self.name
       def getAccessionNumber(self):
           return self.accession
       def getAlternatives(self):
           return self.alternatives
25
       def getGOTerms(self):
           return self.GOs
       # Setter
       def setName(self , name):
30
            self.name = name
       def setAccessionNumber(self, accession):
           {\tt self.accession} \, = \, {\tt accession}
       def addAlternatives(self, alternatives):
           for a in alternatives:
35
                self.alternatives.add(a)
       def addGOTerm(self, go):
            self.GOs.add(go)
       # terminal visualization
40
       def show(self):
           print "Name: \t\t" + self.name
           print "Accession_Number:\t" + self.accession
print "Alternative_Names:"
           counter = 1
45
           for i in self.alternatives:
    print "\t" + str(counter) + "\t" + i
                counter = counter + 1
           print "GO_Term_Annotations:"
           counter = 1
50
           for i in self.GOs:
```

```
\mathbf{print} "\t" + \mathbf{str} (counter) + "\t" + i
                counter = counter + 1
   class Mapping:
       Mapping class: Stores protein objects and provides
       further functionality
60
       \mathbf{def} __init___(self):
            key: protein name = Protein
            self.mapping = dict()
65
       def exists Accession (self, accession):
            Returns true if the mapping contains a protein with
            the\ given\ accession\ number
70
            for name in self.mapping.keys():
                if self.mapping[name].getAccessionNumber() = accession:
                     return True
            return False
75
       def existsName(self, name):
            Returns true if the mapping contains a protein with
80
            the given name
            if name in self.mapping.keys():
                return True
            else:
                return False
85
       def getMainName(self , variant):
            Return the name of a protein for a given alternative name
90
            if \ \ \text{self.existsName(variant)}:
                return variant
                i in self.mapping.keys():
                if variant in self.mapping[i].getAlternatives():
                     return i
95
            return variant
       def addProtein(self, name):
            Expand mapping by one protein with the given name
100
            self.mapping[name] = Protein(name)
       def addAlternatives(self, name, alternatives):
105
            Add an alternative name to the protein with the given name
            Return without action if the protein name does not exist
            if self.existsName(name):
                self.mapping[name].addAlternatives(alternatives)
110
       \mathbf{def}\ \mathrm{addGOTerm}(\ \mathrm{self}\ ,\ \mathrm{name}\,,\ \mathrm{goterm}\,)\colon
            Add a go term to the protein with the given name.
            Return without action if the protein name does not exist
115
            if self.existsName(name):
                self.mapping[name].addGOTerm(goterm)
```

```
def addAccession(self, name, accession):
120
           Add an accession number to the protein with the given name
           Return without action if the protein name does not exist
           if self.existsName(name):
125
                self.mapping[name].setAccessionNumber(accession)
       def show(self):
            Write proteines into the terminal
130
           counter = 1
           for i in self.mapping.keys():
                print str(counter) + ".\t
                self.mapping[i].show()
135
                counter = counter + 1
       def getKeys(self):
            Return mapping keys (in the end, these are the protein names)
140
           return self.mapping.keys
       # Exercise specific functions
145
       def sumNoAnnotation(self):
           Count proteins without go annotations
150
           summ = 0
           for i in self.mapping.keys():
                 if \ len(self.mapping[i].getGOTerms()) == 0 \colon \\
                   summ += 1
           return summ
155
       def rawDistribution(self):
           Compute the minimal, average and maximal number
            of \ go \ annotations \ per \ gene
160
           values = [sys.maxint, 0, 0]
           for i in self.mapping.keys():
                annotations = len(self.mapping[i].getGOTerms())
                if annotations < values[0]:</pre>
                    values [0] = annotations
165
                if annotations > values[2]:
                    values [2] = annotations
                values [1] = values [1] + annotations
           values[1] = float(values[1])/float(len(self.mapping.keys()))
           return values
170
       def rawReverseDistribution(self):
           Compute the minimal, average and maximal number
            of proteins per go annotations
175
           # initialize dict (key: GO term, value: occurrence)
           # store it as field of the class to allow later access
           self.revsum = dict()
           for i in self.mapping.keys():
180
                terms = self.mapping[i].getGOTerms()
                for j in terms:
                    self.revsum[j] = 0
           # count occurrences
185
           for i in self.mapping.keys():
```

```
terms = self.mapping[i].getGOTerms()
                 for j in terms:
                     self.revsum[j] += + 1
190
            # evaluate minimum, average and maximum
            values = [sys.maxint, 0, 0]
            for i in self.revsum.keys():
                temp = self.revsum[i]
                 if temp < values [0]:
                     values[0] = temp
195
                 if temp > values [2]:
                     values[2] = temp
                 values[1] = values[1] + temp
            values [1] = float (values [1]) / float (len (self.mapping.keys()))
            return values
200
       def extendedNumber(self, length, number):
            Append zeros in front of a string, allows a lexicographical
            sorting of different sized numbers
205
            while len(number) < length:
                number = "0" + number
            return number
210
       def nMostFewestAnnotations(self, n):
            Returns the n most and n fewest annotated GO terms
215
            # create a dict, where the keys are a combination of occurrences
            # and GO terms (as string - allows easy sorting)
            sorter = dict()
            for i in self.revsum.keys():
                key = self.extendedNumber(10, str(self.revsum[i])) + i
220
                 sorter [key] = (i, self.revsum[i])
            # sort the keys of the sorter dict in lexicographically increaing
            # order, at the same time the most occurring terms come to the end
            helping = []
            for i in sorter.keys():
225
                helping.append(i)
            helping.sort()
            \# extract the n most and fewest common GO annotations
230
            high = []
            low = []
            for i in helping [0:n]:
                low.append(sorter[i])
                i in helping [(len(helping)-n):len(helping)]:
                high.append(sorter[i])
235
            return [high, low]
   class PPIGONetwork:
240
       Performs part a b c of exercise 5.2
       \mathbf{def} \ \_\mathtt{init}\_\_(\, \mathrm{self} \; , \; \; \mathrm{ppi}\_\mathrm{file} \; , \; \; \mathrm{uniprot}\_\mathrm{file} \; , \; \; \mathrm{go}\_\mathrm{file} \, ) \colon
            \# read in ppi network
            print "1. _Create_Network"
245
            self.ppi = CliqueNetwork(ppi_file, "")
            # create mapping of protein informations
            print "2._Initialize_Mapping"
            self.mapping = Mapping()
            self.initializeMapping()
250
            print "3._Assign_Alternative_Names"
            self.updateMappingNames(uniprot_file)
```

```
print "4._Assign_Go_terms"
           self.assignGOTerms(go_file)
           print "Initialization_completed"
255
       def initializeMapping(self):
            Transfere network proteins into mapping
260
           for n in self.ppi.nodes:
                self.mapping.addProtein\,(n)
       def updateMappingNames(self, uniprot_file):
265
           Read in uniprot file to extend mapping with alternative
           protein names
           if exists(uniprot_file):
                with open (uniprot_file) as openfile:
270
                    for line in openfile:
                        # get entries of a line as list
                        content = line [0:(len(line)-1)].split("\t")
                        if len(content) >= 4:
                            variants = content [4].split("_")
275
                            for temp in variants:
                                self.mapping.addAlternatives(temp, variants)
           else:
                print(filename, "does_not_exist")
280
       def assignGOTerms(self, go_file):
           Extend mapping with GO terms of GO term file
           Add accession number to proteins
285
           if exists(go_file):
               \# "with" closes the file again after reading
                with open(go_file) as openfile:
                    for line in openfile:
                        # get entries of a line as list
290
                        content = line [0:(len(line)-1)].split("\t")

#content = [x \ for \ x \ in \ content \ if \ x \ != ""]

if content [0] = "UniProtKB" and len(content) > 8:
                            if content [8] == "P":
295
                                temp = self.mapping.getMainName(content[2])
                                self.mapping.addGOTerm(temp, content[4])
                                self.mapping.addAccession(temp, content[1])
       def size (self):
           return self.ppi.size()
300
       def overview (self):
           Output information required for subtasks a b c
305
                                         _Overview _-
           print "Total_number_of_proteins:\t\t\t" + str(self.ppi.size())
           no = self.mapping.sumNoAnnotation()
           percent = (float(no)/float(self.ppi.size()))*100.0
310
           print "Number_of_proteines_without_GO_annotation:\t" + str(no) + "_(" + str(percent) +
           distr = self.mapping.rawDistribution()
           print "Smallest_number_of_GO_annotations_per_protein\t" + str(distr[0])
           print "Average_number_of_GO_annotations_per_protein\t" + str(distr[1])
           print "Highest_number_of_GO_annotations_per_protein\t" + str(distr[2])
315
           distr = self.mapping.rawReverseDistribution()
           print "Smallest_number_of_proteins_per_GO_annotations\t" + str(distr[0])
           print "Average_number_of_proteins_per_GO_annotations\t" + str(distr[1])
           print "Highest_number_of_proteins_per_GO_annotations\t" + str(distr[2])
```

Alexander Flohr (2549738) Andrea Kupitz (2550260) Bioinformatics III Fifth Assignment

```
temp = self.mapping.nMostFewestAnnotations(5)

print "5_most_annotated_GO_terms:\t" + str(temp[0])

print "5_fewest_annotated_GO_terms:\t" + str(temp[1])
```