Social Network Analysis using Cultural Algorithms and its Variants

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Social Network Analysis using Cultural Algorithms and its Variants

by

Pooya Moradian Zadeh

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Social Network Analysis using Cultural Algorithms and its Variants

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Declaration of Co-Authorship/Previous Publication

1- Co-Authorship Declaration

I hereby declare that this dissertation incorporates material that is the result of research conducted under the supervision of Dr. Ziad Kobti (my supervisor). This dissertation also incorporates the outcome of a joint research undertaken in collaboration with Mr. Mukund Pandey under the supervision of Dr. Ziad Kobti. The collaboration is covered in Chapters 3 of the thesis. In all cases, the key ideas, primary contributions, experimental designs, data analysis and interpretation, were performed by the author, and the contribution of co-authors was primarily through the proofreading of the published manuscripts. In Chapter 3, Mr. Pandey also contributed in collecting data and explaining the materials.

I am aware of the University of Windsor Senate Policy on Authorship and I certify that I have properly acknowledged the contribution of other researchers to my thesis, and have obtained written permission from each of the co-author(s) to include the above material(s) in my thesis.

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Abstract

Finding relationships between social entities and discovering the underlying structures of networks are fundamental tasks for analyzing social networks. In recent years, various methods have been suggested to study these networks efficiently, however, due to the dynamic and complex nature that these networks have, a lot of open problems still exist in the field. The aim of this research is to propose an integrated computational model to study the structure and behavior of the complex social network.

The focus of this research work is on two major classic problems in the field which are called community detection and link prediction. Moreover, a problem of population adaptation through knowledge migration in real-life social systems has been identified to model and study through the proposed method. To the best of our knowledge, this is the first work in the field which is exploring this concept through this approach.

In this research, a new adaptive knowledge-based evolutionary framework is defined to investigate the structure of social networks by adopting a multi-population cultural algorithm. The core of the model is designed based on a unique community-oriented approach to estimate the existence of a relationship between social entities in the network. In each evolutionary cycle, the normative knowledge is shaped through the extraction of the topological knowledge from the structure of the network. This source of knowledge is utilized for the various network analysis tasks such as estimating the quality of relation between social entities, related studies regarding the link prediction, population adaption, and knowledge formation.

The main contributions of this work can be summarized in introducing a novel method to define, extract and represent different sources of knowledge from a snapshot of a given network to determine the range of the optimal solution, and building a probability matrix to show the quality of relations between pairs of actors in the system. Introducing a new similarity metric, utilizing the prior knowledge in dynamic social network analysis and study the co-evolution of societies in a case of individual migra-
tion are another major contributions of this work. According to the obtained results, utilizing the proposed approach in community detection problem can reduce the search space size by 80%. It also can improve the accuracy of the search process in high dense networks by up to 30% compared with the other well-known methods. Addressing the link prediction problem through the proposed approach also can reach the comparable results with other methods and predict the next state of the system with a notably high accuracy. In addition, the obtained results from the study of population adaption through knowledge migration indicate that population with prior knowledge about an environment can adapt themselves to the new environment faster than the ones who do not have this knowledge if the level of changes between the two environments is less than 25%. Therefore, utilizing this approach in dynamic social network analysis can reduce the search time and space significantly (up to above 90%), if the snapshots of the system are taken when the level of changes in the network structure is within 25%.

In summary, the experimental results indicate that this knowledge-based approach is capable of exploring the evolution and structure of the network with the high level of accuracy while it improves the performance by reducing the search space and processing time.
Dedication

This dissertation is dedicated to my parents, Mohammad and Manijeh, and to my brother, Nima, for their endless love, support, and encouragement.
Acknowledgments

First and foremost, I want to thank my parents for their love and support throughout my life.

I would like to express my deepest appreciation to my supervisor, Dr. Ziad Kobti, for his guidance, advice, and support, throughout this study. Without his guidance and persistent help, this dissertation would not have been possible.

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Chapter 1

Introduction

In recent years, the role of social networks in the evolution of societies has been at the center of attention which is mainly because of the extensive growth of Internet usage and digital connectivity. In general, the social network consists of social actors who are linked together through some kind of relations. Having complex interdependent structures with enormous influences on other systems makes them an attractive and critical research topic for a broad range of scientific fields including sociology, computer, physics, business, medical, and management sciences. Indeed, because of the mutual influence of the network and people, exploring the behavior and structure of these networks have been reviewed by a vast variety of business-oriented, socio-economic, and political approaches. [41, 14, 42, 21, 45]

Consequently, social network analysis (SNA) as an interdisciplinary field has many applications ranging from the study of information propagation and its cascades effects [5, 11, 17, 19, 26] to spread of disease and viruses [13, 10, 6, 37, 18] and from events and disasters detection [47, 57, 40, 38, 25] to prediction of future activities and interactions [48, 23, 4]. Enhancing the marketing and impact maximization techniques [44, 20, 27, 22, 35, 32, 24, 36, 51, 55], identification of high-risk groups and suspicious activities [12, 15, 16, 9, 52, 39, 49], opinion mining and sentiment analysis
2

are other emerging applications of these studies.

Even though various methods have been already proposed to study these networks efficiently, due to the dynamic and complex nature of these networks, a lot of open challenges still exist. In this research work, a novel knowledge-based evolutionary approach is introduced for the investigation of the structure and evolution of complex social networks. Three main research problems in the field have been identified and addressed through this approach. An introduction to this research study is presented in this chapter with the following structure. Basic concepts and characteristics of the social networks are reviewed in the next section. The research problems are defined in section 1.2. The contributions are listed in section 1.3 and the dissertation structure are discussed in section 1.4.

1.1 Social Networks: Basic Concepts

As mentioned before, social networks are social structures made up of a set of actors which are connected to each other through some kind of relations. The common method to represent these networks is mapping them to a graph structure. Therefore, the set of actors forms the set of nodes, and the relations are mapped to a set of links. Consequently, \( G(V, E) \) represents a social network graph where \( V = \{v_1, \ldots, v_n\} \) is a set of nodes and \( E \) is a set of edges that connect the vertices, \( e = (v_i, v_j) \in E \) where \( v_i, v_j \in V \).

Hence, the graph with \( n \) nodes can be described by its adjacency matrix denoted by \( A \) which is an \( n \) by \( n \) matrix where \( A_{ij} \) is 1 if there is a link between \( v_i \) and \( v_j \) and it is 0 if the link does not exist. Furthermore, in a case that the direction of relations is necessary for the analysis a directed graph is used to model the network. The weighted graph also can be utilized in a case that the quality of relationships is not uniform. In other words, edge-weight measures the strength of the connection
between a pair of nodes.

Weights can be assigned to the edges by using various techniques but in principle three main criteria are considered for this issue:

- Similarity between the nodes,
- Distance between the nodes,
- The frequency of relations between two nodes.

In the first case, the level of similarity between a pair of nodes can be considered to calculate the weight of links between them. Different metrics such as the number of shared features, common neighbor nodes or similar attributes can be employed for this process. The second case is focusing on the length of the path between two nodes. Usually, the shorter length leads to have a higher edge-weight. The last one is counting the number of paths between a pair of nodes. Having more paths means stronger connections between the nodes. Combinational methods also can be utilized to make a weighted graph.

On the other hand, in recent years, an emerging concept of multi-layer social network analysis is receiving close review. The main assumption is that the actors are linked to each other through different kind of relations simultaneously and can be members of multiple networks at the same time. In other words, due to the social nature of humans, each actor accepts different social roles (e.g. friend, teacher, manager, mother, child, girl) in the society. Each role has its own particular type of relationship and social etiquette. The fact is, the actor must take some of these roles concurrently which requires multi-membership in societies. The aim of the multi-layer approach is to model these relationships in a realistic manner.

In online social networks also the concept is very critical because users can have profiles on multiple social network websites and perform various activities in more
than one network at once which highlights the need for multi-layer social network analysis.

In this approach, each layer corresponds to a particular network. Thus, each of them has its own members and relations. Due to the multi-membership effect, these layers are connected to each other through their common members. As this is an emerging field, a standard method for representation has not been defined yet. Nevertheless, if \( m \) different layers exist in the system, \( G_i(V, E) \) represents a social graph of the layer \( i \) where \( 1 \leq i \leq m \). The graph can be weighted or unweighted, and directed or undirected.

Consequently, a multi-layer structure system can be represented by a multi-graph denoted by \( M \) as \( M = \{G_1, G_2, \ldots, G_m\} \). Hence, the set of nodes in \( M \) denoted by \( V(M) \) is defined as \( \bigcup_{i=1}^{m} V(G_i) \) where \( V(G_i) \) represents the set of nodes in the layer \( i \). Meanwhile, \( V(G_i) \cap V(G_j) \neq \emptyset, 1 \leq i, j \leq m, i \neq j \) which means that each layer has two sets of members, the independent and the shared ones. The shared members act as a bridge between the societies and have a substantial role in the co-evolution of the layers. As a result, many studies focus on their effects and characteristics in the network.

### 1.1.1 Characteristics of Social Networks

Social Networks as a subset of complex networks have the community structure which is indicated by a high level of clustering coefficient value. Clustering coefficient index in a graph measures the tendency of nodes to cluster together. It is defined as the fraction of node’s neighbors that are neighbors of each other. Meanwhile, social actors are willing to join the communities through their circle of friends and link to similar others which is referred to the homophily effect in social science. [31, 46, 34, 33, 30]

Moreover, social networks demonstrate the small-world phenomenon which means that social actors can be linked together through short chains of intermediate friends.
In other words, any two random nodes in the network can be linked by a short path in the graph. In addition, the degree distribution in these networks follow the power-law distribution. In fact, there exist relatively few nodes in the network with a high degree of connectivity and many nodes with low degree. [3, 1]

1.2 Research Problems and Objectives

This research focuses on the evolution of social networks with emphasis on the role of underlying knowledge in the evolution process. The ultimate goal of this research study is to employ graph theories, network science, and optimization methods to make a computational intelligence framework for describing the functionality of the complex dynamic social systems with the capability of exploring behaviors of these networks.

Utilizing different sources of knowledge extracted from the structure of the network in the analysis process is the main concept that distinguishes this work from the other existing approaches in the field of social network analysis.

Although the scope of social network analysis is very vast, finding relationships between social actors and discovering the underlying structures of the network are its fundamental tasks [7, 43, 21]. Therefore, community detection and link prediction problems which are two classic critical issues in the field are extensively studied in this work. In addition, a real-life problem is introduced and explored which is called population adaption through knowledge migration. To the best of our knowledge, this is the first work in the field that addresses this problem with this approach.

Consequently, the following research problems are addressed in this work through the proposed approach.

Community Detection

Social networks are highly interactive and dynamic systems which are consisted
of interconnected communities. Having knowledge about these communities can shed light on understanding the nature of these social systems which is essential for decision and policy making processes.

Briefly, community detection in social network is an NP-hard problem which deals with finding groups of actors who are more similar or close to each other than other ones in the other groups [50]. In other words, the goal is to find groups of people who have more relations and interactions with each other in the network. Each of this group is called a community. The issue can be seen as an optimization problem where the goal is to maximize the number of relations inside the group and minimize the links to the outside.

The central concept that distinguishes this problem from the classic graph clustering problems is that the number of communities is unknown in advanced while it must be known in the classic clustering problem.

**Link Prediction**

The problem of link prediction in social networks refers to exploring the dynamic nature of the network and its evolution. The link prediction problem can be defined as predicting the structure of a network in the near future by having a snapshot of it at the current time. In other words, given a state of a network at time $t$, the target is to estimate the likelihood of a connection between pairs of unconnected actors at time $t + 1$.

The problem has a broad range of application in recommendation systems, e-commerce, bioinformatic, politics, and security related issues. It can also be used to monitor the evolution of the system and identify the missing links between pairs of identities in the network [29, 2]. Identification of the missing links can be interpreted as finding the hidden connections between pairs of actors which are present in reality but were not observed.
Population Adaptation

Investigating the role of knowledge in the process of individual adaptation is the next goal of this research topic. The key question here is to find out how a population can perform in different environments when it has a prior knowledge about the similar environment? In other words, the question is how a population with prior knowledge about a problem can solve a similar problem and adapt itself to the new situation. Moreover, to what level of similarity between two situations, migrated population can be adapted efficiently? We define this concept as the adaptation process. The result of this research can lead to a remarkable reduction in the search time and space throughout the multiple steps of dynamic social network analysis.

1.3 Research Contributions

To address the research problems, a novel adaptive knowledge-based evolutionary computational model is proposed to study the structure and evolution of social networks by adopting a multi-population cultural algorithm (MPCA). This framework is capable of modeling the above three problems and their associated effects properly. In order to estimate the existence of a relationship between social entities in the network, a unique community oriented approach is defined which forms the core of the framework.

Briefly, the proposed model is designed based on the topological knowledge which is extracted from the structure of the network in each evolution cycle to form the normative knowledge. The knowledge is used to direct and enhance the search process to identify the proper sub-populations (Communities). The extracted knowledge can be employed in various network analysis tasks such as estimating the quality of relation between social entities, related studies regarding the link prediction, population adaptation, and knowledge formation.
To address the problem of community detection, we have proposed a novel knowledge-based evolutionary algorithm using a variant of MPCA [53]. Our first contribution is to propose a novel method to define, extract, and represent normative and domain knowledge sources from a snapshot of the network to determine the range of the optimal solution. The obtained knowledge is stored in a knowledge repository called belief space to guide the search direction and reduce the size of the required search space for finding the optimal/near optimal solution. As our second contribution, a unique data structure has been defined which is based on a probability matrix to form the belief space.

The results of comparison between our approach and other well-known related algorithms clearly show that our algorithm is capable of finding near optimal solution identifying the correct communities faster and more accurately than the others. Meanwhile, the evaluation results show that the search space can be reduced dramatically by 80% as a result of using our approach.

To deal with the link prediction problem in social network, we have proposed a community-oriented knowledge-based computational model which can estimate the next state of a given network with a notably high accuracy in [54]. By identifying the existing communities of the current state of a given network and make use of the belief space in [53], our proposed algorithm calculates the probability of a relationship between each unconnected pair of individuals and estimates the chance of being connected at the next time slot. A unique mapping function and a novel computational model based on the weighted graph have been introduced in this research to estimate the interdependency of each pair of individuals in the network.

We have tested and compared the model on synthetic networks and a big real standard data set from the Stanford large network dataset collection [50]. AUC (Area under the curve) and Precision measurements have been used to evaluate the performance of the model against several well-known other methods. The results show
that our method is able to predict the next state of the network with approximately 80% accuracy.

To cope the problem of population adaptation, the behavior and status of a dynamic social network have analyzed in a case where a population from one network migrates to another similar network and transfers its knowledge to it [56]. In effect, we have attempted to find how a migrated population will adapt itself to a new environment with similar characteristics based on the knowledge that it has learned from the previous network and what the role of this prior knowledge is in its evolution. As a case study we chose the problem of community detection in social networks.

To make a research framework we have adopted our previously proposed MPCA based community detection algorithm [53] for four different knowledge migration scenarios. For each scenario, two cases were examined: one case with individuals with prior knowledge about the similar networks, and another case with individuals without prior knowledge. The results show that when the changes in the structure of networks are less than 25%, trained population can adapt itself to the new network very fast; but when the difference is higher, in the best case they perform like a random population without any training.

As mentioned before, the significance of this contribution in dynamic social networks is that it allows us to use the extracted knowledge from a previous step, stored in the belief space, to detect new communities by eliminating the need for a new search if the similarity of two consecutive network snapshots is within 85%. This method can be generalized to accelerate the search performance in complex dynamic social networks.
1.4 Dissertation Outline

The rest of this dissertation is organized as follows.

In chapter 2, the problem of community detection in social networks as a fundamental task in social network analysis is extensively reviewed. Our proposed approach to deal with this issue is also described in the same chapter which forms the core of our research work.

In chapter 3, to extend the functionality of the community detection algorithm into a dynamic environment the problem of population adaption through knowledge migration are discussed.

In chapter 4, the problem of link prediction in social networks is reviewed. Our unique community-based approach which utilizes the extracted belief space to tackle the issue of link prediction is discussed in this chapter.

Finally, the last chapter will be the conclusion of this research.
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Chapter 2

Community Detection in Social Networks

Social networks can be viewed as a reflection of the real world which can be studied to gain insight into the real life societies and events. During the last decade, community detection as a fundamental part of social network analysis has been explored widely, however, because of the complex nature of the network, it is still an open problem. In this chapter, we propose a knowledge-based evolutionary algorithm to solve this problem by using a multi-population cultural algorithm. In our algorithm, knowledge is extracted from the network to guide the search direction and find the optimal/near optimal solution. Meanwhile, in each step, the knowledge is updated based on the current state of the network. The results of comparison between our method and other well-known algorithms show that our algorithm is capable of finding the true communities faster and more accurately than the others.
2.1 Introduction

Nowadays, more than 1 billion people around the world use online social networks to transfer and share their ideas, thoughts, experiments and willingness. Extracting knowledge from these networks can reveal their structure which has a lot of real-life applications such as marketing, group analysis, and decision making.

Generally, social networks consist of connected communities formed by individuals who communicate with each other. Finding these communities is a fundamental task in social network analysis. However, because of the complex and dynamic nature of these networks, identifying these communities is still an open challenging problem.

The first step to analyze a network is mapping it into a graph, $G(V, E)$, where $V$ is a set of nodes or agents and $E$ is a set of edges or links between agents. Let $A$ be an adjacency matrix for this graph. The entry of $A(i,j)$ is 1 if there is a direct link between nodes $i$ and $j$ otherwise it is 0 if no link exists. Accordingly, community detection in a social network can be seen as an optimization problem where the goal is to find groups of nodes that have more interconnections between each other and fewer intra-links with other nodes. The target is to find the best solution among all possible solutions to the problem \[8, 18\]. As the highlighted problem can be categorized as an NP-Hard problem, many researchers have proposed various methods based on evolutionary algorithms to solve it.

While most of the research are based on genetic algorithms, in this paper we use a different group of evolutionary algorithms which is known as cultural algorithms. The main feature of cultural algorithms that distinguish them from others is employing knowledge \[3, 21\]. In other words, it is a knowledge-based evolutionary algorithm. The cultural algorithm as shown in Fig. 2.1 is a dual inheritance model which consists of two main spaces, population, and culture or belief space. According to the model, in each generation, a group of individuals is selected to update the belief space and the new population is generated based on the parameters which were defined in the
belief space. The belief space in this model acts as a global knowledge repository which is made of information about the individuals and can be used to guide the search direction.

![Figure 2.1: Cultural Algorithm Process](image)

Our proposed algorithm is based on the multi-population cultural algorithm\[10\] which is illustrated in Fig. 2.2. To make the population spaces, a specific number of individuals are generated randomly based on the state space of the network. As the individual(a candidate solution) is composed of a combination of different elements, the state space of the network contains the possible states for each element. After the initial generation, in each population, a group of individuals that have better fitness values is selected to make a belief space. The belief space has a vital role in this algorithm and guides the search direction by determining a range of possible states for each element of the individuals. Hence, the belief space can be seen as a new state space for the network. Consequently, the new individuals in each population are generated based on this belief space. Meanwhile, in each step, the belief space is updated according to the state of the best-selected individuals of each population.
The rest of the paper is organized as follows. In the next section, we review major methods in this area. Section 3 contains the proposed algorithm. Evaluation and results are reviewed in Section 4, and conclusions are represented in section 5.

### 2.2 Related Works

In recent years, different methods have been proposed to solve the community detection problem. One of the most important method which became the base for further research in this field was proposed by Girvan and Newman [6]. In this paper, the concept of modularity was defined, and a divisive method was proposed for the problem. Many researchers proposed different algorithms based on the concept of modularity with various approaches. However, some of these algorithms need prior knowledge of the network, and some others have poor performance on large complex networks [2, 7, 18]. To cope with these drawbacks, researchers have employed evolutionary algorithms by different approaches and techniques. However, the common goal is to detect unknown numbers of communities in the network with a high level of internal connections and low level of external links [1, 2, 6, 7, 8, 9].

Some research focuses particularly on enhancing the fitness function. A fitness function has critical importance in evolutionary algorithms, as it estimates how close
the solution is to the final solution and consequently guides the algorithm direction directly or indirectly. For this purpose, some recent studies have addressed the problem as a multi-objective problem. The first objective aims at maximizing the internal links and the second is minimizing the external connections [1, 2, 4, 6, 9, 13, 14]. Pizzuti [18] proposed a new algorithm to solve the problem by using a genetic algorithm. The author has used the density measure and has defined the new concept of community score as a global measure to partition a given network into clusters. The goal of the algorithm is to maximize this score.

In Facetnet [13], the authors have proposed a new framework to solve the problem by using a multi-objective evolutionary algorithm. In their model, an individual can be a member of more than one community at the same time. They have defined the snapshot quality function and the temporal cost function and an iterative algorithm which uses a function to update rules in order to decrease the value of the cost function uniformly. On the other hand, they have introduced concepts of community membership, community net and evolution net in their framework. Meanwhile, they have proposed a mechanism for adding and removing individuals from communities to cope with the dynamic aspect of the network. A soft modularity function to measure the effectiveness of a community was also employed.

Some recent research uses the NSGA-II (Non-dominated Sorting Genetic Algorithm) as the core of their algorithm [2, 9, 19]. Kim, Mckey, and Moon [9] propose HIGA (hybrid immigrants GA) to cope with the dynamic aspect of the network. The authors have defined an algorithm called Adaptive Immigrants NSGA-II (AI-NSGA II) to give their method dynamic adaptability. The min-max cut and global silhouette index defined as the two objectives of the fitness function. On the other hand, Chen, Wang, and Wei [2] have employed the modularity function and NMI as similarity measures for the first and second objectives. They have also used community score [18] for the solution selection process.
Many studies have also been carried out based on other techniques [1, 6, 7, 20, 15]. Gong et al. [6] proposed a multi-objective algorithm based on the non-dominated neighbor immune algorithm (NNIA). For the first objective they used the modularity function [5] and for the second, they used NMI as a similarity measure. Amiri, Hossain, and Crawford [1] have suggested a multi-objective evolutionary algorithm based on the harmony search algorithm. Jia et al. [7] proposed a Differential Evolution (DE) approach to solving the problem. Modularity function was employed to obtain the fitness function while for the initialization step a particular biased process was used in order to prevent making unreasonable results. Furthermore, for the mutation they used "rand/1" strategy. Qiu and Lin [20] proposed a new algorithm to solve the problem by using a hierarchical structure model. Random walk approach was implemented and the Gaussian Mixture Model (GMM) was used to generate the transition probability matrix to calculate the likelihood of relation between a node and each community.

2.3 Proposed Model for Community Detection

In this section, we describe the proposed algorithm which is a multi-population cultural algorithm for community detection in social networks. The individual representation method and mechanism for crossover and mutation are described in detail in the next part. After that, the structure of belief space will be defined and discussed.

2.3.1 Individual Representation

The representation of an individual in our algorithm is based on a particular locus-based adjacency representation method [17]. The individual or a candidate solution is represented by an array of nodes. The length of the array is equal to the number of nodes in the graph. Each cell of this array is identified by a number which corresponds
to the number of nodes in the graph. For example, cell#5 refers to the node #5. The value of each cell is chosen randomly from the state space of the network denoted by NS, which is formed based on the adjacency matrix of the network graph. Therefore, for each node in the graph, a set of neighbor nodes is defined as follows:

Algorithm 1 Network state
1: procedure MAIN(Adjacency)
2:     for $i \leftarrow 1 : n$ do
3:         for $j \leftarrow i + 1 : n$ do
4:             if $A(i, j) = 1$ then
5:                 $NS(i, j) \leftarrow j$
6:                 $NS(j, i) \leftarrow i$
7:             end if
8:         end for
9:     end for
10: end procedure

As an example, Fig. 2.4 illustrates the network state space of a network graph which has been shown in Fig. 2.3. In addition, Fig. 2.5 illustrates two different random representations of the network based on the network state.

![Figure 2.3: A network graph](image-url)
2.3.2 Initialization

A specific number of individuals are generated randomly based on the individual representation method to form the population spaces. One common problem that usually occurs in the other algorithms in the initialization phase is that some of the individuals are not valid. It means that the individual contains some links which do not exist in the original graph. However in our algorithm, as individuals are generated based on the network state space, existence of the links can be assured because elements of each individual are selected randomly from valid neighbors nodes.

2.3.3 Fitness Function

The objective function is another important factor in the evolutionary algorithms. This function has a key role in guiding the direction of the evolution. However, our
algorithm is independent of it and can work with any form of objective function if it is adaptable with our representation method. Nevertheless, we have employed community score, one of the best-known fitness function which has been defined by Pizzuti [18]. This function can work without prior knowledge of the number and size of the communities, and its goal is to maximize the community score. In this paper, the same fitness function is used for all populations, but it is possible to have different fitness functions in each population.

Let $N = \{C_1, C_2, ..., C_k\}$, denotes the network which consists of different communities. The score of each community is calculated as shown in the following equation, Eq. 2.1.

$$Q(C_k) = \frac{\sum_i \left( \frac{\sum_j a_{i,j}}{|J|} \right)^r}{|I|} \times \sum_{i,j} a_{i,j}$$

(2.1)

Where $i, j \in C_k$ and $a_{i,j}$ denotes a value of the position $(i,j)$ of the adjacency matrix. In addition, $|J|, |I|$ denote the number of $j$ and $i$ in the $C$ respectively. Finally, the community score is the summation of all communities’ scores in the graph then:

$$CS = \sum_{1}^{k} Q(C_k)$$

(2.2)

2.3.4 Belief Space

The core of our algorithm is the belief space which is formed by the selected individuals of each population in every generation to guide the direction of the evolution. We consider that the best solution can be represented by combining elements of the best-selected individuals. In fact, the idea is, instead of searching all possible states and combinations, the search space must be limited to the elements of the selected population. Therefore, in each generation, the belief space defines a range of the best feasible solutions. Consequently, the new population is generated in the range which
has been defined in the belief space. It is expected that in each generation, better solutions are being generated by the algorithm.

We define two different sources of knowledge in the belief space. The first is called BS\_average and stores the best ever average fitness value of the previously selected populations. In each generation, a selected individual can change the belief space if its fitness value is higher than the average value of the previous individuals which influenced the belief space (BS\_average). As shown in Alg. 2, if BS\_average is less than the average fitness value of the currently selected population it must be replaced with the new one. As mentioned before, each individual is represented by an array with the length of n which is the number of nodes. Therefore the selected population can be presented by an s by n matrix where each row of the matrix shows an individual and s is the size of the selected population and n is the number of nodes in the graph. As shown in Eq. 2.4, let SP denotes the selected population which consists of selected individuals (SI) in Eq. 2.3 then the average is computed by calculating the average of fitness values of the selected individuals. It has been represented in Eq. 2.5.

\[ SI = [s_{i_1}, s_{i_2}, ..., s_{i_n}] \]  

\[ SP = \begin{bmatrix} SI_1 \\ \vdots \\ SI_s \end{bmatrix} \quad \rightarrow \quad SP = \begin{bmatrix} s_{i_{1,1}} & \cdots & s_{i_{1,n}} \\ \vdots & \cdots & \vdots \\ s_{i_{s,1}} & \cdots & s_{i_{s,n}} \end{bmatrix} \]  

\[ Average = \frac{\sum_{j=1}^{s} Fitness(Si_j)}{s} \]  

The second source of knowledge is the normative knowledge, BSN, which is represented by an n by n matrix. In each generation, for all individuals of the selected population, the relative frequency of values of all cells are calculated and added into the corresponding entry in the matrix. In fact, as shown in Alg. 2 for all the selected individuals, BSN(j, value(j)) is updated with the relative frequency of the value(j) in
the cell\#j where j is the cell number.

**Algorithm 2 Update Beleif Space**

1: function Update(SP) \[\triangleright\ Selected\ Population\]
2: \( s \leftarrow |SP| \)
3: \( n \leftarrow |SI| \)
4: for \( i \leftarrow 1 \) to \( s \) do
5: \[\text{if } \text{Fitness}(SI_i) > BS\_\text{average} \text{ then}\]
6: \[\quad \text{for } j \leftarrow 1 \) to \( n \) do\]
7: \[\quad BSN(j, \text{value}(j)) \leftarrow \text{relative frequency value}(j) \text{ in row}j\]
8: \[\quad \text{end for}\]
9: \[\quad \text{end if}\]
10: \[\text{end for}\]
11: \[\text{if } BS\_\text{average} < \text{Average} \text{ then}\]
12: \[\quad BS\_\text{average} = \text{Average}\]
13: \[\text{end if}\]
14: end function

For example, in Fig. 3.6, the BSN is formed based on the network with eight nodes and four selected individuals. The first row shows neighbors of the node 1. According to the matrix, the probability of connection between node 1 and node 2 in the final solution is 75\% while it is 25\% for node 3. It means that in the next generation, node 2 will be presented in the first cell of the individuals with a probability of 75\% while node 3 with the probability of 25\%. 
2.3.5 Crossover and Mutation

As populations are generated based on the belief space, the role of Crossover and Mutation operators is completely different in comparison with Genetic algorithms. In fact, these operators help the algorithm to escape from the local maxima.

The algorithm presented in this paper is based on the multi-population spaces. Therefore, each population can have its own crossover and mutation operators. However, to choose a parent for the crossover in the first population, the first individual is selected randomly among all individuals and the second one is randomly chosen among individuals who are not in the selected population. Given these two individuals, a new individual will be generated by combining the parent’s elements. As the parents in these operators are selected among all individuals, the chance of having children with entirely different elements is very high. For the mutation operator, an individual is generated based on the network state space similar to the first generation. It is expected that the algorithm escapes from the local maxima and generates
some solutions outside the current domain by using crossover and mutation

2.3.6 Our proposed algorithm

Our proposed algorithm is started by generating the initial population, after evaluating the fitness of individuals and sorting them, the best groups of individuals of each population are selected based on their fitness function. These groups update the belief space. The new generation of individuals is generated based on the probability matrix of the belief space. Meanwhile, in each iteration with a small probability some individuals are generated by crossover or mutation operators. Each population space in this algorithm can have their own fitness function or operators. The algorithm continues until the last iteration, and the individual with the best fitness function value will be presented as the best solution.
Algorithm 3 MPCA-CD

1: procedure Main(Adjacency)
2:     NS ← MakeNS(Adjacency) \Comment{Initialize the network State}
3:     l ← |Selected Individuals| \Comment{Define the number of selected individuals}
4:     el ← |Elite Individuals| \Comment{Define the number of elite individuals}
5:     Pop ← Represent(NS) \Comment{Initialize the populations}
6:     F ← Fitness(Pop) \Comment{Evaluate the individuals’ fitness values}
7:     Pop ← Sort(Pop, F) \Comment{Sort the individuals by their fitness values}
8:     SP ← Select(Pop, l) \Comment{Select individuals to update the belief space}
9:     Belief = Update(SP) \Comment{Select individuals to update the belief space}
10:    loop \Comment{Start the Loop}
11:     Pop(el : end) ← Represent(Belief) \Comment{generate the populations based on}
     the Belief space
12:     F ← Fitness(Pop)
13:     Pop ← Sort(Pop, F)
14:     SP ← Select(Pop, l)
15:     Belief = Update(SP)
16:     if < stopcriteria > then
17:         Break
18:     end if
19:     end loop
20: end procedure

2.4 Evaluation

To evaluate the effectiveness of the model, we compare it with four well-known algorithms in this field. The first one is GA-Net [18] which is a genetic algorithm,
the second one is the Girvan-Newman algorithm (GN), DECD [7] is the third one which is based on Differential Evolution and the last one is MOGA-Net [19]. We also compare it with variable-CA [24]. To measure the similarity level between the true communities and the detected ones we used Normalized Mutual Information (NMI) [5, 18].

We made 60 artificial networks based on the Newman benchmark [5, 17]. Each network was generated randomly and has 128 nodes which are categorized in 4 same-sized communities with 32 nodes while the degree of each node was 16. Meanwhile, each node is connected to other nodes in its community by internal degree, $Z_{in}$, and to other nodes by external degree, $Z_{out}$ ($Z_{in} + Z_{out} = 16$). The range of $Z_{out}$ of our artificial networks in this experiment is from 1 to 6 where six implies that each node is connected to 6 nodes outside of its community which means that the network is very noisy and fuzzy.

Our proposed algorithm has been implemented in Matlab, and all tests have been performed on a Pentium dual core 2.1GHz with 2.5 GB RAM. In addition, for the first population, crossover and mutation rate were set to 0.8 and 0.2 respectively. The population size was 200, and the number of generations is set to 50 while roulette selection function was used. For the second population, the size was set to 100, and the selection rate was 20% similar to the first one, but the rate of mutation increased to 50% while the roulette wheel selection was used.

As demonstrated in Fig. 2.7, the proposed algorithm can detect actual communities with 100% success when $Z_{out}$ is less than or equal to 5 while none of other algorithms can achieve this rate. For $Z_{out}$ of 5, the average NMI of our algorithm is 1 while the value for GN, GA, DECD, and MOGA-Net are 0.72, 0.77, 0.95 and 0.98 respectively. Even when $Z_{out}$ becomes 6, our algorithm has better performance when compared to the others, and its value was 0.83 while the best value of other algorithms was achieved by MOGA-Net which was 0.67. In addition, this value for
the variable-CA was 0.69.

For the real datasets, we have employed Zachary Karate club \[23\], Dolphin networks \[16\] and American Football \[23\]. Zachary Karate Club dataset was made by Zachary as a result of a study on the friendship of 34 members of a karate club during two years. The group was split into two groups because of some disagreements, and it has 34 nodes in two groups. Our algorithm detected two communities on just three generations on this dataset. The average NMI value for our algorithm in this dataset as shown in Table. 2.1 was 1 which is same as MOGA-NET. The value was 0.82 and 0.69 for the GA and GN algorithms respectively.

For the dolphin dataset which has 62 nodes and was generated based on statistics obtained from seven years of dolphins’ behavior, the average NMI over ten different attempts was 1 for the MOGA-Net and 0.956 for our algorithm and was 0.935 for the GA.

American Football dataset was made based on the United state college football
information and has 115 nodes and 616 edges which were grouped into 12 teams. Our algorithm achieved the highest NMI value among other algorithms in this dataset whose value is 0.923.

<table>
<thead>
<tr>
<th>Dataset/Algorithm</th>
<th>GN</th>
<th>MOGA-NET</th>
<th>GA</th>
<th>MPCA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zakhary</td>
<td>0.692</td>
<td>1.000</td>
<td>0.818</td>
<td>1.000</td>
</tr>
<tr>
<td>Dolphin</td>
<td>0.574</td>
<td>1.000</td>
<td>0.935</td>
<td>0.956</td>
</tr>
<tr>
<td>American Football</td>
<td>0.760</td>
<td>0.796</td>
<td>0.805</td>
<td><strong>0.923</strong></td>
</tr>
</tbody>
</table>

These results clearly highlight that the method presented in this paper gives a better performance than other algorithms and can detect the true communities in noisy networks as well. Another important point is that, as the method is based on belief space, it is expected that the algorithm provides better performance with an increase in the size of the network since more nodes can create a rich belief space to guide the search direction while the search domain narrows down at each step.

**2.5 Discussion and Conclusion**

In this section, the results obtained during the process of community detection are extensively reviewed and discussed. At the first step, we would like to show how the algorithm identifies the optimal/ near optimal solution during the evolution cycles. After that, the role of knowledge in the reduction of search space are discussed. Moreover, two more structures are reviewed to form the belief space. Runtime analysis and the results of running the algorithm on large networks are reported at the end of this section.
2.5.1 Demonstration of the Evolution Process

To demonstrate how the individuals (candidate solutions) are evolved during the process, we ran the algorithm on a random benchmark graph, and we have selected the best individual of each generation. The graph has been generated based on the Newman benchmark with 128 nodes and $Z_{out}$ of 5 which means that it has a complex structure. Therefore, it has four communities where the nodes 1 to 32 are placed in the first community, nodes 33 to 64 forms the second, nodes 65 to 96 shapes the third one and 97 to 128 are placed in the fourth community. The test has been repeated 10 times independently, and the NMI value of each individual has been calculated and illustrated in Fig. 2.8 (the experiments are named 1 to 10). As shown in the figure, the first results of all experiments are very far from the optimal solution, but during the process, they are evolved to reach the optimal point. In fact, the rate of evolution is increased after the 10th iteration. Therefore, this interval can be interpreted as the learning period for the algorithm.

![Figure 2.8: Evolution of individuals in a network with $Z_{out} = 5$](image)

Figure 2.8: Evolution of individuals in a network with $Z_{out} = 5$
To clarify the subject, the detected communities (Decoded best individual) at the end of iterations 1, 10, 18, 20, 23, 24 and 25 of the first experiment are shown in Figures. 2.9, 2.10, 2.11, 2.12, 2.13, 2.14, 2.15.

Fig. 2.9 shows the obtained result after the first iteration, the algorithm found just two communities in the network. For example, nodes 33, 35, 36, 37, 38, 39, 40, 43, 45, 46, 48, 49, 50, 51, 52, 53, 57, 58, 59, 63, 64, 66, and 112 are categorized together in the second community. The NMI value for this solution is 0.25901 as illustrated in Fig. 2.8. The best individuals do not change until the end of the 10th iteration. At this point, as shown in Fig. 2.10, the best individual generated by the algorithm has identified five groups in the network and obtained a higher NMI value.

As illustrated in Fig. 2.11, after the 18th iteration, most of the nodes were clustered correctly, but the number of groups is still incorrect. In the 20th iteration, the algorithm identified four communities. However, some nodes were categorized into wrong communities. For example, as shown in Fig. 2.12, the nodes 61 and 85 have
been clustered in the first community which is not correct.

Figure 2.10: Identified communities at iteration 10

Figure 2.11: Identified communities at iteration 18
Figure 2.12: Identified communities at iteration 20

Figure 2.13: Identified communities at iteration 23
Figure 2.14: Identified communities at iteration 24

Figure 2.15: Identified communities at iteration 25
The evolution cycles still continue to find the optimal/near optimal value. The obtained results at the end of iteration 23 and 24 are illustrated in Fig. 2.13 and Fig. 2.14 respectively. In iteration 24, the node 61 is the only one who has placed wrongly. Finally, the algorithm found the best solution in iteration 25 and identified the four communities correctly as demonstrated in Fig. 2.15.

We have repeated the experiment for 100 times on various synthetic networks, and we observed that the results follow the similar evolution patterns. For example, when $Z_{out}$ is 5, the algorithm can find the optimal value at around iteration 25, and a significant change in the NMI value is expected at around iteration 10.

![Figure 2.16: Evolution of individuals on a synthetic network with $Z_{out} = 5$](image)

To demonstrate the role of knowledge in the evolution process, we ran ten more experiments on a random synthetic graph which was generated based on the Newman method with $Z_{out}$ of 5. The result is demonstrated in Fig. 2.16. After that, we utilized the genetic algorithm (without a knowledge structure) to detect the communities on the same graph. The results show that, this algorithm could not find any communities
in the graph and the NMI values were always 0. It clearly indicates that, the extracted knowledge is a powerful tool which can guide the search process, especially in the complex structures.

In our next attempt, we set a new configuration and changed the value of variable \( r \) in the fitness function from 1.0 to 1.7. The number of iterations has also been set to 30. Again, we ran our MPCA ten times and captured the NMI values of the best individuals of the iterations. After that, the genetic algorithm was executed on the same network for ten times and the NMI values were captured. Fig. 2.17 and Fig. 2.18 show the results of our proposed algorithm and the genetic one respectively.

As demonstrated in the figures, the rate of evolution in the knowledge-based algorithm is higher than the genetic method. In fact, at the first generations, the rate is almost the same for the both algorithms, but from approximately 8th iteration, the cultural algorithm has a faster progress. In addition, MPCA could find the optimal solution in average by 23 iterations while the genetic one could not find it at all.

![Figure 2.17: Evolution of individuals-MPCA- \( Z_{out} = 5 \) - \( r=1.7 \)](image-url)
According to the above experiments, it is clear that, our proposed MPCA can significantly improve the search performance in the both levels of accuracy and run-time compare to the genetic algorithm.

2.5.2 The Role of Knowledge in Search Space Reduction

To show the role of knowledge in the reduction of the search space, a snapshot of the network state space of a graph with $Z_{out} = 5$ has been compared to the obtained belief space. Fig. 2.19 illustrates the neighbors of the first 32 nodes.
Figure 2.19: A snapshot of the network state space in a graph with $Z_{out} = 5$

For example the first node in this graph has 16 neighbors which are nodes 3, 4, 6, 7, 18, 20, 24, 26, 28, 29, 38, 53, 71, 96, and 123. It means that in the initial stage, an individual can randomly choose one of them as the value of its first cell. The situation is similar for the other cells. Hence, the algorithm must search among 16 possible nodes for each cell of the individual’s array.

One of the primary goals of this algorithm is to optimize the search space by using different sources of knowledge. Because of this, the belief space was introduced to enhance the search process. In Fig. 2.20, a snapshot of the belief space for the same set of nodes (1 to 32) in the graph has been illustrated after the 16th iteration. As the figure shows, by using the belief space, the neighbors of the first node has been reduced dramatically from 16 to just 3 (27, 18, 7) nodes. It implies that the search domain became about 5 times smaller than the original network. The situation is almost the same for the other nodes. Hence, instead of searching among 16 neighbors for each cell, the algorithm just need to search between two or three neighbors (the
exception is node 18 which has 4 neighbors in the belief space) for each one to find the optimal/near optimal solution which proves that the belief space can significantly reduce the search domain (above 80%).

\[
\begin{align*}
(1,27) &= 0.5 \\
(1,18) &= 0.3125 \\
(1,7) &= 0.1875 \\
(2,19) &= 0.625 \\
(2,16) &= 0.375 \\
(3,30) &= 0.6875 \\
(3,29) &= 0.3125 \\
(4,5) &= 0.75 \\
(4,17) &= 0.25 \\
(5,19) &= 0.5625 \\
(5,9) &= 0.4375 \\
(6,21) &= 0.5 \\
(6,24) &= 0.4375 \\
(6,26) &= 0.0625 \\
(7,31) &= 0.5 \\
(7,14) &= 0.375 \\
(7,21) &= 0.125 \\
(8,15) &= 0.625 \\
(8,2) &= 0.375 \\
(9,20) &= 0.0375 \\
(9,24) &= 0.0625 \\
(10,8) &= 1 \\
(11,6) &= 0.6875 \\
(11,27) &= 0.3125 \\
(12,27) &= 0.75 \\
(11,20) &= 0.25 \\
(13,15) &= 0.5 \\
(13,29) &= 0.3125 \\
(13,3) &= 0.1875 \\
(14,20) &= 0.5625 \\
(14,17) &= 0.4375 \\
(15,16) &= 0.375 \\
(15,19) &= 0.3125 \\
(15,26) &= 0.3125 \\
(16,19) &= 1 \\
(17,30) &= 1 \\
(18,2) &= 0.5 \\
(18,8) &= 0.3125 \\
(18,32) &= 0.125 \\
(18,10) &= 0.0625 \\
(19,26) &= 1 \\
(20,30) &= 0.875 \\
(20,5) &= 0.0625 \\
(20,31) &= 0.0625 \\
(21,9) &= 0.9375 \\
(21,5) &= 0.0625 \\
(22,15) &= 0.8125 \\
(22,26) &= 0.1875 \\
(23,16) &= 0.625 \\
(23,22) &= 0.375 \\
(24,22) &= 1 \\
(25,21) &= 0.8125 \\
(25,8) &= 0.1875 \\
(26,6) &= 0.5 \\
(26,32) &= 0.4375 \\
(26,1) &= 0.0625 \\
(27,19) &= 1 \\
(28,5) &= 0.5625 \\
(28,9) &= 0.4375 \\
(29,17) &= 0.375 \\
(29,19) &= 0.375 \\
(29,5) &= 0.25 \\
(30,32) &= 0.75 \\
(30,13) &= 0.25 \\
(31,20) &= 0.5625 \\
(31,20) &= 0.4375 \\
(32,13) &= 0.4375 \\
(32,26) &= 0.4375 \\
(32,4) &= 0.125
\end{align*}
\]

Figure 2.20: A snapshot of the belief space after the 16th iteration (the first 32 nodes)

### 2.5.3 The Structure of Belief Space

In section 2.3.4, we introduced a data structure to form the belief space. In this section, two more structures to shape the belief space are discussed.

In the first structure that we call it fixed-size belief space, all individuals of the selected population are involved in the update process. For this purpose, the belief space must be an \( n \) by \( s \) matrix where \( n \) is the number of nodes in the graph, and \( s \) is
the number of individuals in the selected population. In each update process, for all individuals of the selected population, values of all cells are added to the corresponded rows of the matrix.

In this case, to update the culture, the matrix must become empty before the new entries are added to it. Therefore, the size of the belief space is always fixed. As we mentioned before, each individual is represented by an array with the length of \( n \). Therefore, the selected population can be presented by an \( s \) by \( n \) matrix where each row of it shows an individual as shown in the equation 2.4. Accordingly, the belief space can be defined as the transpose matrix of the selected population.

Let \( SP \) denotes the selected population which consists of selected individuals (SI) the belief space, \( BS \), can be defined as:

\[
BS = SP^T = \begin{bmatrix}
s_{i_{1,1}} & \ldots & s_{i_{s,1}} \\
\vdots & \ddots & \vdots \\
s_{i_{n,1}} & \ldots & s_{i_{n,s}}
\end{bmatrix}
\]  

(2.6)

For example, as shown in Fig. 2.21, if the graph has 8 nodes, the belief space is defined as an \( 8 \times s \) matrix where the row \( #i \) in the matrix is filled with values of the cell \( #i \) of the selected individuals.

![Figure 2.21: A sample belief space with the fixed-size structure](image)
The second structure called variable-sized belief space, is very similar to the real situation. In real societies, culture is frequently updated. However, the previous cultural elements are not discarded, and they remain in the system while some of them become stronger, or weaker. Therefore, in each update process, instead of erasing the belief space, new elements are appended to the matrix. To employ this structure the size of the belief space must be equal to $n \times s \times g$ where $g$ is the number of iterations. As an example, if the graph has 100 nodes and the size of the selected population is 20, and the maximum iteration is 30, the belief space size becomes 60000.

We have evaluated the performance of each method using the test framework defined before in Section 2.4. We compared the average number of generations which were occurred to reach the best solution in different algorithms. As Fig. 2.22 shows, all different versions of our algorithm can find the best solution in fewer generations than the other algorithms. For example, when $Z_{out}$ is 5, CA-Var (variable-sized belief space), Ca-Fix (Fixed-size belief space) and MPCA found the correct answers in average by 28, 30, 25 generations respectively while other methods could not find it.

On the other hand, in compare with the fixed-size belief space, the variable approach shows a better performance on the conducted tests. However, according to the results, MPCA has the best performance among them.
2.5.4 Run-Time Analysis

Our proposed algorithm has four principal components which are the individual representation, fitness function evaluation, decoding phase and culture modification. In other words, the run-time of the algorithm is directly dependent on the performance of these components. In this section, the performance of these elements is examined and reviewed.

The algorithm starts with the individual representation. In the locus-based representation, the search space of each node is limited to its adjacent nodes. Therefore, it has a complexity of $O(d^n)$ where $n$ is the number of nodes in the graph, and $d$ is the degree of nodes. Because of the fact that the degree of the nodes in social networks follows the power-law distribution, the value of $d$ is usually much smaller than $n$.

To compute the fitness value, at first, the individual must be decoded. For the decoding phase, we employed and modified a backtracking algorithm which has been
proposed in [22] and demonstrated in Algorithm 4. This algorithm can decode a
given individual in a linear time with the complexity of \(O(n)\). Moreover, the fitness
function is the most critical and time-consuming component of this model which must
be run for all generated individuals and has the complexity of \(O(m)\) where \(m\) is the
number of edges in the network. In addition, the complexity of culture modification
is almost constant and ignorable.

Hence, the run time complexity of each iteration in our proposed algorithm is
\(O(m+n)\), where \(n\) is usually much smaller than \(m\). For the whole algorithm, the time
complexity can be represented as \(O(gs(m+n))\) where \(g\) is the number of iterations,
and \(s\) is the size of the population.

\begin{algorithm}
\caption{Decoding}
\begin{algorithmic}
\Procedure{Decode}{Individual}
\State \(n \leftarrow \text{size}(I)\) \Comment{n = the number of nodes}
\State \(\text{Current}_\text{community} \leftarrow 1\) \Comment{\# of found community}
\State \(\text{Community}_\text{assign}(1 : n) \leftarrow -1\) \Comment{Initialize the array}
\For{i \leftarrow 1 \text{ to } n}
\State \(Ctr \leftarrow 1\) \Comment{Community index}
\If{\(\text{Community}_\text{assign}(i) = -1\)}
\State \(\text{Community}_\text{assign}(i) \leftarrow \text{Current}_\text{Community}\) \Comment{Assign a}
\State \(\text{Community number to the node } \# \ i\)
\EndIf
\EndFor
\State \(\text{Neighbor} \leftarrow I(i)\) \Comment{The entry(i) from the Individual Array becomes}
\State \(\text{neighbor of node } \# \ i\)
\State \(\text{Previous}(Ctr) \leftarrow i\) \Comment{Save the previous community index (used in}
\State \(\text{backtrack process)}\)
\EndProcedure
\end{algorithmic}
\end{algorithm}
Algorithm 4 Decoding - Page 2

11: \textbf{if} $I(i) \neq 0$ \textbf{then}

12: \hspace{1em} $Ctr \leftarrow Ctr + 1$ \t $	riangleright$ Increase the Community index

13: \hspace{1em} \textbf{while} $\text{Community}\_assign(\text{Neighbor}) = -1$ \textbf{do} \t \textbf{While this node}

14: \hspace{2em} \textbf{if} $I(\text{neighbor}) \neq 0$ \textbf{then}

15: \hspace{3em} $Previous(Ctr) \leftarrow \text{Neighbor}$

16: \hspace{3em} $\text{Community}\_assign(\text{Neighbor}) \leftarrow \text{Current}\_Community$ \t \textbf{Assign the same Community number to the neighbor node}

17: \hspace{3em} $\text{Neighbor} = I(\text{neighbor})$ \t \textbf{Change the neighbor}

18: \hspace{3em} $Ctr \leftarrow Ctr + 1$

19: \hspace{2em} \textbf{end if}

20: \hspace{1em} \textbf{end while}

21: \hspace{1em} \textbf{end if}

22: \hspace{1em} \textbf{if} $\text{Community}\_assign(\text{Previous}(Ctr)) \neq \text{Current}\_Community$ \textbf{then}

23: \hspace{2em} $Ctr \leftarrow Ctr - 1$ \t \textbf{Decrease the Community index}

24: \hspace{2em} \textbf{while} $Ctr \geq 1$ \textbf{do} \t \textbf{Backtrack condition}

25: \hspace{3em} $\text{Community}\_assign(\text{Previous}(Ctr)) \leftarrow \text{Community}\_assign(\text{Neighbor})$ \t \textbf{Assign a Community number}

26: \hspace{3em} $Ctr \leftarrow Ctr - 1$

27: \hspace{2em} \textbf{end while}

28: \hspace{1em} \textbf{else}

29: \hspace{2em} $\text{Current}\_Community \leftarrow \text{Current}\_Community + 1$ \t \textbf{Create}

30: \hspace{2em} \textbf{another Community}

31: \hspace{1em} \textbf{end if}

32: \hspace{1em} \textbf{end if}

33: \hspace{1em} \textbf{end for}

34: \hspace{1em} \textbf{end procedure}

In order to analyze the run-time performance of our proposed algorithm, various
experiments have been carried out on 50 synthetic networks ranging from 128 nodes to 10000 nodes. To generate the network, we used LFR benchmark in [12, 11]. By using this benchmark, we can create power-law networks with customized features. Our networks have been generated in seven categories of 128, 512, 1024, 2000, 3000, 5000 and 10000 nodes.

The following parameters have been considered to generate these networks:

- $\beta = 1$, $\beta$ set the exponent for the distribution of community size in the network
- $\gamma = 2$: $\gamma$ set the exponent for the nodes’ degree distribution.
- $\mu = 0.3$, $\mu$ is the mixing parameter which determines the ratio of the number of edges between various communities to the total number of them. The higher number means more complex community structure.
- $D_{Average} = 20$, $D_{Average}$ represents the average degree of each node in the graph.
- $D_{Max} = 50$, $D_{Max}$ set the maximum degree size for each node.

The obtained results show that the runtime of the algorithm is increased linearly on the scale of the network. For example, in Fig. 2.23 the average runtime of the algorithm on a synthetic network has been illustrated. In this example, the algorithm ran on the network with five different population size of 50, 100, 200, 300, and 400. For each of these population, the number of iterations has been set to 10, 20, 30, 40, 50, 60 and 100. The algorithm ran 10 times for each option, and in total 350 independent experiments have been carried out.

As seen in the chart, the rate of increase for all of the iterations is linear. For example, when the population size is 100, and the number of iterations is 30, the runtime is 0.444s. By increasing the size of the population to 300, the runtime increases to 1.32s. In another instance, when the size of the population and the number of iterations are both 50, the runtime is 0.3777s, but when the population
size is changed to 400, the runtime also is increased to 2.8s which clearly indicates the linear nature of the growth.

On the other hand, by increasing the number of iterations, the runtime will increase respectively. For instance, when the population size and the number of iterations were 200 and 10 respectively, the obtained runtime was 0.371, but by changing the iterations number to 50, the runtime gradually increased to 1.445s.

![Runtime analysis on variable population size and iterations](image)

**Figure 2.23:** Runtime analysis on variable population size and iterations

Fig. 2.24 demonstrates the runtime details of each process when the population size is 50. As hypothesized, the fitness evaluation is the most time-consuming component of our algorithm. The decoding process is another element in the system which grows slightly during the process. We have to mention that, the range of obtained values for the individual representation process were slightly different to the decoding process. Moreover, updating the culture is the least time-consuming part of this system. As seen in the chart, in regards to the fitness component, the runtime of other elements
are ignorable.

![Figure 2.24: Runtime analysis- Population size = 50](image)

The obtained results for other populations are also demonstrated in Figures 2.25, 2.26, 2.27, 2.28. The results show that, regardless of the population size and the number of iterations, fitness evaluation plays the major role in increasing the algorithm running time which is predictable. On the other hand, it clearly shows that adding the belief space has a minor impact on the complexity which is ignorable in regards to its significant impact on the performance improvement.
Figure 2.25: Runtime analysis- Population size = 100

Figure 2.26: Runtime analysis- Population size = 200
Figure 2.27: Runtime analysis- Population size = 300

Figure 2.28: Runtime analysis- Population size = 400
Finally in Fig. 2.29 we have illustrated the average overall runtime of the algorithm on synthetic graphs ranging from 128 nodes to 10000 nodes. The obtained results confirm that the runtime of the algorithm increases linearly in regards to the scale of the network.

![Graph showing runtime vs network size](image)

Figure 2.29: Runtime- Network size from 128 to 10000 nodes

### 2.5.5 More Evaluations

In this part we evaluate the performance of our proposed algorithm on more datasets. Our datasets have been generated based on LFR benchmark which has been described in the previous section. For the first experiment, we have created 18 synthetic graphs with variable sizes ranging from 128 to 1000 nodes. The \( \mu \) parameter were set from 0.1 to 0.6. Generally, when \( \mu \) is bigger than 0.4, the number of links between members of the community is almost similar or smaller than the external links. The degrees of the networks also follow the power-law degree distribution. For instance, Fig. 2.30 shows the Probability Density function (PDF) of the degree (in Logarithmic Bins) of the network with 1000 nodes and \( \mu = 6 \).
We ran the algorithm 10 times on each of these networks and compared the obtained NMI values with the results achieved by the genetic algorithm on the same datasets. The results are illustrated in Fig. 2.31. As shown in the chart, our algorithm could find the optimal solution when the $\mu$ is less than 4. For the networks with more complex structure ($\mu \geq 4$) also the algorithm obtained much better values than the genetic one.
In the second experiment, we ran our algorithm on larger networks ranging from 128 to 10000 nodes. The results are shown in Fig. 2.32. According to the results, by increasing the size of the network, the accuracy of the algorithm are slightly decreased. The problem perhaps is related to the fitness function and its limitation. However, we observed that by increasing the number of iterations the quality of the candidate solutions are increased which means that our algorithm can escape from the local optima.
In another experiment, we have collected some data from Twitter on 4th of Nov. 2015 at 3:00 PM. Based on our extracted data, we identified the following list which represents the 10 top trend concepts at that time in Canada.

- #PM23
- #NationalStressAwarenessDay
- Minister
- Jody Wilson-Raybould
- Catherine McKenna
- #KidsToWork
- #EveryEntrepreneur
- #BeyondMarketing
• Maryam Monsef
• Marc Garneau

We searched for the first ranked topic which is ’#PM23’ and extracted 1000 samples from the Tweets. After that, we made a social graph by linking the Screen name of users who mentioned this hashtag in their tweets. In addition, the related hashtags also linked to this tag. Overall, 1000 samples were collected and the network was shaped as shown in Fig. 2.33. The number of nodes in the graph are 1000 with 1515 edges and the cluster coefficient index of 0.243.
We test our algorithm on this social network and illustrated the results. As demonstrated in Fig. 2.34, the algorithm found 55 communities. The first four big communities are #PM23, #pm23, @liberal_party and #swearingIn. We evaluated our information and found that the #PM23 and #pm23 referred to the new Canadian federal cabinet (Prime Minister 23) which were the top trend topics in that day. @Liberal_party was the screen name that tweets about this hashtag and #SwearingIN referred to the swearing-in ceremony in Canada.

![Identified Communities in the social graph](image)

Figure 2.34: Identified Communities in the social graph


2.5.6 Conclusion

In this chapter, we have extensively reviewed the problem of community detection in social networks. We also introduced a novel multi-population cultural algorithm to deal with the problem. The core of the algorithm is belief space which determines the range of the feasible solutions and guides the search direction towards finding the optimal/near optimal solution during the search process.

The performance, runtime, scalability and accuracy of the proposed model have been examined thoroughly on real-life and synthetic networks. The synthetic networks were generated in variable sizes based on Newman and LFR benchmarks. The results show that the proposed algorithm can find the real communities with a high accuracy even when the graph has a very complex and dense structure. Meanwhile, in comparison with other well-known evolutionary approaches in the field, it achieved much better results in fewer evolution cycles. The functionality of our algorithm was also demonstrated on the real-life data extracted from Twitter.

In addition, a comprehensive study has been performed to shed light on the role of knowledge in the evolution process. The obtained results confirm that the extracted knowledge can significantly enhance the process in both levels of accuracy and processing time. In fact, the search space can be reduced dramatically by 80% as a result of using our approach. The runtime analysis also shows that adding the belief space has few impact on the overall complexity of the algorithm. Moreover, in compare to the genetic method, the algorithm has better performance in all the experiments.

On the other hand, we proposed two more structures to shape the belief space. A study has been carried out to compare the performance of these proposed structures. According to the results, the probability based matrix has the better performance in comparison to the other methods.
In summary, the major contributions of our proposed approach can be listed as follows:

- Introduce a unique method to define, extract, and represent Normative and Domain knowledge sources from a snapshot of the network to determine the range of the optimal solution.

- Introduce a novel data structure based on a probability matrix to store the normative knowledge.

- Introduce a method for utilizing the extracted knowledge to guide the search direction.
References


Chapter 3

Population Adaptation in Social Networks based on Knowledge Migration

Social networks can be analyzed from different aspects- micro and macro. If we assume that the main asset of each network is its population and the key difference between populations is their knowledge, then it is the knowledge that drives the evolution of any network.

In this chapter, the behavior and status of a network will be analyzed in a case where a population from one network migrates to another similar network and transfers its knowledge to it. In fact, we are going to find how a migrated population will adapt itself to a new environment with similar characteristics based on the knowledge that it has learned from the previous network and what is the role of this prior knowledge in its evolution. For this purpose, different scenarios are modeled by employing a cultural algorithm with various networks and populations on two different cases: a population with migrated knowledge and a population without it.
The results clearly show that when the changes in the structure of networks are less than 25\%, trained population can adapt itself with the new network very fast but when the difference is higher, in the best case they perform like a random population without any training.

As the case study of this research is community detection in social networks, the obtained results confirm that our proposed community detection algorithm can adapt itself very fast to the various snapshots of dynamic systems if the topological difference between two consecutive snapshots of the network is less than 15\%. It can lead to a remarkable reduction in the search time and space throughout the dynamic social network analysis.

3.1 Introduction

Social networks can be defined as collaborative, open and diverse environments which consist of associated entities. Due to their interactive nature, information and knowledge can be shared among all of their members which can be used to accelerate their evolution process. In a general view, a network consists of two essential layers- structure and content. These two layers are directly under the influence of each other and co-evolve themselves [18, 1, 5, 6]. The structure usually consists of members of the network and their relations. On the other hand, the content layer typically contains information which is transmitted between the network members. In some types of networks, members of the network just transmit data from one point to another, such as computer networks. In some other networks, members of the network not only transmit the information but also generate and produce the content. Consequently, the network member can be considered as the source of information. Examples could be found in networks such as social networks and the Web. In general terms, knowledge and information are the driving engines which guide the evolution of this type
of network.

These networks are not abstract, and they are in constant interactions with their surrounding environment [2]. Having knowledge about their behavior, characteristics and contents can reveal hidden patterns which can be useful to describe characteristics of the environment. Because of the importance and wide range of applications that it has, network analysis is at the center of attention of research centers around the world. In these networks, each member that we call social agent, has some sort of knowledge which comes from different sources including learning from the environments, learning from other members, etc. Meanwhile, all of these social agents taken together are called as population. They follow some unwritten rules and patterns that can be defined as culture. As these individuals interact with each other they transfer their own knowledge to others and update themselves with others’ knowledge; consequently, a network evolves very fast.

In this paper, the target is to find out how a population can perform in different environments when it has a prior knowledge attached to it. If a population knows how it can perform in certain scenarios based on this knowledge, how it will perform if we change the environment? To what level of similarity between two networks, migrated population can be adapted efficiently? We define this concept as an adaptation process. In other words, we are interested in figuring out how populations perform in networks which are related to some extent. What if a population with some knowledge is kept under a different environment? How comfortably will it adapt to this new environment? In addition to that, how will the adaptation vary depending upon how similar is the environment with the one this population is comfortable to?

To find answers to these questions, we choose the problem of community detection in social networks as a case study. As a framework, we have adapted an existing cultural algorithm [17] to find communities in social networks. This is an evolutionary method which works by extracting some knowledge from the structure of the networks.
We define four main scenarios to implement our idea. First is, where knowledge and population both are extracted and migrated to another similar network and the second one, where only the population is extracted and migrated, the third one is when just the knowledge is migrated to another network and the last one is if only the best individuals migrated to the target network. We test the performance of adaptation process for all scenarios in various level of similarity between networks by comparing their adaptation time.

This paper is organized as follows. In the next section, we will review some existing methods in this field. Our model will be proposed in the third section, and the experiments and the results will come after that in the fourth section. The last section will be the conclusion and future works.

3.2 Literature Review

In this section, we will review some existing work related to our research. As our framework is a cultural algorithm, we start by reviewing the architecture of cultural algorithms. After that, we review some research in the field of population migration and adaptation.

Cultural Algorithms are a branch of Evolutionary Algorithms and an extension to the genetic algorithms to find an approximate solution for complex problems. As shown in Fig. 3.1, it encompasses a population component, and in addition to Genetic Algorithms, a knowledge component which is also known as belief space. The belief space updates itself during the runtime of the algorithm and shares the knowledge among the individuals in the population. Knowledge is the beacon that guides the evolution of the population. The role of knowledge is to evolve the next generation with the help of extracted knowledge from the best individuals active in the previous generations [17, 3, 7, 14, 9].
Depending upon various domains of knowledge that a population can have in the cultural algorithms, a belief space is categorized in five different categories—Normative, Situational, Domain, Spatial, and Temporal. Normative knowledge provides standards that are used to define every individual’s behavior in the population. These standards are used as guidelines for mutational adjustments for individuals. In fact, normative knowledge is that stores the knowledge about the acceptable behavior of the agents. Situational knowledge deals with success and failure of events and takes care of the best solutions found in each iteration. Domain specific knowledge, which is similar to situational knowledge, imbues the knowledge about the domain the algorithm is applied to. Spatial knowledge stores the knowledge about the topography of the search space and finally, Temporal knowledge, which stores the history of the search space [17, 3, 14, 9].

The idea of cultural algorithms is inspired by the natural cultural evolution process. It assumes that by using knowledge, generations can evolve faster than normal biological evolution. Because of the comprehensive learning mechanism that it has, cultural algorithms can be used to analyze complex systems in both static and dy-
The dual-inheritance feature of Cultural Algorithms allows the system to both learn and adapt the best features of any population. A belief space stores the knowledge it gains during the runtime of a cultural algorithm. This updated belief space then influences the next population generation process. This next generated population, in return, updates the belief space with the help of best individuals. Furthermore, a fitness function is used to evaluate each individual’s performance.

Like most of the evolutionary algorithms, the cultural algorithm process starts by generating a random population. The quality of generated individuals is evaluated by the fitness function in the next step, and the group of them that have higher fitness values are selected to be candidates for updating the belief space. Based on the problem, some criteria may be set by the accept function which is used to give permission to some of the selected individuals to update the belief space. In the next step, the knowledge is extracted from the individuals and the update function revises the belief space based on them. After that, the new generation of the population is generated based on the rules which are defined in the influence function. The process continues until the predefined stop criteria is met by the algorithm.

To decrease the processing time and increase the efficiency in multi-agent systems recent work has been done on Multi-Population Cultural algorithms (MPCA) which are an extension to the Cultural Algorithms. As shown in Fig. 3.2, in MPCA, a population is divided into many sub-populations, and each one of them is assigned a global belief space common to all of them.
The authors in [8] claim that previous work has not been done to study the performance of MPCA in a case that individuals influence the evolution of other sub-populations. Therefore, they have proposed Transfer-Agent based MPCA. In their proposed model, unlike MPCA, best individuals are not used to update the global belief space. Instead, their positions itself are swapped with each other such that each subpopulation and a foreign individual are introduced to each other. According to their results, transferring individuals between different subpopulations that are having different knowledge, results in better performance in the cases where individuals from a subpopulation with more knowledge are transferred to subpopulations with less knowledge.

The authors in [11], proposed a new population adaptation technique based on a genetic algorithm to reduce the amount of required time to reach an optimal decision for a cognitive engine. They proposed a method that uses information from previous cognition cycle to seed the initial generation by utilizing the optimal decision of the previous cycle to bias the first generation. According to their results, this technique can enhance the required time to reach the solution by up to 480% in comparison to a standard random initialized Genetic Algorithm.

In [7], the authors proposed a new MPCA based on knowledge migration. As
shown in Fig. 3.3, they worked on an MPCA without a global belief space, where in this model, each population has own belief space. They proposed a mechanism for knowledge migration between these sub-belief spaces.

Figure 3.3: Knowledge migration on an MPCA without global belief space

Some research also has been carried out by using the evolutionary algorithm on dynamic environments [18, 1, 5, 6, 15]. The authors in [15], studied the performance of a cultural algorithm in dynamic environments and concluded that the CA is suitable to work on dynamic networks. To the best of our knowledge, there is no empirical research exist to study the potential outcome of migrating trained population for a special problem to another similar issue.

3.3 Problem Statement

As we mentioned before, the main objective of this research is to study the role of knowledge in population adaptation. The question is that, what is the role of prior
knowledge in the process of adaptation? Particularly, as shown in Fig. 3.4, we are interested in determining how adaptation process evolves in a case where a population with prior knowledge about an environment migrates to a new similar environment.

To achieve the goal, the problem of community detection in social networks has been chosen as a case study. We would like to analyze the results of adaptation process by migrating a population which is trained to solve the problem in one network into another network with a similar topology.

![Figure 3.4: Population adaptation process by knowledge migration](image)

The results of this research can be useful to clarify the role of knowledge in the evolution of dynamic networks. In addition, they can be used as a validation measurement for social network analysis and decision-making systems.

### 3.3.1 Community Detection in Social Networks

A social network can be defined as a graph, $G(V, E)$, where $V = \{v_1, v_2, \ldots, v_n\}$, is a set of nodes which represents the network population and $E = \{(v_i, v_j), \ldots\}, (v_i, v_j) \in V$, is a set of edges which can be interpreted as interactions between pair of agents in the population. Consequently, the graph with $n$ vertices can be formed based on
its n-by-n adjacency matrix. Let A denote the adjacency matrix of the unweighted network, an entry of A(i,j) is 0 if nodes i and j are not connected together and A(i,j) is 1 when there is a link between them.

On the other hand, social networks have some distinct characteristics. Having a high value of clustering coefficient value is one of them which indicates that the network consists of connected communities. The clustering coefficient can be calculated based on the number of links between neighbors of a node, and the node degree. If \( N_i \) denotes the number of links between neighbors of the node i, and \( D_i \) denotes the degree of the node i, the cluster coefficient of the node i, \( C_i \), can be computed as \([16]\):

\[
C_i = \frac{(2 \times N_i)}{(D_i \times (D_i - 1))} \tag{3.1}
\]

Community detection, therefore, is one of the fundamental parts of social network analysis, especially if the target is to analyze the structure of the network and its evolution. Community detection can be defined as finding groups of nodes in the network that have more links between each other than outside the group. In recent years, many methods have been proposed to solve the problem efficiently, from classic clustering algorithms to probabilistic models. However, in this paper we are adapting the knowledge-based evolutionary algorithm which has been proposed in [17] as our framework. In the following paragraphs, we briefly review its mechanism. As shown in Fig. 3.5, this algorithm has been defined on two levels — the population and the belief space.
Figure 3.5: Components of the proposed cultural algorithm in [17]

As the algorithm has been defined based on multi-population architecture, in the initial stage, individuals must be generated randomly to form the population. An individual in this algorithm is actually a random subset of the graph and is represented with the help of a unique locus-based adjacency representation [12]. The individual is structured as an array of nodes of the network, and its size is made equal to the number of nodes present in the network. Every cell of this array is direct mapping to its corresponding node in the graph. It means that cell #n in the array represents the nth node in the network. Each cell of this array is filled choosing a random node from the list of its direct neighbors in the network.

<table>
<thead>
<tr>
<th>Population Space:</th>
<th>Belief Space:</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Initial populations</td>
<td></td>
</tr>
<tr>
<td>2. Decoding</td>
<td></td>
</tr>
<tr>
<td>3. Fitness Evaluation</td>
<td></td>
</tr>
<tr>
<td>4. Selection</td>
<td></td>
</tr>
<tr>
<td>5. Update Belief Space</td>
<td></td>
</tr>
<tr>
<td>6. Check the stop criteria</td>
<td></td>
</tr>
<tr>
<td>If met goto 9, else goto 7</td>
<td></td>
</tr>
<tr>
<td>7. Generate new populations</td>
<td></td>
</tr>
<tr>
<td>8. Goto 2</td>
<td></td>
</tr>
<tr>
<td>9. Select the best Individual</td>
<td></td>
</tr>
<tr>
<td>1. Compare the fitness value of the selected candidates to the situational knowledge</td>
<td></td>
</tr>
<tr>
<td>2. Normative Knowledge is updated by the accepted individuals.</td>
<td></td>
</tr>
<tr>
<td>3. Update the Situational Knowledge.</td>
<td></td>
</tr>
</tbody>
</table>

![Graph adjacency matrix and random individuals](image)

Figure 3.6: A sample network and two random individuals

For example, according to the graph adjacency matrix in Fig. 3.6, node #1 is connected to node #2 and #3. Node #2 is connected to nodes #1, #3, and #4. In addition, node #3 is connected to node #1 and #2. Hence, the Random Individual...
1 and 2 have been generated based on the described mechanism in the previous paragraph. As shown in Fig. 3.6, by illustrating the individual the probable communities that form the graph can be extracted. In fact, as a result of decoding an individual, all nodes in the array which are linked together are considered as a separate sub-set of a network which can be interpreted as a community. Consequently, each individual splits the network into some random communities and assigns the network’s nodes randomly to them.

After this step, these individuals will be evaluated with the help of fitness function. The algorithm used the concept of community score which has been proposed in [13] as the fitness function where the higher value of the community score can be interpreted as the better-formed community. When sorted, a group of individuals with better fitness values is selected to be a candidate to update the belief space [17].

The belief space is the core of the algorithm and consists of two sources of knowledge which have been extracted from the accepted individuals. The first one is the situational knowledge which stores the average fitness value of the accepted individuals and the second one is the normative knowledge. The situational knowledge has been used to filter the selected candidates by accepting just those candidates that their fitness values are higher or equal to the situational value. The accepted individuals can update the normative knowledge which is represented by an n-by-n matrix, where n is the number of the nodes in the network. Each row of the matrix is filled with the relative frequency of links between each pair of nodes among accepted individuals [17].

For example, assume the empty 6-by6 matrix and three individuals have been accepted to update it. Let $\text{Individual}_1 = [2, 3, 1, 5, 6, 5]$, $\text{Individual}_2 = [2, 3, 2, 1, 6, 4]$ and $\text{Individual}_3 = [3, 3, 1, 5, 6, 4]$. The result of the update process is a matrix which has been illustrated in Fig. 3.6. In this example, node #2 has been seen in the first position of these individuals, two times out of three. Therefore, the value of entry
(1, 2) of the matrix is set to 0.666. Following the same instructions, the value of entry (1, 3) is 0.333.

Normative knowledge can be interpreted as the weighted adjacency matrix such that the weight shows the level of the relation between each pair of the neighbor nodes among the best individuals. Therefore, it can be useful to limit the search space needed to generate the next generation of individuals. In fact, instead of searching all the neighbors, the random search will be limited just to those neighbors that have been seen together in the best-accepted individuals and the weight determines the chance of selection in the search process.

![Normative knowledge](image)

Figure 3.7: Normative knowledge extracted from the three accepted individuals

After updating the belief space, like as other evolutionary algorithms, if the stop condition has not been met the algorithm starts a loop and generates new populations in every single iteration such that majority of individuals are generated based on this matrix and a minority of them based on the original adjacency matrix [17]. The fitness evaluation, selection, and belief space update process continue until the condition met.

### 3.4 Population Adaptation Based on Knowledge Migration

To study the role of prior knowledge in the adaptation process we propose the following four different scenarios:
- Transferring the adapted populations and their knowledge from a random social network to another network,

- Transferring knowledge obtained during adaptation process to another population which deals with a different network,

- Transferring the adapted population from one network to another,

- Transferring the best individuals of the adapted population to another network.

Before describing the scenarios, the training environment must be defined. To train populations, as we mentioned in the previous section, the community detection algorithm proposed in [17] has been chosen to find the network communities. The input of it, is a random social network graph, and as the outputs, normative knowledge matrix, best individuals and the number of executed iterations to find the correct communities will be extracted. This algorithm starts with generating random populations and stops when the predefined number of iterations has been reached. At first, the belief space is empty, so an individual is generated completely random based on the network adjacency matrix by using the mechanism which has been discussed in the last section. Fig. 3.8 illustrates this process.
To implement our scenarios, we then need to change the network. The new network will be generated based on the current network such that, a particular number of random edges will be added to it. The number of edges that must be added will be determined according to the degree of non-similarity that we want to have between two networks.

For example, let a base network has 100 nodes and 2000 edges, assume the degree of non-similarity=5%. Therefore, the new graph must have the same number of nodes and 2100 edges which means that 100 edges must be added randomly to the network. The only condition that must be set here is that the distribution of these edges must be such that the new graph still remains in the category of social networks and keeps its characteristics [16].

To analyze the performance of the adaptation with or without the prior knowledge, we compare the number of iterations which are needed to find the correct communities.
in the networks (without prior knowledge) with the one obtained from each of the scenarios. For example, let the algorithm needs 10 iterations to find the communities in a training environment. If each of the scenarios can find the correct communities in another network by adapting the knowledge obtained from the training in fewer iterations, it shows the role of knowledge in improving the adaptation process.

Another goal is to change the degree of similarity. We would like to estimate what the role of prior knowledge is in the adaptation process in different degree of similarity between two networks.

### 3.4.1 Scenario 1: Population and knowledge migration

In this scenario, the trained population, and its knowledge will be imported by the algorithm. As shown in Fig. 3.9, instead of generating random populations in the initial phase, the trained population will be used as the initial populations. In addition, the normative knowledge obtained from these populations will be exchanged with the empty normative knowledge matrix. The goal of this scenario is that to know, how the population that has prior knowledge about a similar network can adapt themselves to the new network.
3.4.2 Scenario 2: Knowledge migration

The goal of the second scenario is to analyze the role of knowledge alone in the evolution of the populations. Considering, knowledge about a structure of a network transfers to new populations that do not have any prior knowledge about their network’s structure. As shown in Fig. 3.10, initial individuals will be generated based on the imported knowledge instead of generating them randomly.

The difference between the first and the second scenario is that here the trained populations do not migrate physically and the initial individuals are generated based on their obtained knowledge during the adaptation process.
3.4.3 Scenario 3: Population migration

As shown in Fig. 3.11, in this scenario, the trained population will be migrated to the new environment. The assumption is that the knowledge is an integral part of the populations therefore by migrating them, their knowledge automatically will be transferred to the new network.

Accordingly, the algorithm will import the trained population in the initial phase instead of generating random populations.
3.4.4 Scenario 4: Migration of the best individuals

In the final scenario, instead of transferring the whole of the trained populations, just the selected individuals (best individuals after ranking process) which are actually the elite group of populations will be migrated to the new network. Therefore, they will join the randomly generated individuals in the new algorithm. Fig. 3.12 illustrates this scenario.
3.5 Evaluation

To test our scenarios, we have implemented the algorithm proposed in [17] which has been described before. The size of each population has been set to 200 and the selection rate has been set to 20%. For the fitness function, we used the concept of community score which has been proposed in [13] and used in [17, 13]. The number of iterations was also set on 50. The test has been carried out on a 3.2 GHz core i5 computer with 12 GB RAM. The code has been written in MATLAB R2014r.

To generate the social network graph, 10 base networks have been generated according to the Girvan-Newman benchmark in [4] which is one of the commonly acceptable benchmarks in the field of community detection. Each network has 128 nodes which are grouped in four communities with the size of 32 nodes each. Each node has the average degree of 16, such that $Z_{in} + Z_{out} = 16$, where $Z_{in}$ denotes the number edges that the node has inside the community and $Z_{out}$ denotes the number of links that node has with the other nodes outside its community. By increasing the
value of $Z_{out}$, the complexity of the graph will increase.

In our experiments, the average degree of $Z_{out}$ was set to 5 and the rate of success in finding the correct communities were 100%. To measure the accuracy of the algorithm the Normalized Mutual Information (NMI) [17, 13, 4] value has been used which computes the similarity level between the actual communities and the detected ones. Based on each of the base networks, we generated 50 other networks by using 5 different levels of non-similarity, 5%, 10%, 15%, 20%, and 25% (10 networks for each degree). Therefore, 500 networks have been generated in total. In the following paragraphs, we describe the results obtained in each scenario separately.

### 3.5.1 Scenario 1

In this scenario, we first ran the community detection algorithm (algorithm without prior knowledge) on the base networks and extracted the normative knowledge matrix and the final generated populations. Then we exported them to the algorithm to implement the first scenario. In the next step, this algorithm (first scenario) and the original community detection algorithm ran 10 times for each of the 50 networks. The average numbers of iterations obtained by both algorithms to find the correct communities for each of these networks have been illustrated in Fig. 3.13.

As shown in Fig. 3.13, the community detection algorithm without prior knowledge was needed almost the fix number of iterations in all networks to find the communities. On the other hand, when the degree of non-similarity was 5% (means that there is 95% similarity between the structure of two networks), the algorithm proposed for the first scenario (with the trained populations and prior knowledge) could find the correct communities in the first attempt in all 10 networks. It can be interpreted as the population could adapt themselves very fast and in just one attempt.

By increasing the degree to 10% and 15%, the algorithm still could find the correct communities in almost half or fewer iterations in comparison to the algorithm without
prior knowledge. However, it was observed that, by increasing the non-similarity degree, the role of knowledge in the adaptation process would be vague. Finally, when the degree was set to 25%, we observed that not only the knowledge was less useful but also in some experiments it made more iterations to reach the goal.

Figure 3.13: The results obtained from the first scenario

3.5.2 Scenario 2

Similar to the first scenario, we started running the algorithm on the base networks and extracting the normative knowledge matrix. Then we exported it to the algorithm according to the description of the second scenario. We, then, ran both the algorithms 10 times for each of the 50 networks similar to the first scenario. As shown in Fig. 3.14, akin to the first scenario’s experiments, the average number of iterations needed to find the correct communities in the algorithm without prior knowledge is almost fixed. However, with 5% degree of non-similarity, the algorithm which has been
defined based on the second scenario, found the correct communities in the first run. Moreover, for the degree of 10% and 15%, the average performance of this algorithm was almost similar to the first scenario when the degree of non-similarity was 10% and 15%.

However, for the degree of 20%, the performance of this algorithm was slightly better than the algorithm without prior knowledge. Finally, similar to the first scenario, at the degree of 25%, we observed that knowledge was either not useful at all or even for some experiments increased the number of iteration to reach the desired state. Comprehensively, it seems that transferring knowledge with the trained populations had better performance in comparison to transferring just the obtained knowledge.

Figure 3.14: The average values of the results obtained from the second scenario
3.5.3 Scenario 3

According to the third scenario, the trained populations were imported by the algorithm as the initial populations. As shown in Fig. 3.15, the results obtained from this scenario were almost similar to the results of the first scenario. We believe that the role of the population with prior knowledge is stronger than the role of knowledge by itself.

Regarding the fourth scenario, we extracted the best-selected individuals (80 individuals) and exported them to the populations. The rest of individuals (320 individuals) has been generated randomly based on the graph adjacency matrix.

Figure 3.15: The average values of the results obtained from the third scenario
3.5.4 Scenario 4

As shown in Fig. 3.16, this scenario has better performance in comparison to the third scenario; however, its performance is almost similar to the other scenarios when the degree is above 20%.

![Bar chart showing the average values of the results obtained from the fourth scenario](image)

**Figure 3.16**: The average values of the results obtained from the fourth scenario

The results achieved from the experiments clearly show that the knowledge plays a significant role in the adaptation process especially when the level of difference between the training and target network is less than 25%. It means that prior knowledge can dramatically increase the speed and quality of the adaptation. However, the interesting observation obtained from this study indicates that the prior knowledge is not only helpful when the difference between the training network and the target network is more than 20% but also sometimes can increase the time needed for the adaptation process. Another interesting fact is that migration of trained individuals
has a better impact on the adoption process rather than transferring the obtained knowledge.

3.6 Conclusion and Future Work

In recent years, a lot of work has been carried out on social network analysis and population migration, but most of them have not considered the role of knowledge migration in the population adaptation process. In this research we proposed to analyze the role of knowledge migration in population adaptation, taking community detection in social networks as our case study. We have shown how knowledge, associated with any population, can play a significant role in its adaptation process in social networks with the help of multi-population cultural algorithms. To study the role of prior knowledge, we have proposed the following four different scenarios for transferring knowledge and the trained population from one network to another network.

- Transferring the adapted populations and their knowledge from a random social network to another network,

- Transferring knowledge obtained during adaptation process to another population which deals with a different network,

- Transferring the adapted population from one network to another,

- Transferring the best individuals of the adapted population to another network.

According to the results, we can observe that if two networks are similar to some extent, knowledge can play a very crucial role in the population adaptation process. According to our results, if the degree of non-similarity is less than 25% the influence of knowledge on population adaptation is very significant, but if it is more than 25%
not only its impact will be minimal, but also it may act as an obstacle and make the adaptation process complex.

Another main observation of this research is that the first and the third scenarios had better performances in comparison with the other scenarios. It can be interpreted that, transferring just a source of knowledge from one network to another by itself has less impact on the adaptation process in contrast to transferring the trained individuals.

The significance of this contribution in dynamic social networks is that it allows us to use the obtained knowledge from a previous analysis, stored in the belief space, to identify new communities by eliminating the need for a new search, if the similarity of two consecutive network snapshots is within 85%. This method can be generalized to accelerate the search performance in complex and dynamic social networks.

In the future, we would like to extend our work on the real world problems including single and multiplex networks and implement it to more experimental environments. Our target is to add a variety of complex scenarios to the framework and evaluate the outcome. Studying the role of different types of knowledge in the adaptation process, particularly domain knowledge would be one extension to our future work.
References


Chapter 4

Link Prediction in Social Networks

Social networks have a dynamic nature, so their structures frequently change over time. In this chapter, we propose a new community-oriented knowledge-based evolutionary method to predict the state of a network in the near future by extracting knowledge from its current structure. This method is based on the fact that social networks consist of interconnected communities and their members tend to join these communities. By observing the current state of a given network, the method calculates the probability of a relationship between each pair of individuals who are not directly connected to each other and estimate the chance of being linked in the next time slot.

A unique mapping function and a novel computational model based on the weighted graph have been defined for the estimation process. We have tested and compared the method on various synthetic networks and real datasets. Results show that our method can predict the next state of a network with a notably high rate of accuracy.
4.1 Introduction

People use social networks to interact with others. Regardless of the content, these interactions can reveal valuable information about real societies and individuals. This information can be useful to identify the structure and topology of these networks, which makes it possible to track their evolutions and predict the next state. Naturally, these networks are extremely dynamic, and their rate of evolution is very high. Consequently, their structure changes frequently. Since these networks reflect real life events, having knowledge about their next state can be applied to various domains such as recommendation systems, decision making, marketing and risk analysis [11, 5, 32, 7, 16, 18].

For example, in online social networks, it can be used to recommend new friends or products to the users [14, 2]. Meanwhile, in social media and the stock market, it acts as a powerful tool to predict the upcoming trends, events and behaviors [34, 39, 10, 22, 25]. In professional and academic networks it is helpful to find possible teams of experts for particular tasks [36, 31, 37]. In political science, predicting the outcome of elections or political decisions are challenging tasks which can become possible with the help of this knowledge [6, 13]. In crime networks, it can be seen as a tool which can shed light on the criminal system analysis [3]. In health science, predicting disease and its spread and outbreak are another applications of this important topic [12, 17, 26, 34].

In the field of social network analysis, this problem is known as Link Prediction, which can be defined as estimating the likelihood of a connection between two disconnected entities in a network in the near future [7, 16, 18]. The main idea behind this problem is, the future state of a network is not random and has a dependency on the current state. Therefore, the target is to find the level of dependency and the main factors affecting it.

Social Networks as a subset of complex networks have some particular charac-
teristics such as power-law distribution, the high value of cluster coefficient, and homophily phenomenon. The power-law distribution in social networks means that there exist relatively few nodes in the network with a high degree of connectivity which are called hubs and many nodes with low degree [1]. Identifying the hubs is an important task in analyzing the evolution of social networks.

Cluster coefficient measures another important characteristic of a network which can be interpreted as the tendency of nodes to cluster together. It is defined as:

\[
C = \frac{(3 \times \#\text{triangles})}{(\#\text{connected triplets in the network})}
\]  

(4.1)

Having a high level of cluster coefficient in the network indicates the strong tendency of users to join communities. Meanwhile, regarding the homophily phenomenon, users are willing to join the communities through their circle of friends. Accordingly, in this paper we propose a knowledge-based community-oriented evolutionary framework based on these properties to estimate the state of a network in the near future just by having one snapshot of the network.

Our proposed model is defined based on the similarity approach with two main assumptions. The first is that an individual in a network tends to join a community. The second is that, each individual joins a community through their friends. Hence, the similarity measurement here is defined as having a common community. For example, if a person in a network has 6 friends and 5 of them are members of a community with 30 people, the probability of a friendship between this person and members of the community in the near future is higher than other cases, and the level of this similarity can be estimated approximately.

To estimate this likelihood, a knowledge-based structure which is called belief space has been adapted from the evolutionary cultural algorithm which has been proposed for the community detection problem in [41]. Cultural algorithms are a
specific type of evolutionary algorithm that use knowledge to enhance the search process to find near optimal solutions for a problem [41, 33]. As shown in Fig. 4.1, a cultural algorithm consists of Population and Belief spaces. In fact, the population space is a set of feasible solutions for a given problem which is the community detection problem in this case. The belief space is a knowledge-based structure which guides the population generation process in each iteration, and is evolved by extracting information from the population space [41, 33].

In other words, as illustrated in Fig. 4.1, in each iteration a set of candidate solutions for a given problem are generated in the population space. A fitness function evaluates their performances, and the best group of them are selected. Different types of knowledge will be extracted from this group in the belief space to shape the range of the target solution which is lead to the search space reduction. The new set of solutions is generated based on the obtained range in the belief space.

![Figure 4.1: A cultural algorithm process](image)

In this paper, by focusing on the belief space as a great source of knowledge, we propose an algorithm to determine the level of dependency between each pair of users and estimate their tendency to communicate with each other. The main structure of our proposed algorithm is a directed weighted graph which is generated from the belief space data and demonstrates levels of relationships between all neighbor nodes. For
predictions, a mathematical model is proposed to estimate the likelihood of having a relationship between each unconnected pair of nodes. This model has been defined based on two main concepts, the number of paths between each unconnected pair of nodes and the length of these paths. Having more paths and shorter distances implies a higher chance of connection in the next time slot. Finally, our algorithm calculates the probability of a relation between pairs of nodes which are not directly connected and ranks them.

The main contributions of this research can be summarized into the following categories:

- Introducing a novel concept of observing the quality of links between pairs of nodes. By observing a snapshot of the social graph, an evolutionary approach has been used to determine the level of dependency between each connected pairs of nodes and make a weighted network.

- Proposing a mathematical method to extract information from the structure of a given network as a similarity index.

- Proposing a unique knowledge-based computational model for analyzing the evolution of social systems and estimating the state of the network in the next time slot.

The rest of the paper is organized as follows: In the next section, the problem definition and related works will be reviewed. The detailed description of our model has been presented in section 4.3. After that, in section 4.4 the evaluation of the model will be studied and discussed. Conclusions are presented in the last section.
4.2 Problem Definition and Related Works

If a network maps to a graph, $G(V,E)$, where $V$ is a fixed number of nodes and $E$ represents links between each pair of nodes, an edge is defined as $e = (u,v) \in E$, where $u,v \in V$, at a particular timeslot ($t$). As shown in Fig. 4.2, predicting a state of the graph at time $t + 1$ by having a snapshot of it at time $t$, is defined as the Link Prediction Problem in social networks. In other words, given a network $G_t$ at time $t$, the output of a link prediction algorithm will be a list of edges which are not in $G_t$ and have high probability of appearing in $G_{t+1}$ [7, 16, 18].

![Figure 4.2: Predicting the state of a network at time $t + 1$, given a snapshot of it at time $t$](image)

The problem is closely related to another issue in the field which is defined as finding missing links in the network. The assumption is that the observed network might not thoroughly capture all the links. Therefore, they may exist some hidden links between the users which are not directly visible. Predicting these missing links is a challenging task which can lead to better understanding of the current state of the network [28]. As illustrated in Fig. 4.3.a, the actual network has five nodes with nine edges. However, the observed network which is shown in Fig. 4.3.b just captured seven edges. In this issue, the main task is to identify the missing links demonstrated in Fig. 4.3.c, by estimating the level of interdependence between all the non-observed links.
In recent years, the problem of link prediction in social networks has received extensive attention from researchers due to the wide range of its applications and fast growth of online social networks. Having access to more data about the network and its users in both structure and content levels has had a significant impact on these research works. The current works can be basically categorized into two categories based on their approaches which are similarity-based methods and probabilistic models.

The idea behind the similarity-based approach can be expressed as having a high number of similarities among pairs of users increases their chance of making a link in the near future [16, 18]. Therefore, in this approach, the algorithms calculate the level of similarity between each pair of nodes $x$ and $y$ and assign a score to them. After ranking them, they select the pairs which have higher scores as they have more likelihood to be linked in the next timestep.

The similarity between two nodes can be measured based on the node’s information and attributes or the network’s structure. Profile information, the number of publications or tweets, topics of interest and users’ transactions and activities are some examples which can be used as node’s attributes to measure the similarity [5, 4, 15, 38]. In [4], the authors proposed a tree-based model to estimate the similarity between users based on the semantic dependencies of their profiles’ keywords.

Figure 4.3: Inferring the missing links in a graph with 5 nodes
To find the semantic similarities between the keywords, they proposed a forest model which consists of multiple trees. In this model, each keyword is mapped to a node and will be linked to another node if there is a relation between the corresponding keywords. The hierarchical architecture of the model is used to calculate the distance between the pairs of nodes. They used WordNet to obtain the list of related keywords as the core of their model. After that, they defined two concepts of Weak and Strong similarities to estimate the level of similarity between users.

In another work [15], the authors proposed a model to combine the network structure and node attributes to deal with the problem. They extended the existing model which they called it Social-Attribute Network (SAN) and employed various link prediction algorithms. Their result demonstrated that integrating the both concepts can improve the performance and lead to the more accurate prediction.

There exist many research works which are concentrated on node’s attributes to solve the link prediction problem [5, 18, 16, 4, 15, 38]. However, due to lack of access to the users’ information, extracting the attributes is a big challenge in this approach. Meanwhile, as the main focus of this paper is using structural information to tackle the problem, the topology-based methods will be reviewed in the next paragraphs.

The unsupervised methods for the link prediction mainly rely on the structural similarity of the nodes [11, 16, 18]. Many similarity indexes have been proposed such as the Jaccard similarity coefficient, Katz, Common Neighbors, Leicht-Holme-Newman. Some of these indexes calculate the similarity between a pair of nodes based on their number of common neighbors. Some others are global and compute the similarity based on the existing paths between two nodes [16, 18, 38].

The Common Neighbors index counts the number of shared neighbors of nodes $x$ and $y$. It is formally define as:

$$ C(x, y) = |\Gamma(x) \cap \Gamma(y)| $$

(4.2)
where $\Gamma(x)$ and $\Gamma(y)$ are the lists of neighbors of nodes $x$ and $y$, respectively. The assumption is that the likelihood of a friendship between two unconnected nodes in a network become higher by increasing the number of their shared friends.

The Jaccard similarity coefficient is another important neighbor-based index which measures the number of shared neighbors between two nodes over number of their all unique neighbors. The index is defined as:

\[
J(x, y) = \frac{|\Gamma(x) \cap \Gamma(y)|}{|\Gamma(x) \cup \Gamma(y)|} \quad (4.3)
\]

This index gives a higher similarity score to the nodes which have more common neighbors and less private ones.

Leicht-Holme-Newman is also an neighbor-based index which determines the similarity by calculating the number of common neighbors between nodes $x$ and $y$ relative to the product of their degrees. It assigns a high similarity score to a pair of nodes that have many common neighbors compared to the expected number of such neighbors [21].

\[
L(x, y) = \frac{|\Gamma(x) \cap \Gamma(y)|}{d(x)d(y)} \quad (4.4)
\]

where $d(x)$ and $d(y)$ are the degrees of nodes $x$ and $y$, respectively.

The Resource Allocation Index is another neighbor-based index which performs well on real networks. Consider the situation where node $x$ sends some resources to node $y$ through its mutual neighbors. The similarity between $x$ and $y$ is then defined as the amount of resources received by node $y$. The index is formulated as:

\[
RA(x, y) = \sum_{z \in \Gamma(x) \cap \Gamma(y)} \frac{1}{|\Gamma(z)|} \quad (4.5)
\]

Academic-Adar Coefficient (AA) is a neighbor-based index which was initially proposed to estimate the level of similarity of a pair of web pages. The index assigns
more weight to the rare features in comparison to the common ones. The assumption is that the rare items are more valuable than the ordinary ones. Consequently, more weight is assigned to the common nodes with lower degrees. The metric is defined as:

$$AA(x, y) = \sum_{z \in \Gamma(x) \cap \Gamma(y)} \frac{1}{\log |\Gamma(z)|}$$ \hspace{1cm} (4.6)

Preferential Attachment (PA) is another neighbor-based index which gives a higher similarity score to a pair nodes that have more degrees. The assumption is that the likelihood of a connection between high-degree nodes is greater than low-degree ones. The index has the lowest computational complexity and is defined as:

$$PA(x, y) = |\Gamma(x)| \cdot |\Gamma(y)|$$ \hspace{1cm} (4.7)

In [24, 42], the authors have comprehensively compared some of the indexes on many real networks. According to the results, the resource allocation index has better performance in comparison to the other metrics. However, despite the existence of many neighbor-based indexes, none of them can be used as an absolute universal solution for all types of networks. However, the main challenge in this approach is the issue of scalability. Therefore, a suitable metric must be chosen based on the characteristics of a given social networks [16, 18].

As mentioned before, another approach to calculate the similarity relies on the role of existing paths between a pair of nodes. Consequently, global structural information is required in this approach. Katz and Friend Link are examples of this approach which are reviewed in the following paragraphs [16, 18, 38, 29].

Katz is an important path-based index which estimates the similarity based on the number of paths between two nodes and their distances. The idea is that the more paths between two nodes and the shorter distance imply the higher probability of connection between them. The index counts all paths between a pair of nodes and
assigns more weights to the shorter paths. It is defined as:

$$Katz(x, y) = \sum_{l=1}^{\infty} \beta^l |path^l_{x,y}|$$

(4.8)

where $\beta$ is a parameter to regulate the path’s weights and $path^l_{x,y}$ is the set of all paths between nodes x and y with the length of l.

FriendLink is a path-based index which measures the similarity between two nodes x and y, by counting the number of paths of varying length between them. The assumption is that all the paths regardless of their lengths must be considered in the estimation process. The index is defined as:

$$FL(x, y) = \sum_{i=2}^{l} \frac{1}{(i-1)} \frac{|path^i_{x,y}|}{\prod_{j=2}^{i} (n-j)}$$

(4.9)

Where $n$ is the number of nodes in the network, $l$ is the maximum length of a path, $1/(i-1)$ is the parameter to regulate the paths’ weights based on their length, and $path^i_{x,y}$ is the number of all length-i paths between the nodes.

In addition, maximum likelihood and probabilistic models approaches which are supervised methods are also used to solve the link prediction problem. In [8], the authors proposed a maximum likelihood method for prediction of missing links in networks. Their assumption is that the network has a hierarchical structure. They proposed a method to identify the hierarchical fabric of the network to make a dendrogram called hierarchical random graph. They introduced a method to assign a probability of connection between a pair of nodes in the dendrogram by using sampling. Their ultimate target is to find the dendrogram that best fits the observed network and predict the missing links in such network. Compare to simple similarity indexes, these models can predict the links more accurately, but by increasing the size of the network ($|\text{network}| > 10^4$), they become impractical because of their time complexity [16, 18, 38].
Evolutionary and swarm-based approaches are also used to solve the problem that have been proposed in recent years [5, 32, 7, 35]. In [5], the authors have used the Covariance Matrix Adaption Evolutionary Strategy (CMA-ES) to optimize the prediction accuracy. They suggested a linear model for combining common neighbor’s similarity indexes and nodes specific information by assigning a weight to each index. They mentioned that the limitation of their work is that the optimal model may not be linear. On the other hand, the model does not need any prior information about the network which is the main advantage of this model. In addition, their proposed model can be applied to various networks regardless of the types.

In [7], the authors proposed an algorithm based on ant colony optimization to solve the problem. The model relies on the subgraph evolution. Random walk strategy has been implemented in their algorithm to select paths. The probability is assigned to an edge to help an artificial ant select a better edge. In each iteration, the quality of the paths is evaluated to update the probabilities for the next iterations. Finally, the path with higher quality is selected as a link which has more likelihood to appear. The optimization part consists of three main steps. First, the probability of traversal of an ant from node x to node y in the graph must be computed. The second is releasing the pheromone on the traversed path from the home to the food source (destination). The final step is pheromone evaporation which is essential to find the shortest path in the network.

In [9], the authors proposed a prediction model to estimate the missing links in the network using the obtained information from the community structure. The model, extract the community structures and compute the number of times that a pair of nodes appeared together a community under different resolutions. The likelihood of a connection between them then will be calculated.

Since the future has not come yet, validation and verification of the link prediction algorithms is a challenging task. Therefore, to test and evaluate the performance
of the algorithms two main methods exist. The first one is to use datasets with timestamps which have been obtained from real networks. Consequently, the output of a proposed algorithm is compared with the structure of the network in the next timestep which can be extracted from the datasets. Another option is to divide a network randomly into two subsets, the training set, $E_T$, and the probe set, $E_P$. As the result, $E_T \cup E_P = E$ (the set of the network’s edges) and $E_T \cap E_P = \emptyset$. Here, $E_T$ can be considered as the observed known interactions and $E_P$ as the set of links that must be predicted for testing. In the prediction process, information from $E_T$ must not be used.

To measure the accuracy of the algorithms, two main methods are commonly used, the Area Under the Receiver Operating Characteristic Curve (AUC) and Precision [16, 18].

AUC is defined as:

$$AUC = \frac{(n' + 0.5n'')}{n}$$

(4.10)

where $n$ is the number of independent comparisons and $n'$ denotes the number of times a randomly chosen missing link (a link in $E_P$) had a higher score than a randomly chosen nonexistent link (a link in $U - E$, where $U$ denotes the universal set containing all possible links, of which there are $|V|(|V| - 1)/2$, with $|V|$ the number of nodes in the network). Furthermore, $n''$ denotes the number of times that their score is the same [16, 18]. In fact, AUC is used to estimate the probability that a randomly chosen missing link obtains more score than a randomly chosen nonexistent link. In the implementation phase, to reduce the time complexity, at each time a random pair of a missing link and a nonexistent link is selected for the comparison, instead of comparing all possible states.

As an instance, Fig. 4.3.b can be interpreted as $E_T$ and Fig. 4.3.c as $E_P$. Therefore, pairs of (1,3) and (3,5) are selected as probe links. Provided $E_T$, a link prediction algorithm calculates the likelihood of a relation between the pair of all non-observed edges
in the graph \(((1,3),(3,5)\) and \((1,5)\)). Assume that the assigned scores are \(S(1,3)=0.5\), \(S(3,5)=0.6\), and \(S(1,5)=0.5\). The obtained scores must be compared with each other to measure the accuracy of the results. Hence, as there are just one nonexistent link and two probe links, two possible comparisons are considered: \((S(1,3)=S(1,5))\) and \((S(3,5)>S(1,5))\). Accordingly, the AUC value is equal to \((1 \times 1 + 1 \times 0.5)/2 = 75\%\).

Given the ranked non-observed links, Precision is defined as:

\[
\text{Precision} = \frac{\# \text{ relevant items selected}}{\# \text{ items selected}}
\] (4.11)

In the case that the top-\(L\) links from the predicted links are chosen, if \(L_r\) denotes the number of these links which are in \(E^p\), then Precision can be defined as \(L_r/L\) \([16, 18]\).

### 4.3 Proposed Evolutionary Model

As we mentioned before, community is the core of our model. Thus, in our model we adapt outputs of the evolutionary cultural algorithm which has been proposed to detect communities on social networks in \([41]\). This algorithm is reviewed briefly in section 4.3.1. While the output of this algorithm is the list of communities, the focus of this research is on the belief space. This belief space can be interpreted as a probability matrix which estimates the quality of relationships between each pair of nodes in the network which are directly connected together. Using this belief space which is updated by the extracted information from populations in each iteration, the cultural algorithm limits the search space and enhances the individual evolutions. In our model, we propose using this knowledge repository as a source of information. Our hypothesis is that, this belief space which has designed to capture the relations among the nodes in order to assign them to the true communities, can be used for estimating the next state of the network.
As shown in Fig. 4.4, the belief space will be mapped to a directed weighted graph. The weights indicate the level of dependency between each connected pair of nodes. After that, we propose a method to estimate the likelihood of relationships between pairs of unlinked nodes in the graph. Ranking them will be the last process of this model.

**Figure 4.4: Components of the proposed model**

### 4.3.1 Making the Weighted Graph

In this part, we briefly describe the mentioned community detection algorithm [41]. In this algorithm, an individual can be seen as a feasible solution for the community detection problem. An individual therefore is represented based on a particular locus-based adjacency method [30] stored in an array structure. The length of this array is equal to the number of nodes in the graph. Each cell of this array is addressed from 1 to $n$ (the length of the array) which determines a node in the graph with the same number. E.g., cell #10 corresponds to node #10. The value of each cell #i, is an address of a node which is randomly chosen from the list of neighbors of the node #i.
For example, as shown in Fig. 4.5, if a network has seven nodes, one sample individual can be defined as an array of nodes, shown in Fig. 4.6. As illustrated in Fig. 4.7, this random individual represents a sub-graph with two communities (nodes #1, 5, 6, and 7 in one community and nodes #2, 3, and 4 in another).

Figure 4.5: A sample network

![Figure 4.5: A sample network](image)

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>3</td>
<td>2</td>
<td>2</td>
<td>7</td>
<td>7</td>
<td>6</td>
<td></td>
</tr>
</tbody>
</table>

Figure 4.6: A random individual

![Figure 4.6: A random individual](image)

Figure 4.7: Illustration of the individual in Fig. 4.6 - shows two separate communities (1,5,6,7) and (2,3,4)

![Figure 4.7: Illustration of the individual in Fig. 4.6 - shows two separate communities (1,5,6,7) and (2,3,4)](image)

As mentioned before and presented in Fig. 4.1, in each iteration, a specific number of individuals are generated by the algorithm (to make a population) according to the rules which are set in the belief space. The quality of these individuals is evaluated
using a fitness function which assigns a score to each individual. Consequently, they can be compared with each other. After sorting them, a group of them that have better fitness values are selected to change the belief space. However, to be eligible to update the belief space, they must meet some other conditions.

To update the belief space, each cell of these individuals adds its value to the \( n \) by \( n \) belief space matrix, where \( n \) is the number of nodes. The algorithm will calculate the relative frequency of these values and store them in the matrix as shown in Fig. 4.8. A selected individual denoted by \( SI_i(j) \), \( 1 \leq i \leq k, 1 \leq j \leq n \) where \( k = |\text{selected individuals}| \), is an array of adjacency links consists of \( n \) cells. As mentioned before, it represents a subgraph of a given network which has a high community score. In the belief space matrix, \( R_{x,y} \), \( 1 \leq x \leq n, 1 \leq y \leq n \), represents the number of times that node \( x \) linked to node \( y \). In fact, the matrix demonstrates the relative frequency of the times that two neighbors appeared together in the same community according to the obtained knowledge from the best-selected individuals. With this method, the belief space can be considered as an alternative adjacency matrix for the graph, because it is a weighted sub-graph of the main network that shows the level of dependency between nodes according to the community index. Hence, the next generation of individuals are generated using this adjacency matrix.
Fig. 4.9 shows an example for updating the belief space. Five individuals have been selected to update the belief space of the same network shown in Fig. 4.5. If the matrix had been empty before, then it is populated by the relative frequency of nodes and their neighbors. For example, node #5 was linked to node #1, 20% of times (once out of 5 times). If we illustrate this belief space, a directed weighted graph will be the result as shown in Fig. 4.10.
The belief space plays a key role by setting some rules for generating new generations of individuals. This space collects and saves normative knowledge of the best group of individuals. The assumption is that best individuals are close to an optimal solution, thus the final solution can be generated by combining components of them. In fact, the belief space defines a new state space for the network by storing best individuals. In the subsequent iterations, new generations of individuals are produced mostly based on this state space.

Our main assumption here is, if the number of iterations approaches infinity, the belief space matrix can accurately represent some information about the level of dependency between the connected nodes. Consequently, these relative frequencies can be used as the probability of a relation in the next timeslot based on the community function. By processing a snap shot of an undirected and unweighted network, a weighted directed graph is made which reveals hidden information about the quality of relations in the network.

### 4.3.2 Computing the Probabilities

To compute the probability of relations of a pair of unconnected nodes in this weighted graph, two criteria have been considered. The first is the number of paths between
each pair of unconnected nodes. The second is the distance between them. The assumption is that existence of more paths between a pair of nodes makes a higher probability of connection between them while the distance must be considered to enhance the weighting system. To reduce the complexity, we assume the paths’ length is always two, which means that the probability is computed for those pairs of unconnected nodes that have only one node between themselves. Let $G(V, E, W)$ denote the input weighted graph, where $V$ is a set of nodes and $E$ is a set of edges between each pair of nodes (hence, each edge $e$ is of the form $(i, j)$, with $i, j \in V$. Furthermore, $W$ is a set of weights of edges, with $0 \leq W(i, j) \leq 1$ for all edges $(i, j)$. For each pair of unconnected nodes $(i, k)$, where $i, k \in V$ and $(i, k) \notin E$, if there is a node $j$ with $j \in V$ and $(i, j), (j, k) \in E$, the estimated weight between $i$ and $k$ is computed as follows:

$$\forall j \in V \rightarrow (i, j), (j, k) \in E, (i, k) \notin E,$$

$$W'(i, j, k) = \max(W(i, j), W(j, i)) \times \max(W(j, k), W(k, j)). \quad (4.12)$$

If there would a link between two nodes $i$ and $k$ in the absence of node $j$, then $W'(i, j, k)$ can be interpreted as the estimated weight of that link.

For each similar path this weight must be computed accordingly, and, finally, the probability of a relation between nodes $i$ and $k$ is computed as follows:

$$P(i, k) = 1 - \frac{1}{2(n + \sum_1^n W'(i, j, k))}, \quad (4.13)$$

where $n$ is the number of paths between $i$ and $k$.

For example, in Fig. 4.10, a direct link does not exist between node#1 and #7 but there are two paths of length two between them. Therefore, $n = 2$, and the nodes #5 and #6 represent $j$. We have $W'(1, 5, 7) = 0.8 \times 0.6 = 0.48$ and $W'(1, 6, 7) =$
0.2 \times 0.6 = 0.12, \text{ and } P(1, 7) = 1 - \left(1/(2 \times (2 + 0.6))\right) = 0.6153.

### 4.3.3 Ranking the Probabilities

After calculating all the probabilities, the predicted pairs must be ranked based on their probabilities. Finally, the top-$L$ of them will be selected as the final predicted edges. This process is shown in the following algorithm:

Algorithm CA-LP (G, A, B, L)

**Input:**
- G: an undirected and unweighted graph, $G(V,E)$
- A: adjacency matrix of G
- B: Belief Space matrix
- L: desired number of top predicted links

**Output:**
- O: $n \times n$ matrix of $L$ probabilities where $O(i,j)=P(i,j)$, ($i, j$ are members of V)

**Main:**
1: Map Belief space to a weighted directed Graph
2: Compute $P(i,k)$ by extracting weights from B according to
   \[(12) \text{ and } (13), \text{ for all pairs where } A(i,k)=0 \text{ and } A(i,j), A(j,k)=1\]
3: Store probabilities in a array
4: Sort the array
5: Choose the top-$L$ and store in O where $O(i,k)=P(i,k)$
4.4 Evaluation

To evaluate the performance of our proposed algorithm (CA), we have used various synthetic networks and real large social network datasets. For the first group of synthetic networks, ten graphs were generated randomly based on Newman’s method in [27]. Each of these graphs has 128 nodes with the degree of 16 and 1024 edges. In addition, each graph consists of four same-sized communities where each community has 32 members. Each of these members has $Z_{in}$ links to other members who are inside its own community and $Z_{out}$ links to members from other communities ($Z_{in} + Z_{out} = 16$). The range of $Z_{out}$ in these ten graphs were set from 3 to 5. In fact, a higher value of $Z_{out}$ leads to generate a more complex network.

Table 4.1: Description of the generated synthetic networks based on Girvan benchmark

<table>
<thead>
<tr>
<th>Fraction</th>
<th>#Nodes</th>
<th>#Edges</th>
<th>$E_T$</th>
<th>$E_P$</th>
<th>U</th>
</tr>
</thead>
<tbody>
<tr>
<td>70%</td>
<td>128</td>
<td>1024</td>
<td>717</td>
<td>307</td>
<td>8128</td>
</tr>
<tr>
<td>80%</td>
<td>128</td>
<td>1024</td>
<td>819</td>
<td>205</td>
<td>8128</td>
</tr>
<tr>
<td>90%</td>
<td>128</td>
<td>1024</td>
<td>922</td>
<td>102</td>
<td>8128</td>
</tr>
</tbody>
</table>

As shown in Table 4.1, we have evaluated the accuracy of our model by changing the fraction of observed edges in the network, $E_T$, from 70% to 90%. The belief space which was imported to the algorithm was obtained from the result of running the community detection algorithm proposed in [41]. We tested the effectiveness of the algorithm according to both AUC and Precision methods. Tests were implemented 100 times independently on the top-100 instances. We also compared the results of AUC with five other similarity metrics, Common Neighbors(CN), Jaccard (JC) and Leicht-Holme-Newman (LH), Adamic/Adar (AA), and Resource Allocation (RA). The results are illustrated in figures. 4.11 and 4.12.

The results clearly show that the proposed algorithm has better performance
in comparison with other metrics on this group of synthetic networks. However, by reducing the number of observed edges, the accuracy of the algorithm reduced significantly. The main reason for this issue is that the community detection algorithm can not find the true communities when the system is very noisy. In fact, by removing a considerable amount of edges from the network randomly, the community structure which is the base of this model will be affected.

Another interesting observation is that, by increasing the complexity of the network ($Z_{out} > 4$) the performance of the algorithm reduced significantly. We believe the cause to be the increasing rate of errors in the community detection algorithm when $Z_{out}$ becomes larger.

In addition to Precision, we also compared the top-1 predicted links calculated by the algorithm, $l = |E^P|$, with the probe set, $E^P$. As a result, in average 78.28% of the predicted links were among the probe set when the fraction of the observed network was 90%. It was 77.85% and 75.14% when the fraction were 80% and 70% respectively. It means that the algorithm could predict the correct links with an accuracy of more than 75%.

We have also evaluated the performance of our algorithm on another group of synthetic networks. The previous benchmark is useful to create a network with 128 nodes but does not have a mechanism to generate bigger network. Therefore, we have generated various networks based on LFR benchmark in [20, 19]. By using this benchmark, we can generate power-law networks with customized features. We generated two groups of 1000 and 3000 nodes graphs. For each group, three graphs have been generated at different levels of complexity. In total, six more synthetic networks have been generated based on the LFR benchmark as shown in Table 4.2.
Table 4.2: Description of the synthetic networks generated based on LFR benchmark

<table>
<thead>
<tr>
<th>Label</th>
<th>#Nodes</th>
<th>#Edges</th>
<th>$\mu$</th>
<th>$D_{\text{average}}$</th>
<th>Cluster Coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td>Network#1</td>
<td>1000</td>
<td>9860</td>
<td>0.300+/- 0.025</td>
<td>19.72</td>
<td>0.344</td>
</tr>
<tr>
<td>Network#2</td>
<td>1000</td>
<td>9804</td>
<td>0.400+/- 0.026</td>
<td>19.608</td>
<td>0.232</td>
</tr>
<tr>
<td>Network#3</td>
<td>1000</td>
<td>9885</td>
<td>0.500+/- 0.023</td>
<td>19.77</td>
<td>0.144</td>
</tr>
<tr>
<td>Network#4</td>
<td>3000</td>
<td>29219</td>
<td>0.300+/- 0.026</td>
<td>19.479</td>
<td>0.326</td>
</tr>
<tr>
<td>Network#5</td>
<td>3000</td>
<td>29074</td>
<td>0.400+/- 0.026</td>
<td>19.383</td>
<td>0.225</td>
</tr>
<tr>
<td>Network#6</td>
<td>3000</td>
<td>29561</td>
<td>0.500+/- 0.023</td>
<td>19.707</td>
<td>0.127</td>
</tr>
</tbody>
</table>

The following parameters have been considered to generate these networks:

- $\beta = 1$, $\beta$ set the exponent for the distribution of community size in the network.
- $\gamma = 2$: $\gamma$ set the exponent for the nodes’ degree distribution.
- $\mu =$ vary from 0.3 to 0.5, $\mu$ is the mixing parameter which determines the ratio of the number of edges between various communities to the total number of them. The higher number means more complex community structure.
- $D_{\text{average}} = 20$, $D_{\text{average}}$ represents the average degree of each node in the graph.
- $D_{\text{Max}} = 50$, $D_{\text{Max}}$ set the maximum degree size for each node.
Figure 4.11: Comparison of the algorithms based on AUC over Girvan benchmark
Figure 4.12: Comparison of the algorithms based on AUC and Precision over Girvan benchmark.

As shown in figures 4.13 and 4.14, the generated graphs with LFR benchmark represent the power-law distribution. In addition, they have a high number of cluster coefficient relatively. Meanwhile, the average distance between nodes in all of them is less than four which represents the small-world effect on the social networks. Consequently, these graphs can represent key features of real social networks. Hence, to evaluate the performance of our algorithm we repeat the experiments using them.
Similar to the previous experiments, a group of edges must be removed randomly to make a probe set. For each network, the probe set was extracted randomly ranging from 10% to 30%. Consequently, the performance and the level of accuracy of the algorithm were evaluated.

The results of the evaluation have been illustrated in Fig.4.15. According to the results, our algorithm has better performance in most of the cases. In fact, as long as community detection algorithm finds the true communities with high accuracy, the link prediction algorithm can work fine. As shown in the diagrams, by increasing the size of $\mu$, the algorithm’s accuracy is reduced which is because of the performance of the community detection algorithm. However, even with this weakness, the proposed algorithm can beat the other methods.

On the other hand, the results show that the trend for the graphs with 3000 nodes is not different from the graphs with 1000 nodes and our proposed model can predict the missing links with a high accuracy. Regarding our observations, the performance of the algorithm is almost the same when 90% and 80% of the network are assigned to the observed set, but when it reduced to 70%, the algorithm’s performance reduced significantly. Moreover, randomly removal of many edges from the network, changes the community structure. This issue is a significant obstacle for the community-detection process. Perhaps using datasets with timestamps is a better option to measure the accuracy of the algorithm which can demonstrate more precision.
Figure 4.13: Synthetic networks (Generated based on LFR benchmark (network#1 to 3))
Figure 4.14: Synthetic networks (Generated based on LFR benchmark (network#4 to 6))
Figure 4.15: Comparison of the algorithms based on AUC over LFR benchmark

We also tested the performance of our proposed algorithm on a real-world big dataset, Orkut, with 117,185,083 edges [40]. The dataset obtained from the Stanford Large Network Dataset Repository [23] is a benchmark dataset used by researchers in social network analysis. Another reason for selecting this dataset is that it is a network with ground-truth communities which make us possible to validate our results. Information about this dataset is represented in Table 4.3.

<table>
<thead>
<tr>
<th>#Nodes</th>
<th>#Edges</th>
<th>Cluster Coefficient</th>
<th>$E^T$</th>
<th>$E^P$</th>
<th>U</th>
</tr>
</thead>
<tbody>
<tr>
<td>3072441</td>
<td>117185083</td>
<td>0.1666</td>
<td>105466575</td>
<td>11718508</td>
<td>4719945313020</td>
</tr>
</tbody>
</table>
The procedure for running the experiment on this dataset is similar to the method described before in experimental setup for synthetic networks. The network was divided into two sets, the training set (90%) and the probe set (10%). After ten iterations of independent experiments, the AUC and the precision were calculated. As shown in Fig. 4.16, the algorithm could estimate the correct links with over 68% success based on the precision method. Regarding the size of the network, we believe that it is an acceptable rate for prediction. We have also repeated the experiment by changing the size of the training set. As shown in Fig. 4.17, when the training set is 90%, the algorithm obtained the highest values for both of AUC and Precision. By reducing it to 80%, a slight reduction can be observed in the level of accuracy. However, when it is decreased to 70% a significant change in accuracy is visible. Although the size of the training set is an effective parameter in this process, but according to our observations, the main reason for this reduction is the random removal of edges which reshapes the structure of the network.

Figure 4.16: Obtained results from the Orkut Dataset
Figure 4.17: Obtained results from the Orkut Dataset (range of training set from 70% to 90%)

4.5 Conclusion and Future Work

In this chapter, the problem of link prediction in social network has been extensively discussed. In addition, we proposed a knowledge-based model to predict the state of a network in the near future. The key part of this model is the belief space which is a probability matrix that shows the level of dependency between linked nodes. Assuming it as an adjacency matrix, a weighted directed graph can be made. Consequently, the probability of relation between two disconnected nodes will be computed based on this graph.

The central hypothesis is that if the network has the community structure, each individual can be joined to at least one community through its friends. Therefore, the quality of relations between pairs of unconnected individuals can be estimated through the community-detection process. Consequently, the present computational model has a community-oriented approach.
We evaluated the performance of our algorithm on various synthetic networks and real-world datasets and compared it with three other metrics. AUC (Area under the curve) and Precision measurements have been used to evaluate the performance of the model against several well-known other methods. The results show that our method can predict the next state of the network with approximately 80% accuracy. The comparisons show that the proposed method has better performance in respect of the performance of other methods.

However, we believe that by increasing the number of iterations in the evolutionary model, the quality of prediction will be improved. Meanwhile, since the size of the belief space is fixed to the number of nodes, the complexity of the algorithm will not change based on the number of iterations or the number of edges.

The main contributions of this research can be summarized in the following items:

• Proposing a method for estimating the quality of links between a pair of nodes in the network

• Introducing a novel method to use the concept of community as a similarity index.

• Employing the cultural algorithm as a knowledge-based evolutionary algorithm for the link prediction problem in social networks.

In the future, we would like to observe the performance of the algorithm in different types of social networks and extend our work to multiple networks. Currently, we have tested the algorithm using the common standard procedure of dividing the training and probe set randomly in the ratio of 70-90% and 10-30%, in the future we would like to test the performance on different rates to find the optimal training size.
References


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Chapter 5

Conclusions

In this dissertation, a new knowledge-based approach for social network analysis has been proposed and extensively reviewed. Introducing new methods for representation, extraction, and utilization of knowledge from the structure of social networks are the main contributions of this research work. Throughout this study, the emphasis was on the role of knowledge in the evolution of societies and solutions. According to the experimental results, utilizing the knowledge has a remarkable impact on enhancing the search process to find the optimal/near optimal solution for the problems, and it is significantly effective and robust.

In addition, we found that MPCA has a tremendous potential to model the social systems. It has derived from the real social systems and has a great capacity to be applied to social network analysis. Therefore, we proposed various models and algorithms based on this idea to deal with some major problems in this research field. Community detection, link prediction and knowledge and population migration in social networks were addressed through our proposed approach in this dissertation.

In the community detection problem, the core question was to find the underlying interconnected structure of a given network. The results indicated that the proposed algorithm and methods profoundly improved the accuracy and the run-time of the
search process. Interestingly, while the target of the algorithm was to find the near-optimal solution, the evaluation results show that the algorithm can find the optimal solutions for a variety of complex social structures in fewer evolution cycles compare to the other evolutionary methods. Reduction of the search space by 80% and enhancing the accuracy by up to 30% in comparison with the genetic algorithms are the major achievements of this approach. The run-time analysis also shows that the proposed method has a negligible impact on the complexity.

In the link prediction problem, the target was to estimate the evolution of a given network in the next time step. Our community-oriented approach could get competitive results in this field while the most significant contribution of it, is to define a method for measuring the quality of connections between pairs of actors in the system. In other words, given a snapshot of a undirected, unweighted network, the proposed algorithm estimates the quality of connections between pairs of nodes and represents them as a directed weighted graph. Introducing a new community-oriented similarity index to calculate the likelihood of a relationship between a pair of unconnected nodes is our another contribution. The results demonstrated that the proposed model is capable of predicting the next state of a given network by the average level of 80% accuracy.

The new concept of migration in the both level of knowledge and population and its role in the process of population adaption has been studied in this field. To the best of our knowledge, it is the first research which addresses these problems. The experimental results show that having prior knowledge about a problem or environment can help an individual to adapt faster to the similar situations if the difference level between two cases is less than 25%. If it is higher, not only knowledge is not helpful, but also it may decrease the performance.

To conclude, an integrated knowledge-based computational model has been introduced in this research study using MPCA and based on the community structure of
the social systems to describe the evolution of complex dynamic social networks. The results clearly demonstrate that this model can significantly increase the efficiency and effectiveness of social network analysis in a variety of tasks and problems. The proposed approach not only can be applied to address the classic problems in the field but also it can be employed to cope with the new challenges and issues such as knowledge migration and population adaptation. This proposed model can be used in a broad range of applications including recommendation systems, resource allocation and organizational and social analysis.

We believe that by employing our proposed approach and techniques, social network analysis can be carried out from a new perspective by focusing on the role of knowledge as the primary asset of the system.

In the future, we would like to employ different sources of knowledge to deal with more social issues through this approach. Co-evolution of networks in multi-layer systems, layer selection strategies and the formation of belief space are our next research targets. Measuring the impact of these sources of knowledge on the adaptation process and enhancing its functionality are also our future works in the field.

In addition, we are going to apply this approach to more real-life datasets extracted from online social networks for business and research-oriented projects such as team formation, group identification, and organizational resource optimization.
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