Network Science Homework 2

PageRank, Communities and Subgraph Patterns

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Work can be located in here

It was use a LLM (Gemini 2.5 Pro Preview 05-06) for help to write the questions and answers in a correct format for the Jupyter Notebook

Link Analysis and PageRank

QUESTION 1.

Draw a graph with at least 6 nodes in which one node has a very **high value of PageRank**, although the same node has **low closeness and betweenness centrality** (don't forget to point out the node).**

Answer:

Consider the following directed graph with 6 nodes:

The target node is **A**.

QUESTION 2.

The **damping factor** in PageRank (parameter β , in slides) controls how of often we follow one of the links of the current node vs going to an arbitrary node on the network.

(a) What does $\beta = 0$ mean? What would happen to the PageRank values in that case? Why?

Answer:

The equation that governs if the algorithm follows the outgoing licks is

$$r_i = \sum_{i \to j} \beta \frac{r_i}{d_i} + (1 - \beta) \frac{1}{n}$$

where $\beta \in [0,1]$

In this equation the damping factor β represents the probability of following the outgoing edges.

If $\beta = 0$ then the algorithm will always follow to a random page

(b) What does $\beta = 1$ mean? Can you explain a possible problem with using that value?

Answer:

As explain in the previous question (a) β represents the probability of following the ou going edges, if $\beta = 1$ that means that the algorithm always follow the outgoing edges of the current page.

the problems of this implementation are:

- Dead ends: what happens if there is no more outgoing edges on the current page, if the $\beta=1$ then the algorithm stops without traversing the entire graph.
- Spider traps: the problem of all out-link are within a group, eventually the Spider trap absorbs all importance.

QUESTION 3.

Implement a program (in any programming language) for manually computing the (normalized) PageRank values of a small network using **power iterations** (the "flow" mode). Attach the program to your homework submission with a very short description on how it works.

Answer:

```
import networkx as nx
import numpy as np
import sys
import os
import random
import json
from networkx.readwrite import json_graph

parent_dir = os.path.abspath(os.path.join(os.getcwd(), '..'))
sys.path.append(parent_dir)
from homeWork1_NS.src.load_save_network import load_network_advanced,
save_network_nx, save_graph_network

random.seed(42)
```

This Page Rank was created to use as input a networkx.DiGraph graph

we also create a graph to test results

```
mode)
    Args:
        graph (nx.DiGraph): The input directed graph from NetworkX.
        beta (float): The damping factor (probability of following a
link).
        iterations (int): The maximum number of iterations to perform.
        tolerance (float): The tolerance for convergence. Iteration
stops if the L1 norm
                           of the difference between PageRank vectors
in consecutive
                           iterations is less than this.
        print iteration (bool): If True, prints the PageRank values at
each iteration.
    Returns:
        dict: A dictionary with node IDs as keys and their normalized
PageRank values as values.
    this comment was generated by Gemini 2.5 Pro Preview 05-06
    #Power iterations. (the "flow" mode)
    nodes = list(graph.nodes())
    N = len(nodes)
    if N == 0:
        return {}
    if beta < 0 or beta > 1:
        raise ValueError("Damping factor beta must be in the range [0,
1].")
    if iterations \leftarrow 0:
        raise ValueError("Number of iterations must be a positive
integer.")
    if tolerance <= 0:
        raise ValueError("Tolerance must be a positive number.")
    if not isinstance(graph, nx.DiGraph):
        raise TypeError("Input graph must be a directed graph
(nx.DiGraph).")
    # First iteration
    final iteration = 0
    # create a page rank list pr
    pr = {node: 1.0 / N for node in nodes}
    # Ensure all nodes are added, even if isolated after edge
```

```
directionality
    out degree = dict(graph.out degree())
    dangling nodes = [node for node in nodes if out degree.get(node,
0) == 01
    if (not print iteration) and print check:
        print(f"Computing PageRank with {N} nodes,
{len(graph.edges())} edges, and damping factor {beta}.")
        print(f"Total Nodes (N): {N}")
        print(f"Initial PageRank (sample): { {k: v for i, (k,v) in
enumerate(pr.items()) if i < 3} }")</pre>
        print(f"Dangling Nodes (count): {len(dangling_nodes)}")
        print(f"Damping Factor (beta): {beta}")
        print("-" * 40)
    for iteration in range(1, iterations + 1):
        new pr = \{\text{node: } 0.0 \text{ for node in nodes}\}
        iteration_dangling_pr_sum = 0.0
        for node in nodes:
            if node in dangling nodes:
                # Distribute PageRank of dangling nodes evenly
                iteration dangling pr sum += pr[node]
            else:
                # Calculate PageRank contribution from neighbors
                if graph.in degree(node) > 0:
                    for neighbor in graph.successors(node):
                        new pr[neighbor] += pr[node] /
out degree[node]
                else:
                    new pr[node] += 0
        for node in nodes:
            dangling contribution_to_node = iteration_dangling_pr_sum
/ N
            new pr[node] = (1 - beta) / N + beta * (new <math>pr[node] +
dangling contribution to node)
        # Check for convergence
        diff = sum(abs(new pr[node] - pr[node]) for node in nodes)
        pr = new pr
        if print_iteration and print_check:
            print(f"Iteration {iteration}:")
            for node id in sorted(pr.keys()):
                print(f" Node {node id}: {pr[node_id]:.8f}")
            print(f"Diff: {diff:.6e}\n")
            print("-" * 40)
```

```
final iteration = iteration
        if diff < tolerance:</pre>
            if print check:
                print(f"Converged after {iteration} iterations (L1
Diff: {diff:.6e}).")
            break
    # Normalize the PageRank values (final step, mostly for precision)
    total pr = sum(pr.values())
    if total pr > 0:
        final pr = {node: value / total pr for node, value in
pr.items()}
    else:
        final pr = {node: 0.0 for node in nodes} # Should not happen
if N > 0
    if (not print iteration) and print check:
        print(f"Final PageRank (the first 3 samples): { {k: v for i,
(k,v) in enumerate(final pr.items()) if i < 3} }")</pre>
        print(f"Final PageRank Sum: {sum(final pr.values()):.6f}")
        print("-" * 40)
    return final pr, final iteration
```

Creating a graph to test using watts strogatz graph and converting to a directed graph

```
# Generate an undirected Watts-Strogatz graph
G = nx.watts strogatz graph(n=30, k=4, p=0.3)
# Converting to a directed
Directed G = nx.DiGraph()
Directed G.add nodes from(G.nodes)
Directed G.add edges from((u, v) if random.random() > 0.5 else (v, u)
for u, v in G.edges) # add edges randomly , from u to v
Directed G.add edges from ((a, b)) if random.random() > 0.5 else (b, a)
for b, a in G.edges) # so that we can have both directions
## start
# Ensure all nodes are added, even if isolated after edge
directionality
all original nodes = set(G.nodes())
current digraph nodes = set(Directed G.nodes())
if all original nodes != current digraph nodes:
    Directed G.add nodes from(all original nodes -
current digraph nodes)
## end , this code was completely generated by Gemini 2.5 Pro Preview
05-06
graph data = json graph.node link data(Directed G)
```

```
# Save the graph using json
with open("data/directed_graph.json", "w") as file:
    json.dump(graph_data, file)

# Save the graph using my custom function for easier reading
save_network_nx(Directed_G, "data/directed_graph_nx.txt")

c:\Users\davib\AppData\Local\Programs\Python\Python312\Lib\site-
packages\networkx\readwrite\json_graph\node_link.py:142:
FutureWarning:
The default value will be `edges="edges" in NetworkX 3.6.

To make this warning go away, explicitly set the edges kwarg, e.g.:

nx.node_link_data(G, edges="links") to preserve current behavior, or
nx.node_link_data(G, edges="edges") for forward compatibility.
warnings.warn(
File data/directed_graph_nx.txt already exists.
```

Loading the Graph

```
with open("data/directed graph.json", "r") as file:
    loaded data = json.load(file)
Directed_G_loaded = json_graph.node_link_graph(loaded_data,
directed=True)
print("Loaded Graph Nodes:", Directed G loaded.nodes())
print("Loaded Graph Edges:", Directed G loaded.edges())
Loaded Graph Nodes: [0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14,
15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29]
Loaded Graph Edges: [(0, 1), (0, 2), (0, 28), (1, 3), (1, 0), (1, 26),
(2, 4), (2, 6), (3, 5), (3, 23), (3, 20), (4, 5), (4, 7), (5, 3), (5, 3)
4), (5, 7), (5, 13), (6, 12), (7, 5), (7, 9), (7, 4), (8, 1), (8, 6),
(8, 9), (8, 10), (9, 8), (9, 10), (9, 11), (10, 8), (10, 11), (10, 11)
12), (11, 13), (11, 9), (11, 12), (12, 11), (12, 13), (12, 6), (13,
5), (13, 14), (13, 16), (13, 11), (14, 13), (14, 16), (14, 18), (15,
13), (15, 17), (15, 25), (16, 7), (16, 14), (17, 18), (18, 14), (18,
24), (18, 29), (19, 21), (19, 20), (20, 3), (20, 16), (20, 19), (20,
22), (21, 23), (21, 22), (22, 21), (22, 23), (22, 24), (23, 1), (23,
2), (23, 3), (23, 17), (23, 25), (23, 21), (23, 22), (24, 18), (24,
22), (24, 25), (24, 26), (25, 24), (25, 27), (25, 15), (26, 1), (26,
24), (26, 25), (26, 27), (27, 14), (27, 28), (27, 29), (28, 0), (28,
29), (29, 0), (29, 27), (29, 28)]
c:\Users\davib\AppData\Local\Programs\Python\Python312\Lib\site-
packages\networkx\readwrite\json graph\node link.py:287:
```

```
FutureWarning:
The default value will be changed to `edges="edges" in NetworkX 3.6.
To make this warning go away, explicitly set the edges kwarg, e.g.:
 nx.node link graph(data, edges="links") to preserve current
behavior, or
 nx.node link graph(data, edges="edges") for forward compatibility.
 warnings.warn(
final pr, final iteration = my Page Rank(Directed G loaded, beta=0.85,
iterations=100, tolerance=1.0e-6)
Computing PageRank with 30 nodes, 90 edges, and damping factor 0.85.
Total Nodes (N): 30
Dangling Nodes (count): 0
Damping Factor (beta): 0.85
Converged after 34 iterations (L1 Diff: 8.102533e-07).
Final PageRank (the first 3 samples): {0: 0.03850978495054025, 1:
0.028770181458189277, 2: 0.01975812477323785
Final PageRank Sum: 1.000000
```

test our results are similar to the official implementation

```
n_compare = 5
results = nx.pagerank(Directed_G_loaded, alpha=0.85, max_iter=100,
tol=1.0e-6)
for i in range(n_compare):
    print(f"Node {i}: My PageRank = {final_pr[i]:.8f}, NetworkX
PageRank = {results[i]:.8f}")

Node 0: My PageRank = 0.03850978, NetworkX PageRank = 0.03851562
Node 1: My PageRank = 0.02877018, NetworkX PageRank = 0.02877240
Node 2: My PageRank = 0.01975812, NetworkX PageRank = 0.01976067
Node 3: My PageRank = 0.03375210, NetworkX PageRank = 0.03375269
Node 4: My PageRank = 0.04081663, NetworkX PageRank = 0.04081693
```

QUESTION 4.

Use your program to **compute the PageRank values** of the following network (with β = 0.85).

Show the values of all nodes for each iteration until the computation converges.

ex graph

```
newG = nx.DiGraph()
newG.add nodes from(['A', 'B', 'C', 'D', 'E', 'F', 'G'])
newG.add edges from([
     ('A', 'B'),
    ('B', 'C'), ('B', 'D'), ('B', 'E'), ('B', 'F'), ('C', 'D'), ('D', 'B'),
    ('E',
          'D'),
    ('F',
           'G'),
     ('G', 'F'),
])
print("Nodes in newG:", newG.nodes())
print("Edges in newG:", newG.edges())
Nodes in newG: ['A', 'B', 'C', 'D', 'E', 'F', 'G']
Edges in newG: [('A', 'B'), ('B', 'C'), ('B', 'D'), ('B', 'E'), ('B', 'F'), ('C', 'D'), ('D', 'B'), ('E', 'D'), ('F', 'G'), ('G', 'F')]
final page, final iteration = my Page Rank(newG, beta=0.85,
iterations=100, tolerance=1.0e-7, print iteration=True)
Iteration 1:
  Node A: 0.02142857
  Node B: 0.14285714
  Node C: 0.05178571
  Node D: 0.29464286
  Node E: 0.05178571
  Node F: 0.17321429
  Node G: 0.14285714
Diff: 4.857143e-01
Iteration 2:
  Node A: 0.02142857
  Node B: 0.27187500
  Node C: 0.05178571
  Node D: 0.13982143
  Node E: 0.05178571
  Node F: 0.17321429
  Node G: 0.16866071
Diff: 3.096429e-01
Iteration 3:
  Node A: 0.02142857
  Node B: 0.14027679
  Node C: 0.07920201
  Node D: 0.16723772
  Node E: 0.07920201
  Node F: 0.22256362
```

```
Node G: 0.16866071
Diff: 2.631964e-01
Iteration 4:
 Node A: 0.02142857
 Node B: 0.16358064
 Node C: 0.05123739
 Node D: 0.18588080
 Node E: 0.05123739
 Node F: 0.19459900
 Node G: 0.21060765
Diff: 1.677877e-01
Iteration 5:
 Node A: 0.02142857
 Node B: 0.17942725
 Node C: 0.05618946
 Node D: 0.14329302
 Node E: 0.05618946
 Node F: 0.23520595
 Node G: 0.18683772
Diff: 1.327154e-01
Iteration 6:
 Node A: 0.02142857
 Node B: 0.14322764
 Node C: 0.05955686
 Node D: 0.15507894
 Node E: 0.05955686
 Node F: 0.21836892
 Node G: 0.22135363
Diff: 1.060733e-01
Iteration 7:
 Node A: 0.02142857
 Node B: 0.15324567
 Node C: 0.05186444
 Node D: 0.15311111
 Node E: 0.05186444
 Node F: 0.24001503
 Node G: 0.20704216
Diff: 6.332829e-02
Iteration 8:
 Node A: 0.02142857
```

```
Node B: 0.15157302
 Node C: 0.05399328
 Node D: 0.14216283
 Node E: 0.05399328
 Node F: 0.22997911
 Node G: 0.22544135
Diff: 4.531371e-02
Iteration 9:
 Node A: 0.02142857
 Node B: 0.14226698
 Node C: 0.05363784
 Node D: 0.14542641
 Node E: 0.05363784
 Node F: 0.24526298
 Node G: 0.21691081
Diff: 3.709490e-02
Iteration 10:
 Node A: 0.02142857
 Node B: 0.14504102
 Node C: 0.05166030
 Node D: 0.14284463
 Node E: 0.05166030
 Node F: 0.23603450
 Node G: 0.22990211
Diff: 3.153067e-02
Iteration 11:
 Node A: 0.02142857
 Node B: 0.14284650
 Node C: 0.05224979
 Node D: 0.14007230
 Node E: 0.05224979
 Node F: 0.24766658
 Node G: 0.22205789
Diff: 2.562210e-02
Iteration 12:
 Node A: 0.02142857
 Node B: 0.14049003
 Node C: 0.05178345
 Node D: 0.14060809
 Node E: 0.05178345
 Node F: 0.24053266
 Node G: 0.23194516
```

```
Diff: 2.084612e-02
Iteration 13:
 Node A: 0.02142857
 Node B: 0.14094545
 Node C: 0.05128270
 Node D: 0.13931457
 Node E: 0.05128270
 Node F: 0.24843609
 Node G: 0.22588133
Diff: 1.671770e-02
Iteration 14:
 Node A: 0.02142857
 Node B: 0.13984596
 Node C: 0.05137948
 Node D: 0.13856007
 Node E: 0.05137948
 Node F: 0.24337861
 Node G: 0.23259925
Diff: 1.382294e-02
Iteration 15:
 Node A: 0.02142857
 Node B: 0.13920463
 Node C: 0.05114584
 Node D: 0.13849095
 Node E: 0.05114584
 Node F: 0.24885520
 Node G: 0.22830039
Diff: 1.095317e-02
Iteration 16:
 Node A: 0.02142857
 Node B: 0.13914588
 Node C: 0.05100956
 Node D: 0.13795748
 Node E: 0.05100956
 Node F: 0.24506489
 Node G: 0.23295549
Diff: 9.310196e-03
Iteration 17:
 Node A: 0.02142857
 Node B: 0.13869243
```

```
Node C: 0.05099707
 Node D: 0.13771332
 Node E: 0.05099707
 Node F: 0.24900924
 Node G: 0.22973373
Diff: 7.888696e-03
Iteration 18:
 Node A: 0.02142857
 Node B: 0.13848489
 Node C: 0.05090071
 Node D: 0.13759573
 Node E: 0.05090071
 Node F: 0.24617438
 Node G: 0.23308642
Diff: 6.705392e-03
Iteration 19:
 Node A: 0.02142857
 Node B: 0.13838495
 Node C: 0.05085661
 Node D: 0.13738782
 Node E: 0.05085661
 Node F: 0.24898007
 Node G: 0.23067680
Diff: 5.611379e-03
Iteration 20:
 Node A: 0.02142857
 Node B: 0.13820822
 Node C: 0.05083537
 Node D: 0.13729161
 Node E: 0.05083537
 Node F: 0.24691065
 Node G: 0.23306163
Diff: 4.769672e-03
Iteration 21:
 Node A: 0.02142857
 Node B: 0.13812644
 Node C: 0.05079782
 Node D: 0.13721795
 Node E: 0.05079782
 Node F: 0.24890021
 Node G: 0.23130262
Diff: 3.979113e-03
```

```
Iteration 22:
 Node A: 0.02142857
 Node B: 0.13806383
 Node C: 0.05078044
 Node D: 0.13713673
 Node E: 0.05078044
 Node F: 0.24738767
 Node G: 0.23299375
Diff: 3.382246e-03
Iteration 23:
 Node A: 0.02142857
 Node B: 0.13799479
 Node C: 0.05076714
 Node D: 0.13709388
 Node E: 0.05076714
 Node F: 0.24881182
 Node G: 0.23170809
Diff: 2.848300e-03
Iteration 24:
 Node A: 0.02142857
 Node B: 0.13795837
 Node C: 0.05075246
 Node D: 0.13705659
 Node E: 0.05075246
 Node F: 0.24770434
 Node G: 0.23291862
Diff: 2.421055e-03
Iteration 25:
 Node A: 0.02142857
 Node B: 0.13792668
 Node C: 0.05074473
 Node D: 0.13702392
 Node E: 0.05074473
 Node F: 0.24872555
 Node G: 0.23197726
Diff: 2.042418e-03
Iteration 26:
 Node A: 0.02142857
 Node B: 0.13789890
 Node C: 0.05073799
```

```
Node D: 0.13700402
 Node E: 0.05073799
 Node F: 0.24791866
 Node G: 0.23284529
Diff: 1.736055e-03
Iteration 27:
 Node A: 0.02142857
 Node B: 0.13788199
 Node C: 0.05073209
 Node D: 0.13698667
 Node E: 0.05073209
 Node F: 0.24865058
 Node G: 0.23215944
Diff: 1.463842e-03
Iteration 28:
 Node A: 0.02142857
 Node B: 0.13786724
 Node C: 0.05072849
 Node D: 0.13697304
 Node E: 0.05072849
 Node F: 0.24806401
 Node G: 0.23278157
Diff: 1.244265e-03
Iteration 29:
 Node A: 0.02142857
 Node B: 0.13785566
 Node C: 0.05072536
 Node D: 0.13696380
 Node E: 0.05072536
 Node F: 0.24858969
 Node G: 0.23228298
Diff: 1.051357e-03
Iteration 30:
 Node A: 0.02142857
 Node B: 0.13784780
 Node C: 0.05072290
 Node D: 0.13695601
 Node E: 0.05072290
 Node F: 0.24816344
 Node G: 0.23272981
Diff: 8.936535e-04
```

```
Iteration 31:
 Node A: 0.02142857
 Node B: 0.13784118
 Node C: 0.05072123
 Node D: 0.13695016
 Node E: 0.05072123
 Node F: 0.24854157
 Node G: 0.23236749
Diff: 7.562666e-04
Iteration 32:
 Node A: 0.02142857
 Node B: 0.13783621
 Node C: 0.05071982
 Node D: 0.13694591
 Node E: 0.05071982
 Node F: 0.24823219
 Node G: 0.23268890
Diff: 6.428266e-04
Iteration 33:
 Node A: 0.02142857
 Node B: 0.13783260
 Node C: 0.05071877
 Node D: 0.13694246
 Node E: 0.05071877
 Node F: 0.24850433
 Node G: 0.23242593
Diff: 5.442879e-04
Iteration 34:
 Node A: 0.02142857
 Node B: 0.13782967
 Node C: 0.05071800
 Node D: 0.13693990
 Node E: 0.05071800
 Node F: 0.24828004
 Node G: 0.23265726
Diff: 4.626447e-04
Iteration 35:
 Node A: 0.02142857
 Node B: 0.13782749
 Node C: 0.05071738
 Node D: 0.13693797
```

```
Node E: 0.05071738
 Node F: 0.24847604
 Node G: 0.23246661
Diff: 3.920019e-04
Iteration 36:
 Node A: 0.02142857
 Node B: 0.13782585
 Node C: 0.05071691
 Node D: 0.13693645
 Node E: 0.05071691
 Node F: 0.24831353
 Node G: 0.23263321
Diff: 3.332016e-04
Iteration 37:
 Node A: 0.02142857
 Node B: 0.13782455
 Node C: 0.05071656
 Node D: 0.13693531
 Node E: 0.05071656
 Node F: 0.24845479
 Node G: 0.23249507
Diff: 2.825254e-04
Iteration 38:
 Node A: 0.02142857
 Node B: 0.13782359
 Node C: 0.05071629
 Node D: 0.13693445
 Node E: 0.05071629
 Node F: 0.24833710
 Node G: 0.23261514
Diff: 2.401466e-04
Iteration 39:
 Node A: 0.02142857
 Node B: 0.13782285
 Node C: 0.05071608
 Node D: 0.13693378
 Node E: 0.05071608
 Node F: 0.24843896
 Node G: 0.23251511
Diff: 2.037145e-04
```

```
Iteration 40:
 Node A: 0.02142857
 Node B: 0.13782228
 Node C: 0.05071593
 Node D: 0.13693327
 Node E: 0.05071593
 Node F: 0.24835377
 Node G: 0.23260168
Diff: 1.731573e-04
Iteration 41:
 Node A: 0.02142857
 Node B: 0.13782185
 Node C: 0.05071581
 Node D: 0.13693288
 Node E: 0.05071581
 Node F: 0.24842724
 Node G: 0.23252927
Diff: 1.469408e-04
Iteration 42:
 Node A: 0.02142857
 Node B: 0.13782152
 Node C: 0.05071571
 Node D: 0.13693259
 Node E: 0.05071571
 Node F: 0.24836560
 Node G: 0.23259172
Diff: 1.248996e-04
Iteration 43:
 Node A: 0.02142857
 Node B: 0.13782127
 Node C: 0.05071564
 Node D: 0.13693236
 Node E: 0.05071564
 Node F: 0.24841861
 Node G: 0.23253933
Diff: 1.060247e-04
Iteration 44:
 Node A: 0.02142857
 Node B: 0.13782108
 Node C: 0.05071559
 Node D: 0.13693219
 Node E: 0.05071559
```

```
Node F: 0.24837402
 Node G: 0.23258439
Diff: 9.012099e-05
Iteration 45:
 Node A: 0.02142857
 Node B: 0.13782093
 Node C: 0.05071555
 Node D: 0.13693206
 Node E: 0.05071555
 Node F: 0.24841228
 Node G: 0.23254649
Diff: 7.652152e-05
Iteration 46:
 Node A: 0.02142857
 Node B: 0.13782082
 Node C: 0.05071552
 Node D: 0.13693195
 Node E: 0.05071552
 Node F: 0.24838003
 Node G: 0.23257901
Diff: 6.504329e-05
Iteration 47:
 Node A: 0.02142857
 Node B: 0.13782073
 Node C: 0.05071550
 Node D: 0.13693188
 Node E: 0.05071550
 Node F: 0.24840765
 Node G: 0.23255160
Diff: 5.523908e-05
Iteration 48:
 Node A: 0.02142857
 Node B: 0.13782067
 Node C: 0.05071548
 Node D: 0.13693182
 Node E: 0.05071548
 Node F: 0.24838434
 Node G: 0.23257508
Diff: 4.695322e-05
Iteration 49:
```

```
Node A: 0.02142857
 Node B: 0.13782062
 Node C: 0.05071546
 Node D: 0.13693177
 Node E: 0.05071546
 Node F: 0.24840428
 Node G: 0.23255526
Diff: 3.988245e-05
Iteration 50:
 Node A: 0.02142857
 Node B: 0.13782058
 Node C: 0.05071545
 Node D: 0.13693174
 Node E: 0.05071545
 Node F: 0.24838742
 Node G: 0.23257221
Diff: 3.390008e-05
Iteration 51:
 Node A: 0.02142857
 Node B: 0.13782055
 Node C: 0.05071544
 Node D: 0.13693171
 Node E: 0.05071544
 Node F: 0.24840182
 Node G: 0.23255788
Diff: 2.879892e-05
Iteration 52:
 Node A: 0.02142857
 Node B: 0.13782053
 Node C: 0.05071544
 Node D: 0.13693169
 Node E: 0.05071544
 Node F: 0.24838964
 Node G: 0.23257012
Diff: 2.447908e-05
Iteration 53:
 Node A: 0.02142857
 Node B: 0.13782051
 Node C: 0.05071543
 Node D: 0.13693168
 Node E: 0.05071543
 Node F: 0.24840004
```

```
Node G: 0.23255976
Diff: 2.079780e-05
Iteration 54:
 Node A: 0.02142857
 Node B: 0.13782050
 Node C: 0.05071543
 Node D: 0.13693167
 Node E: 0.05071543
 Node F: 0.24839123
 Node G: 0.23256860
Diff: 1.767813e-05
Iteration 55:
 Node A: 0.02142857
 Node B: 0.13782049
 Node C: 0.05071543
 Node D: 0.13693166
 Node E: 0.05071543
 Node F: 0.24839874
 Node G: 0.23256112
Diff: 1.502091e-05
Iteration 56:
 Node A: 0.02142857
 Node B: 0.13782048
 Node C: 0.05071543
 Node D: 0.13693165
 Node E: 0.05071543
 Node F: 0.24839237
 Node G: 0.23256750
Diff: 1.276777e-05
Iteration 57:
 Node A: 0.02142857
 Node B: 0.13782048
 Node C: 0.05071542
 Node D: 0.13693165
 Node E: 0.05071542
 Node F: 0.24839780
 Node G: 0.23256209
Diff: 1.084941e-05
Iteration 58:
 Node A: 0.02142857
```

```
Node B: 0.13782047
 Node C: 0.05071542
 Node D: 0.13693164
 Node E: 0.05071542
 Node F: 0.24839320
 Node G: 0.23256670
Diff: 9.221995e-06
Iteration 59:
 Node A: 0.02142857
 Node B: 0.13782047
 Node C: 0.05071542
 Node D: 0.13693164
 Node E: 0.05071542
 Node F: 0.24839712
 Node G: 0.23256279
Diff: 7.836830e-06
Iteration 60:
 Node A: 0.02142857
 Node B: 0.13782047
 Node C: 0.05071542
 Node D: 0.13693164
 Node E: 0.05071542
 Node F: 0.24839379
 Node G: 0.23256612
Diff: 6.661305e-06
Iteration 61:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693164
 Node E: 0.05071542
 Node F: 0.24839662
 Node G: 0.23256329
Diff: 5.661021e-06
Iteration 62:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839422
 Node G: 0.23256570
```

```
Diff: 4.811868e-06
Iteration 63:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839627
 Node G: 0.23256366
Diff: 4.089454e-06
Iteration 64:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839453
 Node G: 0.23256540
Diff: 3.476036e-06
Iteration 65:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839601
 Node G: 0.23256392
Diff: 2.954261e-06
Iteration 66:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839475
 Node G: 0.23256518
Diff: 2.511122e-06
Iteration 67:
 Node A: 0.02142857
 Node B: 0.13782046
```

```
Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839582
 Node G: 0.23256411
Diff: 2.134238e-06
Iteration 68:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839491
 Node G: 0.23256502
Diff: 1.814102e-06
Iteration 69:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839568
 Node G: 0.23256425
Diff: 1.541862e-06
Iteration 70:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839503
 Node G: 0.23256490
Diff: 1.310582e-06
Iteration 71:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839559
 Node G: 0.23256435
Diff: 1.113922e-06
```

```
Iteration 72:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839511
 Node G: 0.23256482
Diff: 9.468336e-07
Iteration 73:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839552
 Node G: 0.23256442
Diff: 8.047659e-07
Iteration 74:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839517
 Node G: 0.23256476
Diff: 6.840510e-07
Iteration 75:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839546
 Node G: 0.23256447
Diff: 5.814185e-07
Iteration 76:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
```

```
Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839522
 Node G: 0.23256472
Diff: 4.942057e-07
Iteration 77:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839543
 Node G: 0.23256451
Diff: 4.200604e-07
Iteration 78:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839525
 Node G: 0.23256468
Diff: 3.570513e-07
Iteration 79:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839540
 Node G: 0.23256453
Diff: 3.034852e-07
Iteration 80:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839527
 Node G: 0.23256466
Diff: 2.579624e-07
```

```
Iteration 81:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839538
 Node G: 0.23256455
Diff: 2.192631e-07
Iteration 82:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839529
 Node G: 0.23256465
Diff: 1.863737e-07
Iteration 83:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839537
 Node G: 0.23256457
Diff: 1.584147e-07
Iteration 84:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
 Node E: 0.05071542
 Node F: 0.24839530
 Node G: 0.23256463
Diff: 1.346525e-07
Iteration 85:
 Node A: 0.02142857
 Node B: 0.13782046
 Node C: 0.05071542
 Node D: 0.13693163
```

```
Node E: 0.05071542
Node F: 0.24839536
Node G: 0.23256458
Diff: 1.144530e-07

Iteration 86:
Node A: 0.02142857
Node B: 0.13782046
Node C: 0.05071542
Node D: 0.13693163
Node E: 0.05071542
Node F: 0.24839531
Node G: 0.23256462
Diff: 9.728503e-08

Converged after 86 iterations (L1 Diff: 9.728503e-08).
```

QUESTION 5.

Use the program to do computations varying the β parameter from 0.0 to 1.0 in steps of 0.05 and:

```
import matplotlib.pyplot as plt beta_values = np.arange(0.0, 1.05, 0.05) # from 0.0 to 1.0 inclusive, step 0.05
```

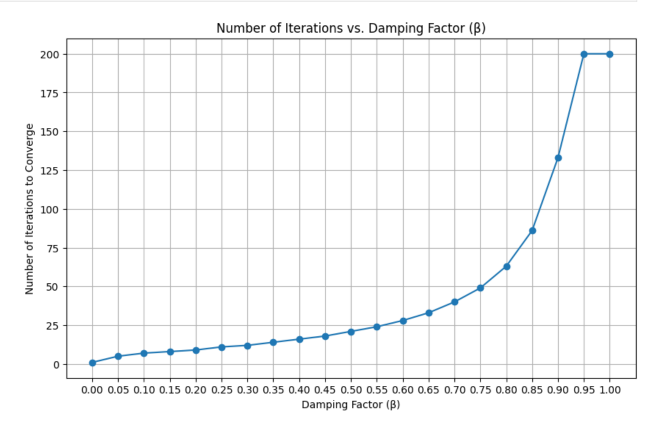
• (a) Show in a plot the number of iterations needed until convergence is reached as you change β. Can you explain what is happening?

```
iteration_counts = []

for values in beta_values:
    # We don't need verbose output for each iteration here, just the count
    _, final_iteration = my_Page_Rank(newG, beta=values, iterations=200, tolerance=1.0e-7, print_check=False)
    iteration_counts.append(final_iteration)
    print(f"Beta = {values:.2f} took {final_iteration} iterations.")

# Plotting
plt.figure(figsize=(10, 6))
plt.plot(beta_values, iteration_counts, marker='o', linestyle='-')
plt.title('Number of Iterations vs. Damping Factor (β)')
plt.xlabel('Damping Factor (β)')
plt.ylabel('Number of Iterations to Converge')
```

```
plt.xticks(beta values)
plt.grid(True)
plt.show()
Beta = 0.00 took 1 iterations.
Beta = 0.05 took 5 iterations.
Beta = 0.10 took 7 iterations.
Beta = 0.15 took 8 iterations.
Beta = 0.20 took 9 iterations.
Beta = 0.25 took 11 iterations.
Beta = 0.30 took 12 iterations.
Beta = 0.35 took 14 iterations.
Beta = 0.40 took 16 iterations.
Beta = 0.45 took 18 iterations.
Beta = 0.50 took 21 iterations.
Beta = 0.55 took 24 iterations.
Beta = 0.60 took 28 iterations.
Beta = 0.65 took 33 iterations.
Beta = 0.70 took 40 iterations.
Beta = 0.75 took 49 iterations.
Beta = 0.80 took 63 iterations.
Beta = 0.85 took 86 iterations.
Beta = 0.90 took 133 iterations.
Beta = 0.95 took 200 iterations.
Beta = 1.00 took 200 iterations.
```



- When β is low (close to 0.0), the random jump component dominates
- When β is high (close to 1.0), the algorithm relies more heavily on the link structure
- β close to 1.0 can be problematic. If there are dangling nodes (pages with no out-links), PageRank "leaks" out of the system unless handled. If there are spider traps, they can absorb all the PageRank. The my_Page_Rank function handles dangling nodes by redistributing their PageRank, which mitigates complete leakage for β=1,
- my Page Rank doesn't accept 0 or 1

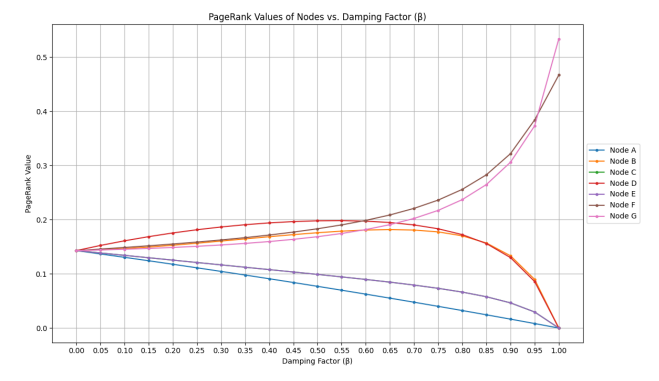
Conclusion:

Since there are spider traps and dangling nodes in the graph ex F and G we can state that the less the random jump component of the equation is prevalent the more interactions are needed to converge

• **(b)** Show in a plot the different **PageRank values of all nodes as your change** β. Can you divide the nodes into different curve behaviors? Can you explain what is happening?

```
pageRank results = {node: [] for node in newG.nodes()}
for values in beta values:
    pr_dict, _ = my_Page_Rank(newG, beta=values, iterations=200,
tolerance=1.0e-7, print check=False)
    for node in newG.nodes():
        pageRank results[node].append(pr dict.get(node, 0.0))
    print(f"PageRanks for beta = {values:.2f} (sample A, B, C, D, E,
F, G): A = \{pr_dict.get('A',0):.3f\}, B = \{pr_dict.get('B',0):.3f\}, 
C={pr_dict.get('C',0):.3f}, D={pr_dict.get('D',0):.3f},
E=\{pr \ dict.get('E',0):.3f\}, F=\{pr \ dict.get('F',0):.3f\},
G={pr dict.get('G',0):.3f}")
# Plotting
plt.figure(figsize=(12, 7))
for node id in sorted(pageRank results.keys()): # sorted for
consistent leaend order
    plt.plot(beta values, pageRank results[node id], marker='.',
linestyle='-', label=f'Node {node id}')
plt.title('PageRank Values of Nodes vs. Damping Factor (β)')
plt.xlabel('Damping Factor (β)')
plt.ylabel('PageRank Value')
plt.xticks(beta_values)
plt.legend(loc='center left', bbox to anchor=(1, 0.5))
plt.grid(True)
plt.tight layout()
plt.show()
```

```
PageRanks for beta = 0.00 (sample A, B, C, D, E, F, G): A=0.143,
B=0.143, C=0.143, D=0.143, E=0.143, F=0.143, G=0.143
PageRanks for beta = 0.05 (sample A, B, C, D, E, F, G): A=0.137,
B=0.144, C=0.138, D=0.152, E=0.138, F=0.146, G=0.144
PageRanks for beta = 0.10 (sample A, B, C, D, E, F, G): A=0.130,
B=0.147, C=0.134, D=0.161, E=0.134, F=0.149, G=0.145
PageRanks for beta = 0.15 (sample A, B, C, D, E, F, G): A=0.124,
B=0.149, C=0.130, D=0.169, E=0.130, F=0.152, G=0.147
PageRanks for beta = 0.20 (sample A, B, C, D, E, F, G): A=0.118,
B=0.153, C=0.125, D=0.175, E=0.125, F=0.155, G=0.149
PageRanks for beta = 0.25 (sample A, B, C, D, E, F, G): A=0.111,
B=0.156, C=0.121, D=0.181, E=0.121, F=0.159, G=0.151
PageRanks for beta = 0.30 (sample A, B, C, D, E, F, G): A=0.104,
B=0.160, C=0.117, D=0.186, E=0.117, F=0.162, G=0.153
PageRanks for beta = 0.35 (sample A, B, C, D, E, F, G): A=0.098,
B=0.164, C=0.112, D=0.191, E=0.112, F=0.167, G=0.156
PageRanks for beta = 0.40 (sample A, B, C, D, E, F, G): A=0.091,
B=0.168, C=0.108, D=0.194, E=0.108, F=0.172, G=0.160
PageRanks for beta = 0.45 (sample A, B, C, D, E, F, G): A=0.084,
B=0.172, C=0.103, D=0.196, E=0.103, F=0.177, G=0.164
PageRanks for beta = 0.50 (sample A, B, C, D, E, F, G): A=0.077,
B=0.176, C=0.099, D=0.198, E=0.099, F=0.183, G=0.168
PageRanks for beta = 0.55 (sample A, B, C, D, E, F, G): A=0.070,
B=0.179, C=0.094, D=0.198, E=0.094, F=0.190, G=0.174
PageRanks for beta = 0.60 (sample A, B, C, D, E, F, G): A=0.062,
B=0.181, C=0.090, D=0.197, E=0.090, F=0.199, G=0.182
PageRanks for beta = 0.65 (sample A, B, C, D, E, F, G): A=0.055,
B=0.182, C=0.085, D=0.195, E=0.085, F=0.209, G=0.191
PageRanks for beta = 0.70 (sample A, B, C, D, E, F, G): A=0.048,
B=0.181, C=0.079, D=0.190, E=0.079, F=0.221, G=0.202
PageRanks for beta = 0.75 (sample A, B, C, D, E, F, G): A=0.040,
B=0.177, C=0.073, D=0.183, E=0.073, F=0.236, G=0.217
PageRanks for beta = 0.80 (sample A, B, C, D, E, F, G): A=0.032,
B=0.170, C=0.066, D=0.172, E=0.066, F=0.256, G=0.237
PageRanks for beta = 0.85 (sample A, B, C, D, E, F, G): A=0.024,
B=0.157, C=0.058, D=0.156, E=0.058, F=0.283, G=0.265
PageRanks for beta = 0.90 (sample A, B, C, D, E, F, G): A=0.016,
B=0.133, C=0.046, D=0.130, E=0.046, F=0.322, G=0.306
PageRanks for beta = 0.95 (sample A, B, C, D, E, F, G): A=0.008,
B=0.090, C=0.030, D=0.086, E=0.030, F=0.384, G=0.373
PageRanks for beta = 1.00 (sample A, B, C, D, E, F, G): A=0.000,
B=0.000, C=0.000, D=0.000, E=0.000, F=0.467, G=0.533
```



- When β = 0.0: All nodes have (or are very close to) the same PageRank value, equal to 1/N (where N is the number of nodes). This is because the random surfer always teleports to a random page, ignoring the link structure.
- As β increases towards 1.0: The link structure becomes more prevalent in the Page Rank calculation.
 - Nodes that are "important" according to the link structure (many incoming links from other important pages, or part of a structure that accumulates rank) will see their PageRank values increase.
 - Nodes that are less "important" (few incoming links, or primarily link out to non-reciprocating structures) will see their PageRank values decrease relative to the more important nodes.

In this graph we can also observe the absorption of "importance" by spider trap (F,G), forcing the other nodes to have a lower page rank the more "importance" the spider trap has.

Community Discovery

QUESTION 6

For this exercise you will be asked to analyze a set of undirected networks depicting the "social networks" (character co-occurrences in a scene) of movies.

You should download the following zip file: movies.zip. It contains:

- Nodes and edges (csv format) for 773 different movies from 1915 to 2012
- a movies.csv file with meta data, indicating which movie name, IMDB id, release year, number of nodes and edges

A readme file describing the original source of the dataset

Start by opening the files on a text editor to see how they internally look like.

```
from networkx.algorithms.community.quality import modularity
import networkx as nx
from task6.src import utils, modularity_utils
import pandas as pd
%load_ext autoreload
%autoreload 2
```

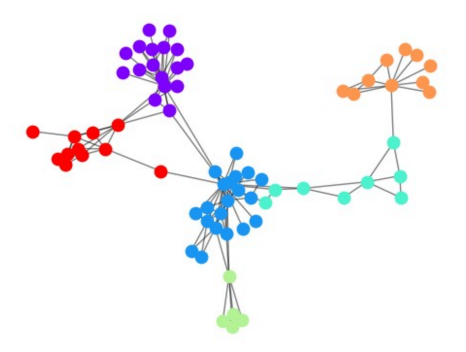
QUESTION (a)

Select any six networks of the dataset and using Gephi, networks or any other platform/library, you should run **Louvain Algorithm** to find the best possible communities and **create a table showing**: id of the dataset, name of the movie, number of nodes and edges, number of communities found and modularity for those communities. Give a brief comment on which networks seem to present **community structure**, and why.

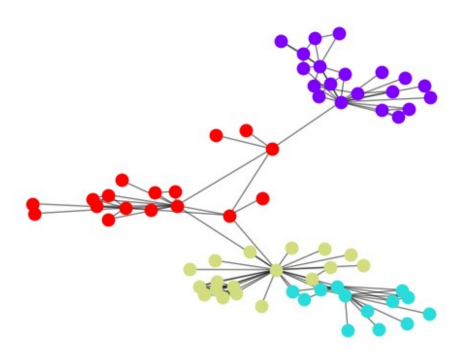
```
movies_data = pd.read_csv("./task6/movies/movies.csv")
table data = []
table columns = ['ID', 'Movie Title', 'Nodes', 'Edges', 'Communities',
'Modularity']
# Create a DataFrame to store the results
for index, row in movies data.iterrows():
   id = row['ID']
   # print(f"Processing {id}...")
   name = row['Title']
   num nodes = row['Characters'] # This is the number of nodes
   num edges = row['Edges']
   nodes_df, edges_df = utils.get_network dataframe(id)
   # print(f'nodes columns: {nodes df.columns}')
   # print(f'edges columns: {edges df.columns}')
   G = utils.create graph from dataframes(nodes df, edges df)
    communities, community_data = utils.compute louvain communities(G,
edges df)
   # print(f"Number of communities:
{community_data['n communities']}")
   # print(f"Modularity: {community data['modularity score']:.4f}")
   num communities = community data['n communities']
   modularity score = community data['modularity score']
   # Create a new row with the data
```

```
row = (id, name, num nodes, num edges, num communities,
modularity score)
    table data.append(row)
    # print(row)
# Create a DataFrame from the list of rows
table df = pd.DataFrame(table data, columns=table columns)
table df.sort values(by='Modularity', ascending=False, inplace=True)
top 6 = table df.head(6)
top 6
                          Movie Title Nodes Edges
                                                     Communities
      ID
Modularity
74
      92
                                Babel
                                          71
                                                154
                                                                6
0.690504
711 837
                              Traffic
                                          68
                                                131
                                                                4
0.668580
                                                                8
447 523
                             Magnolia
                                          82
                                                239
0.662979
342 402
                           Highlander
                                          59
                                                108
                                                                5
0.606953
328 386 He's Just Not That Into You
                                          51
                                                114
                                                                4
0.603339
                                                124
                                                                6
88
     110
                       Batman Returns
                                          51
0.579572
for index, row in top 6.iterrows():
    id = row['ID']
    # print(f"Processing {id}...")
    name = row['Movie Title']
    num nodes = row['Nodes']
    num edges = row['Edges']
    nodes df, edges df = utils.get network dataframe(id)
    # print(f'nodes columns: {nodes df.columns}')
    # print(f'edges columns: {edges df.columns}')
    G = utils.create_graph_from_dataframes(nodes df, edges df)
    communities, community data = utils.compute louvain communities(G,
edges df)
    # print(f"Number of communities:
{community data['n communities']}")
    # print(f"Modularity: {community data['modularity score']:.4f}")
    utils.visualize communities(G, communities, name)
```

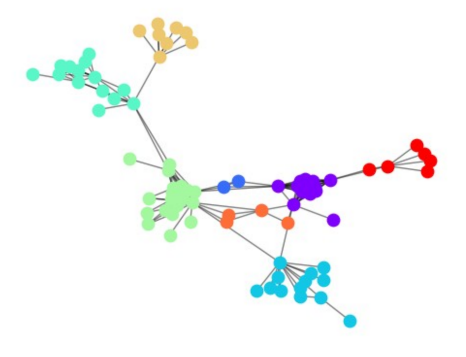
Babel - Louvain Communities



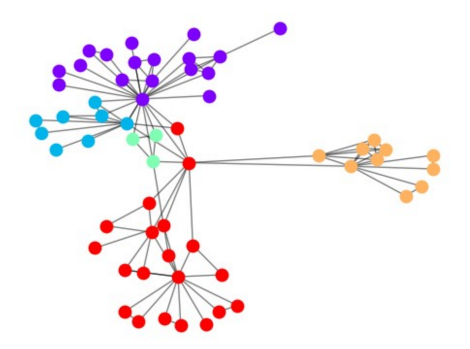
Traffic - Louvain Communities



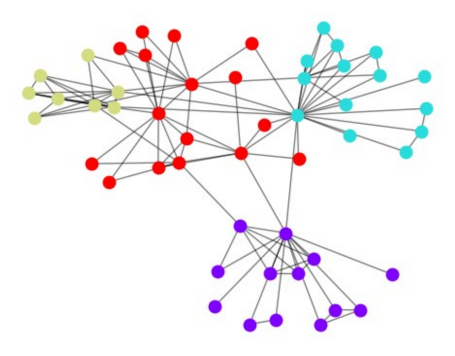
Magnolia - Louvain Communities



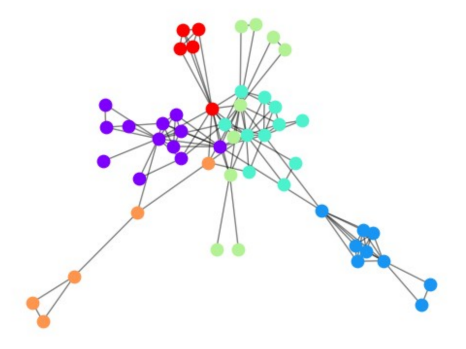
Highlander - Louvain Communities



He's Just Not That Into You - Louvain Communities



Batman Returns - Louvain Communities



R: We approached the problem by applying the Louvain Algorithm to all the networks and sorting them by descending modularity. The top 6 modularity scores are for "Babel", "Traffic",

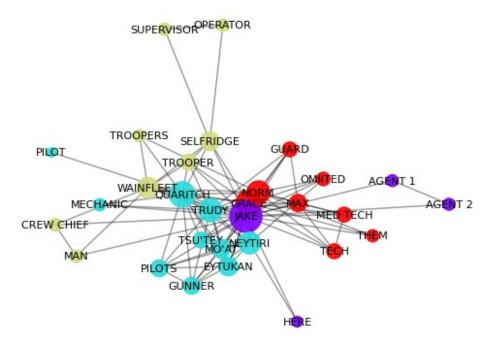
"Magnolia", "Highlander", "He's Just Not That Into You" and "Batman Returns" ordered by higher modularity to lower. When analyzing the networks we can see community structure mainly for the 3 networks, since the nodes are clearly well connected and separated from the other clusters exposing the community structures. For the last 3 as the modularity decreases, harder is to notice the community structures. In these last cases we have more separated nodes without clear clustering or aggregation in general. In some cases there are clusters or defined communities but not at entire network level.

QUESTION (b)

Choose any two of the movies (that are not different chapters of the same universe) and produce visualizations for the networks, labeling the nodes with their character names, using colors to represent communities and the size of the nodes to represent PageRank values. Try to make the picture as aesthetically pleasing as possible, reinforcing the community structure (and explain how you created the layout). Give a brief informal description on the meaning of the communities in the context of the movie (are they what you were expecting? are they meaningful? choose movies that you are familiar with and that you believe should have community structure).

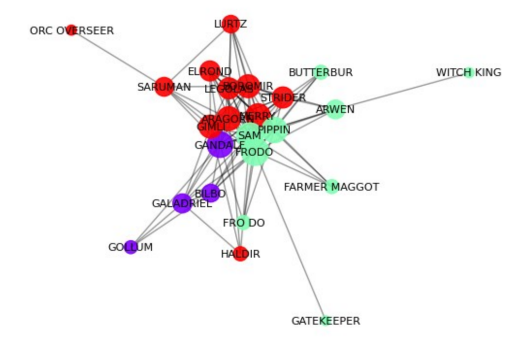
```
nodes_df, edges_df = utils.get_network_dataframe(88)
G = utils.create_graph_from_dataframes(nodes_df, edges_df)
communities, _ = utils.compute_louvain_communities(G, edges_df)
pagerank_scores = utils.compute_pagerank(G, edges_df)
labels = {node: G.nodes[node].get('label') for node in G.nodes()}
# print(labels)
utils.visualize_communities_pgrank(G, communities, title="Avatar",
pagerank_scores=pagerank_scores)
```

Avatar - Louvain Communities



```
nodes_df, edges_df = utils.get_network_dataframe(512)
G = utils.create_graph_from_dataframes(nodes_df, edges_df)
communities, _ = utils.compute_louvain_communities(G, edges_df)
pagerank_scores = utils.compute_pagerank(G, edges_df)
labels = {node: G.nodes[node].get('label') for node in G.nodes()}
# print(labels)
utils.visualize_communities_pgrank(G, communities, title="Lord of Rings", pagerank_scores=pagerank_scores)
```

Lord of Rings - Louvain Communities



R: For this task the network was constructed using the node labels, the pagerank value obtained by NetworkX.pagerank function, the colors for the nodes were determined using community ids generated by the Louvain Algorithm. We selected the films Avatar (88) - node degree 0.27 - and The Lord of Rings (512) - node degree 0.11.

For Avatar we can notice some understandable community aggregations in terms of colors, but they are not well separated. For example we have the scientists (Grace, Norm, Max...) aggregated in the same community. Jake is the central point of the history and have high betweenes in the network. He is classified in a separated community and makes sense because of his dynamic role in the history construction. The Na'vi characters (Tsu'Tey, Neytiri, Mo'At..) a are also aggregated in the same community. We don't have for any of these communities a clear community structure.

For Lord of the Rings network we also don't findclear community structure defined. However, in the communities we can notice the representation aggregations that are reflected in the film history. The hobbities are classified in the same community (Sam, Pippin, Frodo). They are closelly linked to Gandlaf, which belongs to same community of Galladriel. In other hand there is a aggregation of knights (Gimli, Aragorn, Legolas, Boromir) but having in the same community the Saruman. This community reflects some ambiguity and not clear definition.

QUESTION (c)

Implement a program (in any programming language) for manually computing the (normalized) **modularity of a network when given a partition**. Test it on one movie of your choice and the on the partitions you produced on the previous questions (and report if the value seems ok). Attach the program to your homework submission with a very short description on how it works and how I could run it.

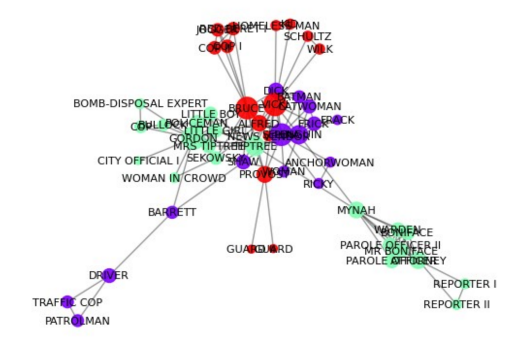
The modularity can be computed as:

$$Modularity = \frac{1}{2m} \left(\sum_{i,j \in V} \left(A_{ij} - \frac{K_i K_j}{2m} \right) \delta \left(C_i, C_j \right) \right)$$

Where A is the adjacency matrix of the graph, Ci is the community to which node i belongs, ki is the degree of node i, m is the total number of edges and V is the set of nodes.

```
nodes df, edges df = utils.get network dataframe(110)
G = utils.create_graph_from_dataframes(nodes df, edges df)
comm 2, progress scores = modularity utils.greedy agglomerative(G)
pagerank scores = utils.compute pagerank(G, edges df)
labels = {node: G.nodes[node].get('label') for node in G.nodes()}
# print(labels)
utils.visualize communities pgrank(G, comm 2, title="Batman Returns",
pagerank scores=pagerank scores)
modularity local implementation =
sum(modularity utils.get intra community modularity(G, comm) for comm
in comm 2)
print(f"Modularity (local implementation):
{modularity_local_implementation:.4f}")
# Example usage of the networkx modularity function
communities, community data = utils.compute louvain communities(G,
edges df)
modularity score = modularity(G, communities)
print(f"Modularity (networkx): {modularity score:.4f}")
```

Batman Returns - Louvain Communities



Modularity (local implementation): 0.4822 Modularity (networkx): 0.4749

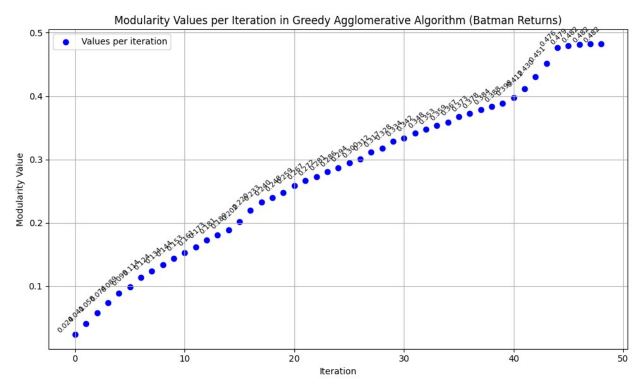
R: We found similar results for modularity but not the same values. At the end it impacted also in the communities composition. In this example we had 0.48 as modularity result whereas the NetworkX implementation found 0.57. To execute the program it is needed to:

- 1. execute the ./task6/src/main.py using the "python main.py" command. The program will ask inputs and output the modularity results of self implemented algorithm
- QUESTION (d)

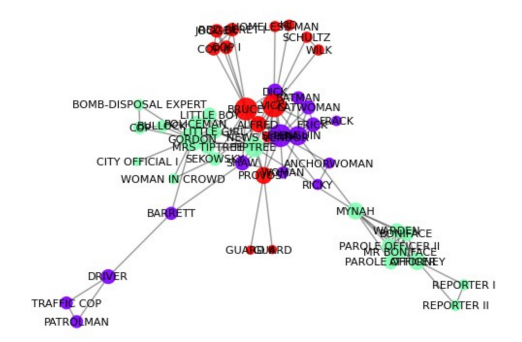
Implement (in any programming language) a **simple greedy agglomerative algorithm**: start with each node being a separated community and then do successive iterations in which you try all possible changes for one node (that is, for each node $i \in V$, try changing its community to all possible communities $j \in C$), and apply the change that produces the best gain in modularity (if there is ties, choose any possible). Attach the program to your homework submission with a very short description on how it works.

Using one of the movies from from the previous questions, make a **plot showing the modularity increase** as you are making more iterations until you reach you a "local maximum", and report the communities you found (as a visualization), comparing them to the communities found previously.

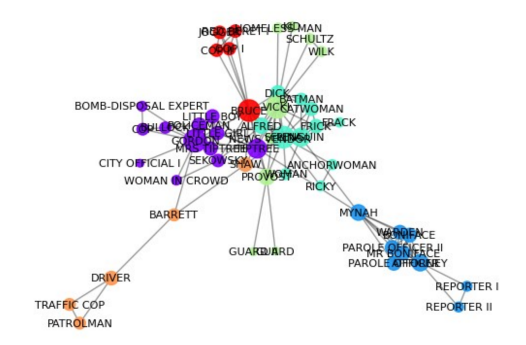
```
# print(progress scores)
# Create iteration index
iterations = list(range(len(progress scores)))
# Plot
plt.figure(figsize=(10, 6))
plt.scatter(iterations, progress_scores, color='blue', label='Values
per iteration')
# Add labels
for i, val in enumerate(progress scores):
    plt.text(i, val, f"{val:.3f}", fontsize=8, ha='right',
va='bottom', rotation=45)
plt.title('Modularity Values per Iteration in Greedy Agglomerative
Algorithm (Batman Returns)')
plt.xlabel('Iteration')
plt.vlabel('Modularity Value')
plt.grid(True)
plt.legend()
plt.tight_layout()
plt.show()
utils.visualize communities pgrank(G, comm 2, title="Batman Returns -
self implemented", pagerank scores=pagerank scores)
utils.visualize communities pgrank(G, communities, title="Batman"
Returns - NetworkX implemented", pagerank scores=pagerank scores)
```



Batman Returns - self implemented - Louvain Communities



Batman Returns - NetworkX implemented - Louvain Communities



R: To execute the program use the command "python task6/src/main.py". It will ask for inputs, for example to the user to indicate the relative path to src folder. Afterwards it requires input to

enter a example ID of movie to calculate the communities using Louvain greedy algorithm. It outputs the score of self-implemented modularity and also NetworkX implemented modularity. The program displays both (self-implemented and NetworkX implementation) networks of communities and the plot of modularity increase in Greedy Algorithm iterations. About the results, as previously mentioned we did not find exactly the same results for modularity, what at the end impacted the communities discovery. But we still possible to generate the communities in several cases.

• QUESTION (e)

Using your previous program as a basis, explain how could you obtain a **larger quantity of communities**? And how could you obtain **less communities**?

Answer:

R: Since in the Louvain algorithm we have an agglomerative approach, we start with the each node of network being one community. As the iterations goes on, the communities are merged and the number of communities in the network reduces. To have larger quantity of communities we should do an early stop. In other hand, to have less communities we should do a late stop, ending with more communities merged (even without improve of modularity).

Network Motifs

QUESTION 7.

For this part of your homework it is highly advisable that you use the gtrieScanner tool. You should download, unzip and compile this version: gtrieScanner_src_01.zip (it is the same as the version online with a newly added "-raw" option to help you on the homework plus some precomputed g-tries)

Your first task is to be able to compile the source code. You will need a C++ compiler and make tools. If you have Linux you can simply use g++ and make available on any common distribution. If you use Windows we suggest you use WSL or Cygwin to have a shell with Linux-like functionality.

Counting subgraphs

QUESTION (a)

Consider the following undirected network:

network_diagram

The frequency (number of occurrences of size 3) of subgraphs of size 3 in this network is:

Subgraph	Frequency
	18
	2

You could obtain these results by running (for instance) one of the following commands:

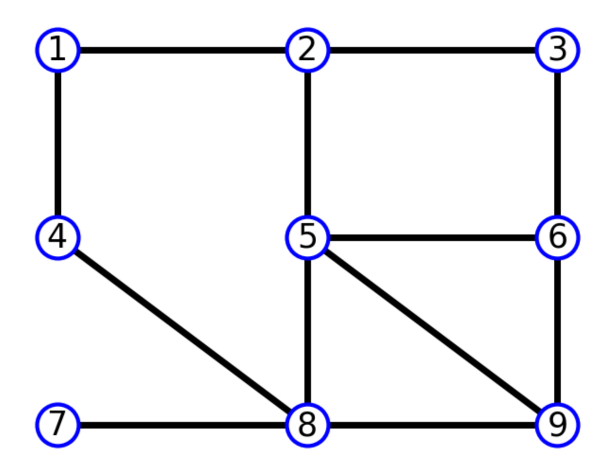
```
./gtrieScanner -s 3 -m esu -g network.txt -f simple
./gtrieScanner -s 3 -m gtrie undir3.gt -g network.txt -f simple
```

supposing that **network.txt** is a text file containing the description of the network as an adjacency list: one line per edge, each line containing two integers separated by a space, the endpoints of the respective edge (the file should have 12 lines, the first of which could be 12, for example).

Your task here is to determine the number of occurrences of all subgraphs of size 4 in this network. You should put in the report a table like the one shown above (the html version of the output is "broken", so you should produce your own images of the subgraphs)

```
import networkx as nx
# Step 1: Create an undirected graph with no parallel edges
G = nx.Graph()
# Add the nodes and edges
G.add nodes from([1,9])
G.add\_edges\_from([(1,2),(1,4),
                  (2,1), (2,5), (2,3),
                  (3,2), (3,6),
                  (4,1), (5,2), (5,6), (5,8), (5,9),
                  (6,3), (6,5), (6,9),
                  (7,8), (8,4), (8,5), (8,9)]
# Position like the image:
pos = {
    7: (0, 0), 8: (1, 0), 9: (2, 0),
    4: (0, 1), 5: (1, 1), 6: (2, 1),
    1: (0, 2), 2: (1, 2), 3: (2, 2)
}
options = {
    "with labels": True,
    "font size": 25,
    "node_size": 1000,
    "node color": "white",
    "edgecolors": "blue",
    "linewidths": 3,
    "width": 5,
}
# Show the nodes and edges
print(G.nodes)
print(G.edges)
```

```
# Draw the graph
nx.draw(G, pos, **options)
[1, 9, 2, 4, 5, 3, 6, 8, 7]
[(1, 2), (1, 4), (9, 5), (9, 6), (9, 8), (2, 5), (2, 3), (4, 8), (5, 6), (5, 8), (3, 6), (8, 7)]
```



```
# Write to file in gtrieScanner-compatible format
with open("network.txt", "w") as f:
    for u, v in G.edges():
        f.write(f"{u} {v}\n")

import matplotlib.pyplot as plt
import numpy as np
import os

adj_matrix_1 = np.array([
    [0, 1, 1, 0],
    [1, 0, 0, 1],
    [1, 0, 0, 0],
    [0, 1, 0, 0]
```

```
]) \# freq = 21
adj_matrix_2 = np.array([
    [0, 1, 1, 1],
    [1, 0, 1, 0],
    [1, 1, 0, 0],
    [1, 0, 0, 0]
]) # freq = 5
adj_matrix_3 = np.array([
    [0, 1, 1, 1],
    [1, 0, 0, 0],
    [1, 0, 0, 0],
    [1, 0, 0, 0]
]) # freq = 4
adj_matrix_4 = np.array([
    [0, 1, 1, 0],
    [1, 0, 0, 1],
    [1, 0, 0, 1],
   [0, 1, 1, 0]
]) # freq = 1
adj matrix 5 = np.array([
    [0, 1, 1, 1],
    [1, 0, 1, 1],
    [1, 1, 0, 0],
    [1, 1, 0, 0]
]) # freq = 1
pos1 = {
    2: (0, 1), 3: (1, 1),
    0: (0, 2), 1: (1, 2),
}
options2 = {
    "with_labels": False,
    "node_size": 150,
    "node color": "black",
    "edgecolors": "black",
    "width": 3,
}
adj_matrices = [adj_matrix_1, adj_matrix_2, adj_matrix_3,
adj_matrix_4, adj_matrix_5]
# Create output directory if it doesn't exist
output dir = "subgraph images"
os.makedirs(output dir, exist ok=True)
```

```
# Draw all graphs
for i, mat in enumerate(adj_matrices, 1):
    G = nx.from_numpy_array(mat)
    plt.figure(figsize=(1.5,1.5))
    plt.title(f"Subgraph {i}", fontsize=8)
    nx.draw(G, pos1, **options2)
    plt.axis("off")  # Hide axis
    plt.tight_layout()
    filepath = os.path.join(output_dir, f"subgraph{i}.png")
    plt.savefig(filepath, dpi=200)
    plt.close()
```

To generate the subgraphs, we plotted the network and saved its adjacency list to a .txt file containing 12 lines, each representing a pair of connected nodes (as shown in the code above). After generating the file, we ran the following command in the shell to search for subgraphs of size 4 within the network:

```
gtrieScanner -s 4 -m esu -g network.txt -f simple
```

We got 5 subgraphs as shown in the table below:

Subgraphs Table:

Subgraph	Frequency
	21
	5
	4
	1
	1

A bit of math: subgraphs in purely random networks

QUESTION (b)

Imagine you have a G_{np} undirected Erd"os–R'enyi random network. What is its expected number of triangles ()? And what about the expected number of chains ()? Justify your answer.

Note that you can test your theory by generating Erd''os-R' envi networks and counting the subgraphs using gtrieScanner, but your answer should be stated as formulas involving n and p.

- ☐ Expected Subgraph Counts in Erdős–Rényi Graph (G(n, p))
 - Expected number of triangles:

$$E[\text{Triangles}] = \binom{n}{3} \cdot p^3 = \frac{n(n-1)(n-2)}{6} \cdot p^3$$

• Expected number of 3-node chains (Path of length 2):

$$E[\text{Chains}] = {n \choose 3} \cdot 3 p^2 (1-p) = \frac{n(n-1)(n-2)}{2} \cdot p^2 (1-p)$$

For testing the hypothesis we generated 2 networks containing 500 nodes and 1000 nodes and evaluated the error (%). After creating the networks, we used Gtrie to count the subgraphs with 3 nodes (chains and triangles), using the code below (example for the file with 1000 nodes):

```
gtrieScanner -s 3 -m esu -g Expected1000 p0.1.txt -f simple
```

```
from IPython.display import Markdown

def generate_erdos_renyi_and_save(n, p, filename):
    G = nx.erdos_renyi_graph(n, p)
    # Save as edge list (one edge per line, undirected)
    with open(f"{filename}.txt", "w") as f:
        for u, v in G.edges():
            f.write(f"{u+1} {v+1}\n")

def expected_triangles(n, p):
    return (n * (n - 1) * (n - 2) / 6) * (p ** 3)

def expected_chains(n, p):
    return (n * (n - 1) * (n - 2) / 2) * (p ** 2) * (1 - p)
```

Lets try 500 nodes

```
n = 500
# Generate the adjency files (Only load the next cell when
necessary!!!)
#for p in [0.01, 0.05, 0.1]:
    #generate erdos renyi and save(500, p, f"Teste Expected500 p{p}")
# Calculate the expected frequencies
expected triangles 1 = \text{expected triangles}(n, 0.01)
expected chains 1 = \text{expected chains}(n, 0.01)
expected triangles 2 = \text{expected triangles}(n, 0.05)
expected chains 2 = expected chains(n, 0.05)
expected_triangles_3 = expected_triangles(n, 0.1)
expected chains 3 = expected_chains(n, 0.1)
# Real frequencies (calculated by Gtrie)
real1 triangles = 19
real1 chains = 6642
real2 triangles = 2784
```

```
real2 chains = 151989
real3 triangles = 20197
real3 chains = 550481
from IPython.display import Markdown
def percentage error(expected, real):
   return abs(expected - real) / expected * 100 if expected != 0 else
0
Markdown(f"""
### Subgraph Frequencies in Erdős-Rényi \( G(n=500, p) \) Networks
| Subgraph Type | p = 0.01 (Expected) | p = 0.01 (Real) | Error
(%) | p = 0.05 (Expected) | p = 0.05 (Real) | Error (%) | p = 0.1
(Expected) | p = 0.1 (Real) | Error (%) |
{real1_triangles} | {percentage_error(expected_triangles_1,
reall_triangles):.1f}% | {expected_triangles_2:.2f}
{real2_triangles} | {percentage_error(expected_triangles_2,
real2_triangles):.1f}% | {expected triangles 3:.2f}
{real3 triangles} | {percentage error(expected triangles 3,
real3 triangles):.1f}%
| **Chain (Path-3)** | {expected_chains_1:.2f} | {real1_chains}
fpercentage error(expected chains 1, real1 chains):.1f}%
{expected chains 2:.2f} {real2 chains}
{percentage error(expected_chains_2, real2_chains):.1f}% |
{percentage error(expected chains 3, real3 chains):.1f}% |
Le's try 1000 nodes for testing if we can obtain lower error values!
""")
<>:41: SyntaxWarning: invalid escape sequence '\('
<>:41: SyntaxWarning: invalid escape sequence '\('
C:\Users\davib\AppData\Local\Temp\ipykernel 18844\2596487662.py:41:
SyntaxWarning: invalid escape sequence '\('
<IPython.core.display.Markdown object>
```

n = 1000

```
n = 1000
```

```
# Generate the adjency files (Only load the next cell when
necessary!!!)
#for p in [0.01, 0.05, 0.1]:
   #generate erdos renyi and save(1000, p, f"Expected1000 p{p}")
# Calculate the expected frequencies
expected_triangles_1 = expected_triangles(n, 0.01)
expected chains 1 = \text{expected chains}(n, 0.01)
expected triangles 2 = \text{expected triangles}(n, 0.05)
expected chains 2 = expected chains(n, 0.05)
expected triangles 3 = \text{expected triangles}(n, 0.1)
expected chains 3 = \text{expected chains}(n, 0.1)
# Real frequencies (calculated by Gtrie)
real1 triangles = 151
real1 chains = 48076
real2 triangles = 20717
real2 chains = 1184104
real3 triangles = 164933
real3 chains = 4467024
from IPython.display import Markdown
def percentage error(expected, real):
   return abs(expected - real) / expected * 100 if expected != 0 else
0
Markdown(f"""
### Subgraph Frequencies in Erdős-Rényi \( G(n=1000, p) \) Networks
| Subgraph Type | p = 0.01 (Expected) | p = 0.01 (Real) | Error
(%) \mid p = 0.05 \text{ (Expected)} \mid p = 0.05 \text{ (Real)} \mid \text{Error (%)} \mid p = 0.1
(Expected) | p = 0.1 (Real) | Error (%) |
-----|
| **Triangle** | {expected_triangles_1:.2f} |
{real1 triangles} | {percentage error(expected triangles 1,
real1 triangles):.1f}% | {expected triangles 2:.2f}
{real2 triangles} | {percentage error(expected triangles 2,
real2 triangles):.1f}% | {expected triangles 3:.2f} |
{real3_triangles} | {percentage_error(expected_triangles_3,
real3 triangles):.1f}% |
| **Chain (Path-3)** | {expected_chains_1:.2f} | {real1_chains}
{percentage_error(expected_chains_1, real1_chains):.1f}% |
{expected_chains_2:.2f} {real2_chains}
```

Back to empirical findings: uncovering motifs in bacteria

• QUESTION (c)

Your task is now to find some network motifs of the transcriptional regulation directed network of the bacteria Escherichia coli. Start by downloading the network as a weighted adjacency list: ecoli.txt (each line is an edge in the format start node end node weight)

This directed network is ready for being fed to gtrieScanner. For example you could run:

```
./gtrieScanner -s 3 -d -m gtrie dir3.gt -g ecoli.txt
```

This would compute the frequency of all possible 13 types of size 3 subgraphs, and it should show you that the most frequent one is the following, appearing 250 times:

Now, if you add the "-r n" option, it should produce n networks with the same degree sequence and it will you show you how often each subgraph appears on it. For example:

```
./gtrieScanner -s 3 -m gtrie dir3.gt -d -g ecoli.txt -r 500 -raw
```

Check the results and report on what is the more overrepresented subgraph, including its z-score (Z), frequency on the original network (real), average number of occurrences (avgR) and standard deviation (stdevR)) on the randomized networks.

```
The z-score of subgraph i is computed as Z_i = \frac{(real_i - avgR_i)}{stdevR_i} as in (Milo et al. 2004).
```

Notice how the most frequent subgraph is not the most significant one. Check if your very simplistic analysis is consistent with the known literature (Milo et al. 2002) (Shen-Orr et al. 2002), that is, if the motif you found is also reported (what is the name given to this motif?)

```
adj_matrix_1 = np.array([
       [0, 1, 1],
```

```
[0, 0, 1],
    [0, 0, 0]
]) # freq = 130
adj matrix 2 = np.array([
    [0, 0, 1],
    [1, 0, 0],
    [0, 0, 0]
]) # freq = 250
adj matrix 3 = np.array([
    [0, 1, 1],
    [0, 0, 0],
    [0, 0, 0]
1) \# freq = 168
adj_matrix_4 = np.array([
    [0, 0, 0],
    [1, 0, 0],
   [1, 0, 0]
]) # freq = 126
options2 = {
    "with labels": False,
    "node size": 150,
    "node color": "black",
    "edgecolors": "black",
    "width": 3,
}
adj matrices = [adj matrix 1, adj matrix 2, adj matrix 3,
adj_matrix_4]
# Create output directory if it doesn't exist
output dir = "motifs images directed"
os.makedirs(output dir, exist ok=True)
# Draw all graphs as directed
for i, mat in enumerate(adj matrices, 1):
    G = nx.from_numpy_array(mat, create_using=nx.DiGraph) # <-- Make</pre>
it directed
    plt.figure(figsize=(1.5,1.5))
    plt.title(f"Subgraph {i}", fontsize=8)
    nx.draw(G, pos1, arrows=True, **options2) # arrows=True to show
direction
    plt.axis("off")
    plt.tight_layout()
    filepath = os.path.join(output dir, f"motif{i}.png")
    plt.savefig(filepath, dpi=200)
```

```
plt.close()
Markdown(f"""
### □ Motif Analysis of *E. coli* Transcriptional Regulatory Network
#### Summary of gtrieScanner Results
We analyzed the directed network using `gtrieScanner` to identify all
3-node subgraphs (motifs) with the code bellow:
`gtrieScanner-s 3-m gtrie dir3.gt-d-g ecoli.txt-r 500-raw`
After comparing the real network to 500 randomized networks with the
same degree sequence, we obtained the following results:
| Subgraph Type (ID) | Structure
Real Frequency | Random Avg | Std Dev | Z-Score |
_____
   | **ID: 011-001-000** | <img
src="task7/Task7 NS/motifs images directed/motif1.png" width="100"/> |
130 | 12.05 | 3.22 | **36.63** |
| **ID: 001-100-000** | <img
src="task7/Task7_NS/motifs_images_directed/motif2.png" width="100"/> |
250
              | 345.19 | 13.91 | **-6.84** |
| **ID: 011-000-000** | <img
src="task7/Task7_NS/motifs_images_directed/motif3.png" width="100"/> |
              | 278.06 | 7.12 | **-15.46** |
168
| **ID: 000-100-100** | <img
src="task7/Task7_NS/motifs_images_directed/motif4.png" width="100"/> |
              | 237.74 | | 5.99 | | **-18.66** |
126
As supported by the literature, the most statistically significant
motif is the **Feedforward Loop (FFL)** (see picture of subgraph 1):
> *"A transcription factor X regulates a second transcription factor
Y, and both jointly regulate one or more operons Z_1...Z_n. An example
of a feedforward loop is the L-arabinose utilization system."*
- *Milo et al., 2004* """)
<IPython.core.display.Markdown object>
```

Characterizing families of networks using motifs

Start by carefully reading the following paper:

Milo et al. "Superfamilies of evolved and designed networks." Science 303.5663 (2004)

The idea here is to perform a very similar analysis, even using some of the same networks!

QUESTION (d)

Download this set of 8 directed networks: networks.zip (inside the zip there is a README.txt explaining what is each network). Use gtrieScanner to compute motif fingerprints of all networks. You should produce and include in the report the following:

- Plot(s) showing the (normalized) significance profile (SP) of all 13 directed motifs of size 3 for each network. Try to expose the similarity between groups of networks. It should be clear to which subgraph corresponds each data point (ex: see figure 1 of the paper).
- One heat map of 8 × 8 cells showing the correlation between the SPs of all pairs of networks (ex: see figure 2 of the paper).
- A visual description of the main characteristic motifs of each group of networks (that is, you should draw them). Can you give an interpretation on why are they so significant?

You should use at least 100 random networks for each original network and you can opt to ignore subgraphs that occur only once in the original network (attributing a z-score of zero to them).

For normalizing the z-scores use the suggested formula: $SP_i = \frac{Z_i}{\sqrt{Z_i} \cdot Z_i}$

For the heat map you can use any sofware. R and Python have several possible packages, but even Excel or LibreOffice will suffice (use range conditional formatting). You even have some possible online alternatives. If you know about it, you can even use a clustering algorithm to produce a dendrogram showcasing the relationship between the families of networks.

Answer:

For completing this task first we ran the following code for each of the 8 networks:

```
gtrieScanner -s 3 -d -m gtrie dir3.gt -g yeast.txt -r 100 -raw
```

After that we extracted the Z scores of all 13 patterns and created a vector with that. An important key to have in mind here is that the motifs are in the same order in the output of Gtrie algorithm.

Then we normalize all the vectors and plotted them.

```
# First we loaded the Z-scores vectors
z vector circuit1 = np.array([
   9.82, # motif 1
   1.88.
            # motif 2
         # motif 3
   1.60,
   -2.76, # motif 4
           # motif 5
   0.0,
   0.0,
           # motif 6
   0.0,
           # motif 7
   0.0,
           # motif 8
           # motif 9
   0.0,
   0.0,
           # motif 10
            # motif 11
   0.0,
```

```
0.0, # motif 12
    0.0
              # motif 13
])
z vector circuit2 = np.array([
    18.00,
               # motif 1
    1.76,
               # motif 2
    1.65, # motif 3
    -6.40, # motif 4
    0.0, # motif 5

0.0, # motif 6

0.0, # motif 7

0.0, # motif 8

0.0, # motif 9
    0.0, # motif 10
0.0, # motif 11
0.0, # motif 12
0.0 # motif 13
])
z vector ecoli = np.array([
     32.94,
             # motif 1
     -7.83,
                # motif 2
     -16.82,
                # motif 3
     -21.81, # motif 4
    0.0, # motif 5
          # motif 6
# motif 7
# motif 8
    0.0,
    0.0,
    0.0,
    0.0, # motif 9
0.0, # motif 10
0.0, # motif 11
0.0, # motif 12
             # motif 13
    0.0
])
z vector english = np.array([
    34.09,
             # motif 1
    29.78,
             # motif 2
             # motif 3
    28.90,
             # motif 4
    23.65,
             # motif 5
    20.96,
             # motif 6
   -14.55.
             # motif 7
   -18.66,
   -22.14, # motif 8
   -24.23, # motif 9
             # motif 10
   -24.55,
   -31.21, # motif 11
   -33.74, # motif 12
-35.17 # motif 13
```

```
1)
z vector french = np.array([
         26.48,
                             # motif 1
         22.87,
                              # motif 2
                          # motif 3
         19.89,
         15.14, # motif 4
    15.14, # motif 4

4.85, # motif 5

-9.23, # motif 6

-12.98, # motif 7

-19.35, # motif 8

-23.15, # motif 9

-26.22, # motif 10

-27.53, # motif 11

-27.73, # motif 12

-42.95 # motif 13
])
z_vector_highschool = np.array([
         186.95, # motif 1
   186.95, # motif 1
42.97, # motif 2
15.37, # motif 3
14.24, # motif 4
11.75, # motif 5
5.45, # motif 6
3.86, # motif 7
3.05, # motif 8
-5.29, # motif 9
-5.66, # motif 10
-14.80, # motif 11
-16.13, # motif 12
-17.94 # motif 13
1)
z_vector_residence = np.array([
         907.\overline{12}, # motif 1
         164.64,
                                 # motif 2
                            # motif 3
         145.86,
         31.01, # motif 4
        18.22, # motif 5
17.31, # motif 6
16.86, # motif 7
3.14, # motif 8
      -17.91, # motif 9

-25.81, # motif 10

-46.94, # motif 11

-49.09, # motif 12

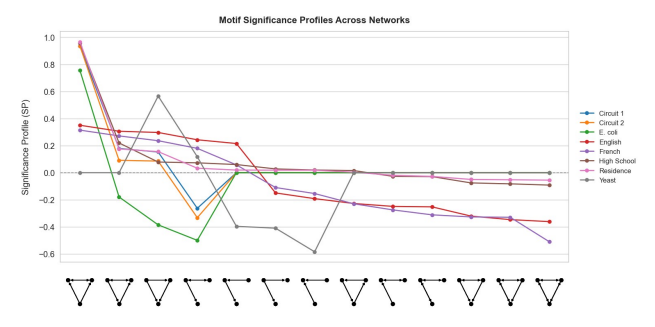
-51.75 # motif 13
])
```

```
z_vector_yeast = np.array([
    0.0,
             # motif 1: 010-100-110 → inf
    0.0,
             # motif 2: 011-100-010 → inf
    13.98.
             # motif 3: 011-001-000
    2.91,
             # motif 4: 011-100-000
   -9.79,
             # motif 5: 001-100-000
             # motif 6: 011-000-000
  -10.13,
  -14.44.
           # motif 7: 000-100-100
             # motif 8: 001-100-010 → nan
   0.0,
   0.0,
             # motif 9: 010-100-100 → nan
   0.0,
             # motif 10: 011-100-100 → nan
   0.0,
            # motif 11: 011-101-000 → nan
            # motif 12: 011-101-100 → nan
   0.0,
   0.0
            # motif 13: 011-101-110 → nan
])
def normalize Z(vector):
    vector = np.array(vector)
    norm = np.linalg.norm(vector) # sqrt(sum of squares)
    return vector / norm if norm != 0 else vector
z vector circuit1 norm = normalize Z(z vector circuit1)
z vector circuit2 norm = normalize Z(z vector circuit2)
z vector ecoli norm = normalize Z(z vector ecoli)
z vector english norm = normalize Z(z vector english)
z vector french norm = normalize Z(z vector french)
z vector highschool norm = normalize Z(z vector highschool)
z_vector_residence_norm = normalize_Z(z_vector_residence)
z vector yeast norm = normalize Z(z vector yeast)
# Define your 13 directed 3-node motif adjacency matrices
adj matrices = [
    np.array([[0, 1, 0], [1, 0, 0], [1, 1, 0]]),
    np.array([[0, 1, 1], [1, 0, 0], [0, 1, 0]]),
    np.array([[0, 1, 1], [0, 0, 1], [0, 0, 0]]),
    np.array([[0, 1, 1], [1, 0, 0], [0, 0, 0]]),
    np.array([[0, 0, 1], [1, 0, 0], [0, 0, 0])),
    np.array([[0, 1, 1], [0, 0, 0], [0, 0, 0])),
    np.array([[0, 0, 0], [1, 0, 0], [1, 0, 0]),
    np.array([[0, 0, 1], [1, 0, 0], [0, 1, 0]]),
    np.array([[0, 1, 0], [1, 0, 0], [1, 0, 0]),
    np.array([[0, 1, 1], [1, 0, 0], [1, 0, 0]]),
    np.array([[0, 1, 1], [1, 0, 1], [0, 0, 0]]),
    np.array([[0, 1, 1], [1, 0, 1], [1, 0, 0]]),
    np.array([[0, 1, 1], [1, 0, 1], [1, 1, 0]])
]
# Drawing options
options2 = {
    "with labels": False,
```

```
"node size": 100,
    "node color": "black",
    "edgecolors": "black",
    "width": 2.5,
}
# Fixed position layout
pos1 = {
    0: (0, 1), 1: (1, 1),
    2: (0.5, 0)
}
# Output folder
output dir = "motifs pngs 3"
os.makedirs(output dir, exist ok=True)
# Draw and export as PNG
for i, mat in enumerate(adj matrices, 1):
    G = nx.from numpy array(mat, create using=nx.DiGraph)
    plt.figure(figsize=(1, 1))
    #plt.title(f"Motif {i}", fontsize=8)
    nx.draw(G, pos=pos1, arrows=True, **options2)
    plt.axis("off")
    #plt.tight layout()
    filepath = os.path.join(output dir, f"motif{i}.png")
    plt.savefig(filepath, format='png', dpi=200)
    plt.close()
import numpy as np
import matplotlib.pyplot as plt
import matplotlib.image as mpimg
from matplotlib.offsetbox import OffsetImage, AnnotationBbox
from matplotlib import gridspec
import seaborn as sns
sns.set(style="whitegrid", context="notebook", font scale=1.3)
# SP matrix (replace with your normalized vectors)
sp matrix = np.array([
    z vector circuit1 norm,
    z vector circuit2 norm,
    z vector ecoli norm,
    z vector english norm,
    z vector french norm,
    z vector highschool norm,
    z vector residence norm,
    z vector yeast norm
])
network labels = [
```

```
"Circuit 1", "Circuit 2", "E. coli", "English", "French", "High School", "Residence", "Yeast"
1
motif image paths = [f"motifs pngs 3/motif{i+1}.png" for i in
range(13)]
# Create figure with two subplots: top for SP, bottom for motif images
fig = plt.figure(figsize=(16, 9))
gs = gridspec.GridSpec(2, 1, height_ratios=[5, 1], hspace=0.05)
# --- Top subplot: SP plot ---
ax0 = plt.subplot(qs[0])
palette = sns.color palette("tab10", n colors=len(sp matrix))
for i, sp in enumerate(sp matrix):
    ax0.plot(range(13), sp, marker='o', label=network_labels[i],
             linewidth=2.0, markersize=6, color=palette[i], zorder=3)
ax0.axhline(0, color='gray', linestyle='--', linewidth=1.2, zorder=1)
ax0.set xlim(-0.5, 12.5)
ax0.set xticks([]) # No ticks because we'll use images
ax0.set_ylabel("Significance Profile (SP)", labelpad=15)
ax0.set_title("Motif Significance Profiles Across Networks",
fontsize=16, weight='bold', pad=15)
ax0.legend(loc='center left', bbox to anchor=(1.01, 0.5),
frameon=False, fontsize=12)
# --- Bottom subplot: Motif images ---
ax1 = plt.subplot(qs[1])
ax1.axis("off") # Hide axis completely
for x, img path in enumerate(motif image paths):
    try:
        img = mpimg.imread(img path)
        imagebox = OffsetImage(img, zoom=0.25)
        ab = AnnotationBbox(imagebox, (x, 0.5), frameon=False,
box_alignment=(0.5, 0.5))
        ax1.add artist(ab)
    except FileNotFoundError:
        print(f"Image not found: {img path}")
ax1.set xlim(-0.5, 12.5)
ax1.set ylim(0, 1)
plt.tight layout()
plt.show()
```

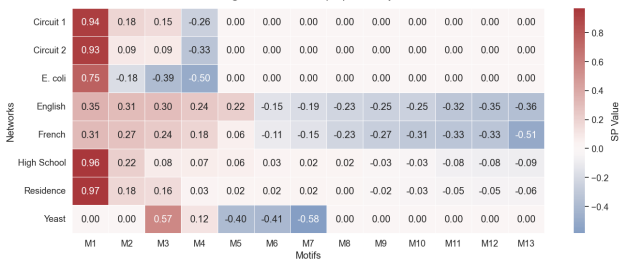
C:\Users\davib\AppData\Local\Temp\ipykernel_18844\2132356139.py:65:
UserWarning: This figure includes Axes that are not compatible with
tight_layout, so results might be incorrect.
 plt.tight_layout()



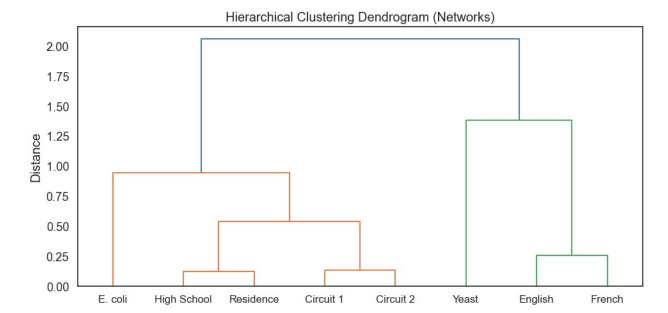
```
# SP matrix: shape (8 networks, 13 motifs)
sp matrix = np.array([
    z vector circuit1 norm,
    z_vector_circuit2_norm,
    z vector ecoli norm,
    z vector english norm,
    z vector french norm,
    z vector highschool norm,
    z vector residence norm,
    z_vector_yeast_norm
1)
network labels = [
    "Circuit 1", "Circuit 2", "E. coli", "English", "French", "High School", "Residence", "Yeast"
1
motif labels = [f''M{i+1}''] for i in range(13)]
plt.figure(figsize=(14, 6))
sns.set(style="white", context="notebook", font scale=1.2)
ax = sns.heatmap(sp matrix, cmap="vlag", center=0, linewidths=0.5,
                  xticklabels=motif labels, yticklabels=network labels,
                  annot=True, fmt=".2f", cbar_kws={'label': 'SP
```

```
Value'})
ax.set_title("Significance Profile (SP) Heatmap", fontsize=16,
weight='bold', pad=15)
plt.xlabel("Motifs")
plt.ylabel("Networks")
plt.tight_layout()
plt.show()
```

Significance Profile (SP) Heatmap



```
from scipy.cluster.hierarchy import dendrogram, linkage
from scipy.spatial.distance import pdist
# Assuming sp matrix and network labels are already defined
# sp matrix shape: (8 \text{ networks}, \overline{13} \text{ motifs})
\# Rows = networks
# Perform hierarchical clustering
linked = linkage(sp matrix, method='ward') # or 'average', 'complete'
# Plot the dendrogram
plt.figure(figsize=(10, 5))
dendrogram(linked,
           labels=network labels,
           orientation='top',
           distance sort='ascending',
           show leaf counts=True)
plt.title("Hierarchical Clustering Dendrogram (Networks)")
plt.vlabel("Distance")
plt.tight_layout()
plt.show()
```



• QUESTION (e)

Your task is to find the "family" of the three "unknown" networks given in unknown.zip You should justify your answer by computing and plotting their motif significance profiles and by adding them to the previous heatmap. Each network will clearly belong to one of the groups discovered on the previous question.

```
z_vector_netA = np.array([
    43.93, # motif 1
   1.99, # MOCI.
1.50, # motif 3
    -15.18,
             # motif 4
             # motif 5
    0.0,
            # motif 6
    0.0,
    0.0,
             # motif 7
    0.0,
             # motif 8
             # motif 9
    0.0,
    0.0,
             # motif 10
    0.0,
             # motif 11
             # motif 12
    0.0,
    0.0
             # motif 13
])
z_vector_netB = np.array([
    0, # motif 1
    6.50,
            # motif 2
    4.77,
           # motif 3
          # motif 4
    2.92,
          # motif 5
    2.49,
    1.68,
              # motif 6
```

```
1.14, # motif 7
     0.95,
                # motif 8
     -1.55, # motif 9

-2.00, # motif 10

-4.03, # motif 11

-4.83, # motif 12

-4.83 # motif 13
])
z_vector_netC = np.array([
     9.23, # motif 1
     8.53, # motif 2
8.05, # motif 3
5.56, # motif 4
    0.41, # motif 5

-3.84, # motif 6

-6.31, # motif 7

-7.00, # motif 8

-7.89, # motif 9

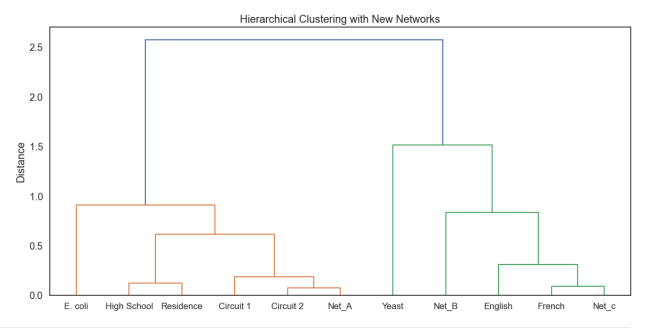
-8.93, # motif 10

-11.05, # motif 11

-11.24, # motif 12

-16.45 # motif 13
1)
z vector netA norm = normalize Z(z vector netA)
z vector netB norm = normalize Z(z vector netB)
z_vector_netC_norm = normalize_Z(z_vector_netC)
new sp matrix = np.array([
     z vector netA norm,
     z vector netB norm,
     z vector netC norm
])
# Stack them vertically
combined_matrix = np.vstack([sp_matrix, new_sp_matrix])
# Perform hierarchical clustering
linked_all = linkage(combined_matrix, method='ward')
# Combined labels (original + new)
combined labels = network labels + ["Net A", "Net B", "Net c"]
# Plot dendrogram
plt.figure(figsize=(12, 6))
dendrogram(linked_all,
              labels=combined labels,
              orientation='top',
              distance sort='ascending',
```

```
show_leaf_counts=True)
plt.title("Hierarchical Clustering with New Networks")
plt.ylabel("Distance")
plt.tight_layout()
plt.show()
```



Significance Profile (SP) Heatmap Circuit 1 0.18 0.15 -0.26 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 - 0.8 Circuit 2 0.09 0.09 -0.33 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 E. coli -0.18 -0.39 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.00 - 0.6 English 0.35 0.31 0.30 0.24 0.22 -0.15 -0.19 -0.23 -0.25 -0.25 -0.32 -0.35 -0.36 - 0.4 French 0.31 0.27 0.24 0.18 0.06 -0.11 -0.15 -0.23 -0.27 -0.31 -0.33 -0.33 High School Residence SP Value 0.22 0.08 0.07 0.06 0.03 0.02 0.02 -0.03 -0.03 -0.08 -0.08 -0.09 Residence 0.18 0.16 0.03 0.02 0.02 0.02 0.00 -0.02 -0.03 -0.05 -0.05 -0.06 - 0.0 0.00 0.00 0.12 -0.40 -0.41 0.00 Yeast 0.00 0.00 0.00 0.00 0.00 - -0.2 0.04 0.03 -0.33 0.00 0.00 0.00 0.00 0.00 0.00 Net_A 0.00 0.00 0.00 Net B 0.00 0.38 0.24 0.20 0.14 0.09 0.08 -0.16 -0.32 -0.39 -0.39 -0.12 - -0.4 0.29 0.27 0.25 0.17 0.01 -0.12 -0.22 -0.35 Net_c -0.20 -0.25 -0.28 -0.35

M7

Motifs

M8

M9

M10

M11

M12

M13

As shown above, the unknown networks belong to different clusters:

M5

M4

M1

M2

МЗ

• Net_A shows similar patterns to *circuit1* and *circuit2*, suggesting it may be a circuit network.

M6

- Net_B shows simmilar patterns to *english or yeast*, indicating it could be either a language or a biological network.
- Net_C shows simmilar patterns to english and french so it could be a language network.