Bioinformatics III Second Assignment

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Exercise 2.1: The Scale-Free network

(a) Implement the algorithm given in the lecture to set up a scale-free network according to the Barabási-Albert model (see Lecture 2, slide 8). Start from the first three connected nodes and add each new node with a given number of links. Connect the new links with increasing preference to nodes that have higher degrees. This ScaleFreeNetwork-class should again use the abstract network class that you wrote in the first assignment. To obtain a much faster implementation and full points, think of a method to map the probabilities to connect to nodes somehow instead of computing them from scratch in each iteration.

Note: We first generated the probability distribution en each iteration and use the function $random.choices(pop,\ prob\)$ to select a node according to its probability (Method 1). After an intense reflection and the understanding of what what said during the tutorial, we tried a second option (Method 2) commented in the listing 1. The benchmark below (figure 2) shows that the first option seems more time-efficient and thus we used this one to execute the program with the requested amount of nodes (100'000).

```
Debug: 1000 nodes and 2 links.... creating ScaleFree network
Network created -> Time elapsed: 0.008204527695973714 minutes
Debug: 10000 nodes and 2 links.... creating ScaleFree network
Network created -> Time elapsed: 0.6214408477147421 minutes
```

Figure 1: Time of execution with method 1, with 1000 and 10'000 nodes

```
Debug: 1000 nodes and 2 links... creating ScaleFree network
Network created. Size: 1000 Total Degree: 3994
Network created -> Time elapsed: 0.03091550668080648 minutes
Debug: 10000 nodes and 2 links... creating ScaleFree network
Network created. Size: 10000 Total Degree: 39994
Network created -> Time elapsed: 3.095263167222341 minutes
```

Figure 2: Time of execution with method 2, with 1000 and 10'000 nodes

Implementation of the missing methods for the ScaleFreeNetwork-class. Listing 1 shows source code of ScaleFreeNetwork.py and listing 2 shows how the class has been tested.

Listing 1: ScaleFreeNetwork.py

```
o \#Bioinformatics 3 : Wiebke Schmitt \& Thibault Schowing
  import random
  from Node import Node
  from AbstractNetwork import AbstractNetwork
  class ScaleFreeNetwork(AbstractNetwork):
      """ Scale-free\ network\ implementation\ of\ AbstractNetwork"""
      def __createNetwork__(self , amount_nodes , amount_links):
10
          Create a network with an amount of n nodes, add m links per iteration
              step
          for n nodes:
               for m links:
                   link node to other nodes
15
          def symetricConnection(node1, node2):
               node1.addLinkTo(node2)
               node2.addLinkTo(node1)
20
          random.seed()
          print("Debug: ", amount_nodes, "_nodes_and", amount_links, "links....
              _creating_ScaleFree_network")
          # Initial m0 nodes connected to each other
25
          m0 = 3
          \# Create Nodes
          for i in range (0, m0):
               self.appendNode(Node(i))
30
          \# Connect Nodes to each other
          for i in range(0,amount_links):
               for j in range (i+1, m0):
                   symetricConnection(self.getNode(i), self.getNode(j%3))
35
          # Method 1
          \# In a first attempt we used the code below.
40
          def genProbList():
              prob_list = []
              sumkj = self.degreeSum()
               for key, node in self.nodes.items():
45
                   ki = node.degree()
                   prob_list.append(ki / sumkj)
              return prob_list
50
          # new nodes id (without the 3 initial nodes)
          for new_node_id in range(3, amount_nodes - 3):
               new_node = Node(new_node_id)
55
               self.appendNode(new_node)
               population = list(range(0, self.size()))
              # Generate probability list of existing nodes
               prob_list = genProbList()
60
               for i in range (amount_links):
```

```
while (True):
                         # choose the neighbour according to its probability
65
                         chosen\_neighbour = random.choices (population , weights =
                             prob_list, k = 1)[0]
                         # if it's a new link and it's not a self-connection
                         if not new_node.hasLinkTo(chosen_neighbour) and not
                             chosen_neighbour == new_node.id:
70
                             symetricConnection(new_node, self.getNode(
                                  chosen_neighbour))
                             break
           # Method 2
75
            # In a second attempt, we used the code below
              # the initial network contains 3x2 links
              network\_degree = 6
           #
              # next node ID
80
           #
              id = 3
           #
              while id < amount\_nodes:
           #
           ..
#
#
                  new\_node = Node(id)
85
                  self.\,appendNode\,(new\_node)
           #
                  remaining\_links = amount\_links
           ..
#
#
                  while \ remaining\_links:
90
           ###
                       rand\_node = random.choice(self.nodes)
                       if(id != rand\_node.id and not rand\_node.hasLinkTo(new\_node))
           #
           ######
                           node\_prob = rand\_node.degree() / network\_degree
                           random\_prob = random.random()
95
                           if(node\_prob > random\_prob):
                               rand_node.addLinkTo(new_node)
                               new\_node. addLinkTo(rand\_node)
100
           ####
                               network\_degree += 2
                                remaining\_links = 1
                  id \neq 1
           #
              print("Network created. Size: ", len(self.nodes), "
                                                                          Total Degree:
                  ", network_degree)
```

(b) Determine the degree distributions for scale-free networks of 10 000 and 100 000 nodes (each with two new links per iteration), respectively, and plot them with double logarithmic axes. A new pre-implemented method in Tools.py will help you with that. What are the differences? Next, compare one of the distributions to the degree distribution of an equally sized random network (play around with the plot-scaling). What are the major differences?

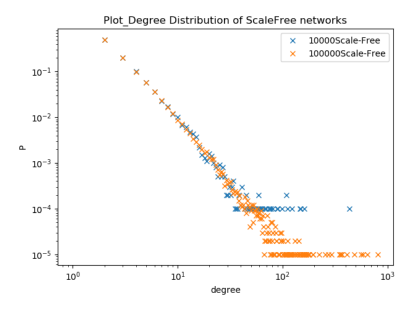


Figure 3: Two scale-free network, one with 10000 nodes and one with 100000 nodes. We can observe that the minimal probability depends highly on the network's size -¿ the bigger the network, the lower the probability can go.

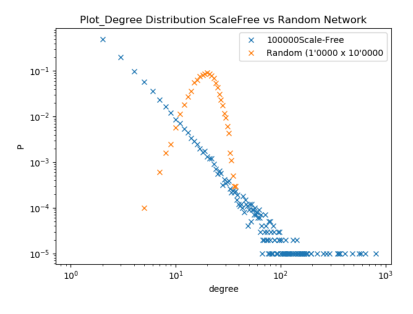


Figure 4: We can observe that the Scale-Free distribution follow a power-law style, as it is almost a straight line when plotted with log-log axes. On the other hand, the random network is more Poisson distributed as seen in Assignment 1.

(c) The degree distribution of a scale-free network follows a power law, which has the form P(k) $k^{-\lambda}$ To simplify the exercise, we assume P(k) $Ck^{-\lambda}$ with C being a fixed normalization constant to obtain a proper distribution. Try to fit this theoretical distribution to the degree distribution of a random network using the Kolmogorov-Smirnov distance.

Listing 2: ScaleFreeTest.py

```
o #!/usr/bin/python
  #Bioinformatics 3: Wiebke Schmitt & Thibault Schowing
  from ScaleFreeNetwork import ScaleFreeNetwork
  from DegreeDistribution import DegreeDistribution
  from RandomNetwork import RandomNetwork
5 import time
  import matplotlib.pyplot as plt
  import Tools as Tools
10 if __name__= "__main__":
      # TASK 2.1 a AND b
      # Number of nodes and link per node
      SMALL = 10000
      BIG = 100000
15
      NB_LINK = 2
      # Create first network
      time1 = time.time()
      sf_net = ScaleFreeNetwork(SMALL, NB_LINK)
20
      time2 = time.time()
      print("Network_created_->_Time_elapsed:_", (time2 - time1)/60, "_minutes")
      # Create second network
      time1 = time.time()
      sf_net2 = ScaleFreeNetwork(BIG, NB_LINK)
      time2 = time.time()
      print("Network_created_->_Time_elapsed:_", (time2 - time1)/60, "_minutes")
      # Create random network
      rand_net = RandomNetwork(10000, 100000)
      # Network's normalized distributions
      sf_degree = DegreeDistribution(sf_net).getNormalizedDistribution()
      sf\_degree2 = DegreeDistribution(sf\_net2).getNormalizedDistribution()
35
      rand_degree = DegreeDistribution(rand_net).getNormalizedDistribution()
      # Plot the degree distributions
      # Small vs Big scale-free network
      legend1 = str(SMALL) + "Scale-Free"
legend2 = str(BIG) + "Scale-Free"
40
      Tools.plotDistributionComparisonLogLog([sf_degree, sf_degree2],[legend1,
           legend2]\,,\ "Plot_Degree\_Distribution\_of\_ScaleFree\_networks"\,)
      \# Big scale-free vs random network
      Tools.plotDistributionComparisonLogLog([sf_degree2, rand_degree], [legend2
45
            "Random_(1'0000_x_10'0000"]," Plot_Degree_Distribution_ScaleFree_vs_
           Random_Network")
      # TASK 2.1 c
      \# Find lambda - 10'000 nodes - 2 links
      sf_net_c = ScaleFreeNetwork(10000,2)
      sf_net_c_degree = DegreeDistribution(sf_net_c).getNormalizedDistribution()
50
      k = len(sf_net_c_degree)
      gamma_distance = []
      \# Foreach gamma, calculate the KS distance steps = [x * 0.1 \text{ for } x \text{ in range}(10, 30)]
55
      for gamma in steps:
           theoretical_dist = Tools.getScaleFreeDistributionHistogram(gamma, k)
           distance \ = \ Tools.simpleKSdist(theoretical\_dist\ , \ sf\_net\_c\_degree)
60
           gamma_distance.append((gamma, distance))
      # Sort the distances-gamma tuples
```

Listing 3: Tools.py

```
o import matplotlib.pyplot as plt
  import math
  from itertools import accumulate
{\tt 5} \ \ \textbf{def} \ \ plot Distribution Comparison (\, histograms \,, \ legend \,, \ title \,) \, :
        Plots a list of histograms with matching list of descriptions as the
           legend
       # determine max. length
10
       max\_length = max(len(x) for x in histograms)
       # extend "shorter" distributions
       for x in histograms:
            x.extend([0.0]*(max_length-len(x)))
15
       # plots histograms
       for h in histograms:
            plt.plot(range(len(h)), h, marker = 'x')
       # remember: never forget labels!
plt.xlabel('degree')
plt.ylabel('P')
20
       # you don't have to do something stuff here
       plt.legend(legend)
25
       plt.title(title)
       plt.tight_layout()
       plt.show()
  def plotDistributionComparisonLogLog(histograms, legend, title):
        Plots a list of histograms with matching list of descriptions as the
        legend,
       fig = plt.figure()
35
       ax = plt.subplot()
       # determine max. length
       \max_{\text{length}} = \max(\text{len}(x) \text{ for } x \text{ in histograms})
       # extend "shorter" distributions
40
       for x in histograms:
            x.extend([0.0]*(max_length-len(x)))
       ax.set_xscale("log")
       ax.set_yscale("log")
45
       # plots histograms
       for h in histograms:
            ax.plot\left(\mathbf{range}(\mathbf{len}\left(h\right)\right),\ h,\ marker\ =\ 'x',\ linestyle='')
50
       # remember: never forget labels!
plt.xlabel('degree')
plt.ylabel('P')
55
       \# you don't have to do something stuff here
       plt.legend(legend)
       plt.title(title)
       plt.tight_layout()
       # Uncomment the line below to display normally
60
       # plt.show()
       # Comment the 2 lines below to display normally
```

```
filename = title + ".png"
          fig.savefig(filename)
 65
    \mathbf{def} \ \ \mathtt{getScaleFreeDistributionHistogram} \left( \mathtt{gamma}, \ k \right) \colon
          Generates a Power law distribution histogram with slope gamma up to degree
 70
          histogram = []
          # NORMALISATION_CONSTANT \
          # Todo here or in ScaleFreeTest.py
 75
          for i in range (1, k+1):
                histogram.append(i**-gamma)
          \#Normalisation
 80
          norm\_hist = [i / sum(histogram) for i in histogram]
          \textbf{return} \quad \texttt{norm\_hist}
    def simpleKSdist(histogram_a, histogram_b):
          Simple \ \ Kolmogorov-Smirnov \ \ distance \ \ implementation
          histograms = [histogram_a, histogram_b]
 90
          max_len = max(len(x) for x in histograms)
          for x in histograms:
                x. extend ([0.0] * (max_len - len(x)))
 95
          \begin{array}{lll} \textbf{for} & i & \textbf{in} & \textbf{range} \left(0\,,\;\; 2\right) \colon \;\; \# \;\; accumulative \;\; distribution \\ & & \text{histograms} \left[\,i\,\,\right] \;=\; \textbf{list} \left(\, accumulate \left(\, \text{histograms} \left[\,i\,\,\right]\,\right)\,\right) \end{array}
100
          ksdist = []
          for i in range(max_len):
                ksdist.append(abs(histogram_a[i] - histogram_b[i]))
          return max(ksdist)
105
```

Use the KS distance to determine a γ (between 1 and 3, 0.1 steps sufficient) that fits best to the degree distribution of a scale-free network with 10 000 nodes and two new links per iteration. Compare the empirical distribution of the network to the theoretical distribution with optimal γ in a double-log. plot. Comment on the quality of your fit, reason why it may fail and how it could be vastly improved.

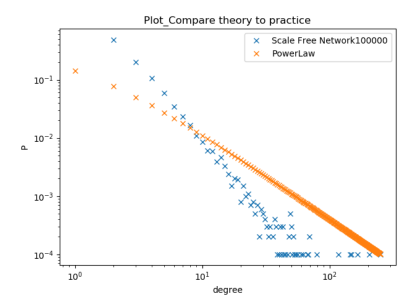


Figure 5: We can affirm that the scale-free distribution follows the power law at least asymptotically. For large values, the probability P(k) for nodes having k connections to other nodes goes as P(k) $k^{-\lambda}$. The only difference is speed at which the probability goes down when increasing the degree, as we can easily notice on the plot. To improve the fit, it would be a good idea to ignore the high degree nodes and fit the power law to the majority (fit the slope).

Exercise 2.2: Classify real-world network examples

(a) File sharing services

The two first listed services, Megaupload and Rapidshare, are more server oriented. The servers host the content, and the client download it. This is more like a scale-free network with a few big central servers around the world (so also like a clustered-network). About directions, each client can upload and download files (media like music and movies are certainly the most famous example). Of course, people uploading files are rarer than people downloading the contend. The traffic, so the directions, are more oriented from the hubs to the leaves/final client.

(b) Social networks

These are undirected networks. Two people are friends or not, but there is no directionality to the relation. A social network can be considered to be a scale-free network, because people with many friends are more likely to make new friends than people who are not as active socially. It can therefore also be considered to be a clustered network, because there tend to be people that are much more connected than others for geographical reasons. Celebrities can also form some kind of hub with many many connected people and form a local small-world.

(c) Broadcasting networks

This is hierarchical networks. Main TV/Radio companies send contents over cables or satellite connexion. For cables, city-relays, neighbourhood-relays or other structure can transmit the information stream from the central node, to the final one (TV or radio). The signal is directed from the broadcasting company to the client, so is the network.

Exercise 2.3: Real interaction networks

(a) Here is the implementation of the BioGRIDReader-class

Listing 4: BioGRIDReader.py

```
o import operator
  from GenericNetwork import GenericNetwork
  from DegreeDistribution import DegreeDistribution
  import Tools as Tools
5 class BioGRIDReader:
        ''Reads BioGRID tab files'''
      def __init__(self , filename):
           Initialization, read in file and build any data structure that makes
              you happy
10
           content_start = False
          # Temporary tab -> contains one line
15
           line\_tab = []
           self.INTERACTOR\_A = [
           self.INTERACTOR_B = [
           self.OFFICIAL_SYMBOL_A =
           self.OFFICIAL_SYMBOL_B = []
20
           self.ALIASES\_FOR\_A = []
           self.ALIASES\_FOR\_B = [
           self.EXPERIMENTALSYSTEM = []
           self.SOURCE = []
           self.PUBMED_ID = []
25
           self.ORGANISM\_A\_ID =
           self.ORGANISM\_B\_ID = []
           with open(filename, "r") as f:
               for line in f:
30
                   if line.startswith("INTERACTOR_A"):
                        content_start = True
                       continue
                   if \verb| content_start: \\
                       # Process data
35
                       line = line.rstrip()
                       line_tab = line.split('\t')
                       \verb|self.INTERACTOR_A.append(line\_tab[0])|\\
                       self.INTERACTOR.B.append(line_tab[1])
40
                       self.OFFICIAL_SYMBOL_A.append(line_tab[2])
                       self.OFFICIAL_SYMBOL_B.append(line_tab[3])
                       self.ALIASES_FOR_A.append(line_tab[4])
                       self.ALIASES_FOR_B.append(line_tab[5])
                       self.EXPERIMENTAL.SYSTEM.append(line_tab[6])
45
                       self.SOURCE.append(line_tab[7])
                       self.PUBMED\_ID.append(line\_tab[8])
                        self.ORGANISM\_A\_ID.append(line\_tab[9])
                       self.ORGANISM_B_ID.append(line_tab[10])
50
          # The file has now been read and all infos are in lists
          # Tuples can store pairwise interactions
55
      def getMostAbundantTaxonIDs(self, n):
           interact = \{\}
           organism_pairs_list = zip(self.ORGANISM_A_ID, self.ORGANISM_B_ID)
60
```

```
for A, B in organism_pairs_list:
               if not A in interact:
                    interact[A] = 1
               else:
65
                    interact[A] += 1
               # If both are the same, the interaction must be counted only once
               if A != B:
                    if not B in interact:
                        interact[B] = 1
70
                    else:
                        interact [B] += 1
           # Sort the dict to retrieve the n first
           \#\ https://stackoverflow.com/questions/613183/how-do-i-sort-a-
               dictionary-by-value
75
           sorted_interact = sorted(interact.items(), key=operator.itemgetter(1))
           nFirst = []
           for i in range (1, n+1):
80
               nFirst.append(sorted_interact[-i])
           return nFirst
85
       def getHumanInteraction(self):
           # Search for human-human interactions
           nb_human_human_interact = 0
           organism_pairs_list = zip(self.ORGANISM_A_ID, self.ORGANISM_B_ID)
           for A, B in organism_pairs_list:
90
               if A == B == "9606":
                    nb_human_human_interact += 1
           print("HUMAN_INTERACTIONS\n")
           print("\nNumber_of_Human_Human_interactions_(human_id_=_9606):_",
95
               nb_human_human_interact)
           # Now we need the indices of the human - human interactions
           \# The code below extract the indices where ORGANISM A / ORGANISM B are
                human and take the intersection
           \#\ Order\ dict:\ https://stackoverflow.com/questions/16772071/sort-dict-
               by-value-python
100
           # Get Indexes
           indexesA = [i for i, x in enumerate(self.ORGANISM_A.ID) if x == '9606'
           indexesB = [i for i, x in enumerate(self.ORGANISM_B_ID) if x == '9606'
               ]
           \# Get intersection
105
           indexes = list(set(indexesA).intersection(indexesB))
           proteins = [self.OFFICIAL_SYMBOL_A[i] for i in indexes]
           proteins.extend([self.OFFICIAL_SYMBOL_B[i] for i in indexes])
110
           proteins\_count = \{\}
           for prot in proteins:
               if prot not in proteins_count:
                   proteins_count[prot] = 1
115
               else:
                    proteins_count[prot] += 1
           proteins_count = sorted(proteins_count.items(), key=lambda x:x[1])
           # Obtain the n most used proteins
           n = 10
120
           nFirst = []
```

```
for i in range(1, n + 1):
                 nFirst.append(proteins_count[-i])
            print("\nThe_", n, "_proteins_with_the_highest_degree_are:_")
125
            print(nFirst)
        def writeInteractionFile(self, taxon_id, filename):
             organism_pairs_list = zip(self.ORGANISM_A_ID, self.ORGANISM_B_ID)
130
             file = open(filename, "w+")
            # Get Indexes
            indexesA = [i for i, x in enumerate(self.ORGANISM_A_ID) if x ==
            indexesB = [i for i, x in enumerate(self.ORGANISM_B_ID) if x ==
135
                 taxon_id]
            # Get intersection
            indexes = list(set(indexesA).intersection(indexesB))
             for i in indexes:
140
                 \mathbf{file} . \, \mathbf{write} \, (\, \mathbf{self} \, . \mathbf{OFFICIAL.SYMBOL.A} \, \lceil \, \mathbf{i} \, \rceil)
                 \begin{array}{l} \textbf{file}.\,write\,("\backslash t")\\ \textbf{file}.\,write\,(\,self\,.OFFICIAL\_SYMBOL\_B\,[\,i\,]\,) \end{array}
                 file.write("\n")
145
             file.close()
   if __name__= "__main__":
150
        path = "../../../Bioinformatics3\_data/assignment2/BIOGRID-ALL-3.4.159.
            tab.txt"
        bio = BioGRIDReader(path)
        abundantTaxon = bio.getMostAbundantTaxonIDs\left(5\right)
        print("The_most_abundent_TaxonIDs_are_(id, _qty):_", abundantTaxon)
        bio.getHumanInteraction()
155
        # Export human interactions to a file
        EXPORT_FILE_NAME = "humanFile.txt"
       EXPORT_ORGANISM = "9606"
        bio.writeInteractionFile(EXPORT_ORGANISM, EXPORT_FILE_NAME)
160
        # Create GenericNetwork with previously exported file
        gen = GenericNetwork(EXPORT_FILE_NAME)
        print(str(gen))
        print("The_network_has_", gen.degreeSum(), "_links.\n")
165
        # Get distribution
        gen_degree = DegreeDistribution(gen).getNormalizedDistribution()
        # Plot the degree distribution
        Tools.plotDistributionComparisonLogLog([gen_degree], ["Human's_proteins_
170
             interactions"], "Plot_Degree_Distribution_Generic_Network_")
```

(b) The class getMostAbundantTaxonIDs(n) is listed in the listing 4 above.

The most abundent TaxonIDs are (id, qty): [('559292', 704012), ('9606', 414501), ('316407', 184023), ('284812', 72149), ('7227', 67935)]

- 559292: Saccharomyces cerevisiae (Baker's Yeast)
- 9606: Human (Homo sapiens)
- 316407: Escherichia coli
- 284812: Schizosaccharomyces pombe (Fission Yeast)
- 7227: Drosophila melanogaster

The order is not surprising at all. All those organism are the most studied in history so it is not a surprise to find many results concerning them.

(c) How big is the human interaction network and which are the 10 proteins with the highest degree? Take one of them as an example and briefly explain the biology behind the connectivity.

Number of Human-Human interactions (human id = 9606): 386192

The 10 proteins with the highest degree are: [('TP53', 3024), ('TRIM25', 2559), ('APP', 2454), ('EGFR', 2134), ('UBC', 2042), ('NTRK1', 2002), ('MDM2', 1939), ('BRCA1', 1876), ('ELAVL1', 1840), ('HDAC1', 1646)]

The gene/protein P53 is the most present in the data. The protein's full name is "Cellular Tumor Antigen p53". "p53 has many mechanisms of anticancer function and plays a role in apoptosis, genomic stability, and inhibition of angiogenesis." This protein interacts with many cellular processes and thus, has many interactions with many other genes/proteins.

In our case, the human interaction network has 17087 nodes and 772384 links.

(d) Generic network distribution and implementation

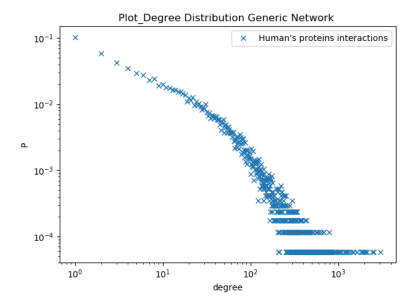


Figure 6: Human interaction degree distribution

 $^{^{1}}$ https://en.wikipedia.org/wiki/P53

We can observe that there is indeed a mix between the scale free straight slope and flat probability as it reaches big degrees and the slight curvature as if it was mixed with a bell-curve distribution. Of course, this is closer to a scale free network. Important molecules/proteins play a role in many reaction and more complex ones might have only a few purposes. The bump on the line is the small part of randomness in the nature that makes it different from maths.

Listing 5: GenericNetwork.py

```
o from AbstractNetwork import AbstractNetwork
  from Node import Node
  # from standard library module
  from itertools import islice
5 import sys
  class GenericNetwork(AbstractNetwork):
      \label{def_def} \mbox{\tt def} \ \mbox{\tt \_init\_\_(self, filename):}
10
           Create a network from a file
           self.nodes = \{\}
15
           # We first need to create all Nodes (unique)
           allEntries = []
           pairs = []
           with open(filename) as f:
20
               # Run through the entire file to make a set of entries
               for line in f:
                    line = line.rstrip()
                    line\_tab = line.split('\t')
25
                    pairs.append(line_tab)
                    allEntries.extend(line_tab)
               allUniqueEntries = set(allEntries)
               for n in allUniqueEntries:
30
                    self.appendNode(Node(n))
               for pair in pairs:
                    self.getNode(pair[0]).addLinkTo(self.getNode(pair[1]))
                    self.getNode(pair[1]).addLinkTo(self.getNode(pair[0]))
```