An Evolutionary and Local Refinement Approach for Community Detection in Signed Networks

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An approach to detect communities in signed networks that combines Genetic Algorithms and local search is proposed. The method optimizes the concepts of modularity and frustration in order to find network divisions far from random partitions, and having positive and dense intra-connections, while sparse and negative inter-connections. A local search strategy to improve the network division is performed by moving nodes having positive connections with nodes of other communities, to neighboring communities, provided that there is an increase in signed modularity. An extensive experimental evaluation on randomly generated networks for which the ground-truth division is known proves that the method is competitive with a state-of-art approach, and it is capable to find accurate solutions. Moreover, a comparison on a real life signed network shows that our approach obtains communities that minimize the positive inter-connections and maximize the negative intra-connections better than the contestant methods.

Keywords: Evolutionary computation; community detection; multiobjective clustering; signed networks; local search.

1. Introduction

In the last few years the rapid diffusion of Internet and social networking has allowed people to connect and exchange opinions and information. The representation of such connections through the concept of network, where a node denotes an individual, and an edge denotes the link between two individuals, has primarily been interpreted as positive. Thus relationships have typically expressed collaboration,
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common interests, membership to the same group, friendship. However, since the primary studies on structural balance theory of Heider, it has been pointed out that relationships between individuals can be either positive or negative, such as like-dislike, friends-enemies, love-hate, trust-distrust. Signed networks are an extension of networks to include the additional information of positive and negative links. Thus positive links denote friendly relations, while negative links represent antagonistic relations. Detecting community structure on these kind of networks is an important research topic since it allows to determine instability inside relationships, and, consequently, to predict changes in group organization.

Approaches to find dense groups of nodes for unsigned networks are mainly based on the optimization of the concept of modularity. The extension of such concept to signed networks has been introduced by Gómez et al.

As regards signed networks, Doreian and Mrvar were the first that proposed a partitioning method by introducing the concept of frustration, which expresses the number of positive ties among different groups and the number of negative links inside the same group.

In this paper a method that combines Genetic Algorithms and a local refinement strategy to detect communities in signed networks is proposed. The method, named SN-MOGA (Signed Networks with MultiObjective Genetic Algorithms) optimizes the concepts of modularity and frustration by applying a Multiobjective Genetic Algorithm. The maximization of modularity allows to detect network divisions far from random divisions, while the minimization of frustration guarantees to have as few negative intra-connections and positive inter-connections as possible. The SN-MOGA algorithm evolves a population of candidate solutions by trying to obtain the best trade-off between high modularity and low frustration. At the end of the evolutionary process a solution from the Pareto front is chosen, and a local search strategy is performed to improve signed modularity, by moving those nodes having positive connections with nodes of other communities, to neighboring communities, provided that there is an increase in signed modularity.

The idea of modeling community detection in both signed and unsigned networks as a problem of optimizing multiple objectives is not new. Moreover, studies on which kinds of objectives for unsigned networks should be selected to improve the performance of a method, along with the advantages of using multiobjective optimization when objective functions are negatively correlated, have been discussed by Shi et al. In this context, the main contributions of our work consist in coupling multiobjective optimization with a local search strategy to improve the solution obtained from the Pareto front. A correlation analysis of the two objective functions employed by the method shows that the two objectives are negatively correlated, thus suitable for multiobjective optimizations, according to the observations reported by Shi et al. that negatively correlated objectives lead to better performances compared with single-objective or positively correlated objectives.
An extensive experimentation on synthetic networks and real life networks shows that our multiobjective approach optimizing signed modularity and frustration is capable to divide signed networks in groups of nodes having high accuracy and low edge misclassification. Comparison with other state-of-the-art methods indicates that SN-MOGA obtains network partitioning more meaningful and closer to the ground truth division.

The paper is organized as follows. In the next section an overview of existing approaches to community detection in signed networks is given. In Section 3 preliminary definitions are introduced, and the problem is clearly stated. Section 4 presents the algorithm. Section 5 describes the evaluation measures adopted for assessing the method results. Section 6 evaluates the performance of our method on synthetic generated networks for which the ground truth division is known, and a real life network. Moreover, a comparison with existing state-of-the art methods is reported. Section 7 compares SN-MOGA with the Particle Swarm Optimization method of Gong et al. Section 8 analyzes the running time of the method. Section 9, finally, concludes the paper.

2. Related Work

In this section we give an overview of the main proposals to find communities in signed networks.

Signed networks originate from the studies of Heider on structural balance theory. The idea underlying balancing is that if two people $i$ and $j$ belonging to the same group like each other, then their evaluation regarding other people should be consistent, that is if $i$ and $j$ like each other, then they both either dislike or like the same people, and if $i$ and $j$ dislike each other, they disagree in evaluating others. A triad is defined balanced if the product of its edge signs is positive. If all the triads in a network are balanced, the network is balanced. It has been proved that in a balanced network the set of vertices can be divided into two clusters such that positive links are only within clusters, while negative links are between clusters. However, rarely a network has a 2-way partitioning, thus Davis extended the concept of balance to $k$-balance. A network is $k$-balanced if it can be divided into $k$ groups such that, edges within groups are positive and edges between groups are negative. In such a case the network is also said partitionable or clusterable, while the term balanced is generally used for 2-way balance. $k$-balancing is an important research topic since balancing assures stability, while imbalance generates tension inside a group.

One of the first partitioning approaches to structural balance has been proposed by Doreian and Mrvar. The method randomly divides the nodes in a fixed number $k$ of clusters, and then tries to optimize a criterion function by moving nodes among neighboring partitions. The criterion function they proposed is frustration (see next section for a formal definition). The neighbors of each partition are computed, then, either a node is moved to a neighboring cluster, or two nodes are exchanged between two neighboring groups. These neighbors are examined at random, and, if the new
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partition has a lower value of frustration, the new solution is accepted. The main
drawbacks of the method are that the number of groups must be given as input
parameter, and that it disregards the density of links, which is one of the main
characteristics exploited in unsigned community detection methods.

More recently, because of the increasing interest in signed networks, several
approaches have been proposed. Many of these methods extend concepts used to
detect communities in unsigned networks, to take into account the sign of links.

Yang et al.\textsuperscript{11} proposed an algorithm that uses the concept of random walk and
adopts an agent-based heuristic to extract communities. The method starts with an
arbitrary node. From this node an agent performs a random walk for a number of
steps by visiting one of the neighboring nodes on the base of the transition probabil-
ity computed from the network connectivity degree. The method, named FEC, is
composed of two main phases. The FC (Find a Community) phase transforms the
adjacency matrix of the graph by applying iterative operations, in order to compute
aggregate transition probabilities, and sorts them for each row. The EC (Extract
the sink Community) phase divides the transformed matrix in two blocks by ap-
plying a cutoff criterion. One of the two blocks is identified as a community, called
the sink community, while the remaining block is recursively processed in the same
way. A main problem is the definition of the cutoff value. To this end the authors
proposed a variation of the cut concept used in spectral clustering\textsuperscript{12} that takes into
account the sign of edges. FEC needs as input parameter the number \( l \) of steps the
agent performs before arriving to a destination node. A sensitivity analysis of this
parameter shows that when \( l \) is greater than a range of values between 10 and 20,
the FC phase is insensitive to this parameter.

Approaches that extend the concept of modularity of Newman and Girvan\textsuperscript{3} have
been proposed by Traag and Bruggeman\textsuperscript{13} and Gómez et al.\textsuperscript{4} The former extended
modularity with negative signs and formalized the concept as a Potts model.\textsuperscript{14}
They defined a Hamiltonian for the positive part and another for the negative one,
by extending the approach of Reichardt and Bornholdt.\textsuperscript{14} Then minimizing the
Hamiltonian is shown to be equivalent to maximizing modularity. To this end the
authors modified the simulated annealing approach of Reichardt and Bornholdt,\textsuperscript{14}
and showed an application of the method to a network of conflicts and alliances
between countries. Gómez et al.,\textsuperscript{4} instead, generalized the concept of modularity
to signed networks and proposed to maximize signed modularity to detect commu-
nities. They applied the approach to a real network related to retail stores in the
city of Lyon, and found that the results they obtained were better when compared
with the classification provided by a public institution.

Spectral graph theory is another important concept extensively used to find com-
munities. In this context, Kunegis et al.,\textsuperscript{15} in order to deal with signed networks,
defined the signed Laplacian matrix of a graph, investigated its properties, gave a
definition of signed ratio cut, and proposed a spectral approach to find a clustering
by minimizing ratio cut. Moreover, they exploited these concepts also for graph
visualization and link prediction. Chiang et al.\textsuperscript{16} proposed a multilevel clustering
algorithm that introduces new $k$-way objectives and kernels. These objectives are shown to be equivalent to a general weighted kernel $k$-means objective, thus the optimization of these objectives can be performed by using a kernel $k$-means like algorithm. Furthermore the authors show that the approach of Kunegis et al.,\cite{15} presents some weakness when directly generalizes the signed Laplacian to $k$-way clustering. Anchuri and Magdon-Ismail\cite{17} proposed a two step spectral approach to detect communities in signed social networks. An input parameter to fix the value of the leading eigenvector, used to assign nodes to communities, must be given. They consider the concepts of frustration\cite{5} and modularity\cite{3} and detect communities by optimizing only one of these two objectives at a time. After that, they try to improve the chosen objective by moving nodes among communities. The authors formalize the problem of minimizing frustration and maximizing modularity to that of maximizing the mathematical form $f(M, s) = s^TMs$, where $s \in \{-1, +1\}^n$ is an $n$-dimensional vector. When $s$ is equal to the eigenvector corresponding to the maximum eigenvalue of $M$, it maximizes $f(M, s)$. The top eigenvector is computed by using the Power Iteration method. Since the method finds a partitioning in two communities, it can be extended to a higher number of communities by iteratively dividing communities until the objective cannot be improved any more. Experiments on two real-life signed networks show that when the objective function is modularity, the two step approach obtains the minimum frustration value with respect to modularity maximization without improvement, and $k$-means approaches. Same results are obtained when minimizing frustration with and without improvement.

A different approach based on simulated annealing has been presented by Bogdanov et al.\cite{18} The authors proposed a framework for building signed networks from content generation flow. Validation is performed on two case studies of articles extracted from Wikimedia download site. Since the number $k$ of clusters to find must be provided in input, the authors vary the value of $k$ from 2 to 10, and compute the criterion they optimize to obtain a partitioning. They choose the value of $k$ for which a higher value does not increase the optimization criterion.

One of the main limitations of these approaches is that the number $k$ of clusters must be given as input parameter. Thus some strategy must be introduced to determine $k$, such as executing the method for a range of $k$ values, and then choosing the $k$ giving the best value of the criterion that the method optimizes.

Recently, however, methods based on evolutionary computation, that automatically determine the number of partitions, have been proposed. Li et al.\cite{19} presented and compared two evolutionary algorithms, named $EA$-$SN$ and $CSA$-$SN$, and two memetic algorithms, named $EA_{HC}$-$SN$ and $CSA_{HC}$-$SN$. The latter two differ with respect to the formers since they include a hill-climbing strategy. All the algorithms adopt the character string encoding of individuals, i.e. each node is associated with the label of the cluster it belongs to, and use as objective function to optimize the improved modularity and the improved modularity density. The first one is the modularity extended by Gómez et al.\cite{4} to signed networks, while the latter is a
generalization to networks with signs of the modularity density concept, proposed by the same authors for unsigned networks. Experiments on different networks show that the memetic approaches outperform the evolutionary approaches.

Liu et al.\textsuperscript{20} proposed a multiobjective evolutionary method to find communities in signed networks, named MEA\textsubscript{s-SN}. The two objectives to optimize are based on the concepts of positive and negative cluster similarity. The authors extend the definition of similarity of Huang et al.\textsuperscript{21} between two neighboring nodes to signed links, and define the first objective as the positive internal and external similarity of a community structure, while the second objective as the negative internal and external similarity of a community structure. Moreover, they propose a representation of individuals consisting of two components. The first component is a node permutation, the second component denotes the cluster label the node belongs to. In order to determine this label, MEA\textsubscript{s-SN} performs a community detection method that starts by an empty cluster and adds a node, provided that a criterion, named signed tightness, increases. This approach allows the method to assign a node to multiple communities. The method has been compared with the algorithm FEC of Yang et al.,\textsuperscript{11} with CSA\textsubscript{HC-SN} of Li et al.,\textsuperscript{19} and an extension of the Blondel et al.\textsuperscript{22} method. The authors showed that their approach outperforms the competitors.

A different bio-inspired approach has been proposed by Gong et al.\textsuperscript{9} They introduced a multiobjective discrete particle swarm optimization algorithm, called MODPSO, to solve the network clustering problem by optimizing two objective functions, the kernel k-means and the ratio-cut. Though the method is proposed for unsigned networks, the authors extended the two fitness functions for signed networks, and presented results also on 4 small sized networks, used by Yang et al.\textsuperscript{11} to evaluate the FEC algorithm.

The method we propose, analogously to Liu et al.,\textsuperscript{20} is based on multiobjective optimization. However the two approaches are different in many aspects. First of all MEA\textsubscript{s-SN} uses an individual representation that combines both cluster label and node permutation, SN-MOGA, instead, as will be clear in Section 4, adopts the locus-based representation. The objective functions the two algorithms optimize are also different. MEA\textsubscript{s-SN} adapts the community fitness introduced by Lancichinetti et al.\textsuperscript{23} to signed networks, while SN-MOGA uses signed modularity and frustration. In the experimental result section we compare SN-MOGA with MEA\textsubscript{s-SN} on synthetic networks, and with MEA\textsubscript{s-SN} and Chiang et al.\textsuperscript{16} method on the real-life network Wikipedia. Moreover, a comparison with MODPSO is also reported on four popular signed networks. We show that SN-MOGA is very competitive with respect to these methods.

3. Notation and Definitions

A signed social network can be modeled as a graph $G = (V, E, W)$, where $V$ is the set of $n$ nodes (vertices) and $E$ is the set of $m$ edges. $W : V \times V \rightarrow \{-1,0,1\}$
is a function which assigns +1 to edges connecting positively a pair of nodes, −1 to edges that connect negatively a pair of nodes, and 0 if an edge does not exist between the nodes.

Let \( A \) denote the weighted adjacency matrix associated with \( G \), i.e. \( A_{i,j} = W(i,j) \). The matrix \( A \) can be split into two adjacency matrices corresponding to positive and negative edges by setting \( A_{i,j}^+ = A_{i,j} \) if \( A_{i,j} > 0 \), zero otherwise, and \( A_{i,j}^- = -A_{i,j} \) if \( A_{i,j} < 0 \), zero otherwise. Thus

\[
A = A^+ - A^-
\]  

(1)

Given a node \( i \in V \), \( a_i^+ \) and \( a_i^- \) are defined respectively as the positive degree and the negative degree of \( i \).

Now consider a division \( C = \{C_1, \ldots, C_k\} \) of the graph \( G \) into \( k \) communities.

Frustration \( F(C) \) of a network partition \( C = \{C_1, \ldots, C_k\} \) is defined as the sum of the number of positive edges between nodes belonging to different communities and the number of negative edges between nodes inside the same community.\(^5\)

\[
F(C) = \sum_{i,j \in V} \alpha A_{i,j}^- \delta(c_i, c_j) + (1 - \alpha) A_{i,j}^+ (1 - \delta(c_i, c_j))
\]  

(2)

where \( c_i \) (\( c_j \)) is the community of node \( i \) (\( j \)) and \( \delta(c_i, c_j) \) is the Kronecker delta function which takes the value 1 if nodes \( i \) and \( j \) belong to the same community, 0 otherwise, and \( 0 \leq \alpha \leq 1 \) is a parameter that allows to give a different weight to positive and negative links. In the following we do not differentiate the importance of links, thus we consider frustration without this parameter.

Frustration can be rewritten as:

\[
F(C) = \sum_{r=1}^{k} \left( \sum_{i,j \in C_r} A_{i,j}^- + \sum_{i \in C_r, j \notin C_r} A_{i,j}^+ \right)
\]  

(3)

now let

\[
l_r^- = \sum_{i,j \in C_r} A_{i,j}^-
\]  

(4)

and

\[
\gamma_r^+ = \sum_{i \in C_r, j \notin C_r} A_{i,j}^+
\]  

(5)

Then frustration can be expressed as:

\[
F(C) = \sum_{r=1}^{k} (l_r^- + \gamma_r^+)
\]  

(6)

The concept of modularity has been introduced by Newman and Girvan in Ref. 3. Intuitively, it is the difference between the fraction of edges inside a community, and the expected value of the fraction of edges that would be in the network if edges fell at random without regard to community structure. For signed networks
the definition of modularity is modified to take into account the contribution of both positive and negative edges.

Signed modularity can be defined as:

\[ Q^S = \frac{1}{2m^+ + 2m^-} \sum_{i,j \in V} \left( A_{i,j} + \frac{a_i^- a_j^-}{2m^-} - \frac{a_i^+ a_j^+}{2m^+} \right) \delta(c_i, c_j) \]  

(7)

where \( m^+ \) and \( m^- \) are the number of positive and negative entries in \( A \), respectively. Let \( m = m^+ + m^- \). Since the Kronecker function \( \delta(c_i, c_j) \) is equal to 1 if the nodes \( i \) and \( j \) are in the same community, similarly to frustration, the signed modularity can be reformulated as:

\[ Q^S = \sum_{r=1}^{k} \left[ \sum_{i,j \in C_r} A_{i,j} + \sum_{i \in C_r} (a_i^-)^2 - \sum_{i \in C_r} (a_i^+)^2 \right] \]  

(8)

Now let

\[ l_r = \sum_{i,j \in C_r} A_{i,j} \]  

(9)

\[ d_r^+ = \sum_{i \in C_r} (a_i^+)^2 \]  

(10)

\[ d_r^- = \sum_{i \in C_r} (a_i^-)^2 \]  

(11)

Signed modularity can be rewritten as:

\[ Q^S = \sum_{r=1}^{k} \left[ \frac{l_r}{2m} + \frac{(d_r^-)^2}{2m^- (2m)} - \frac{(d_r^+)^2}{2m^+ (2m)} \right] \]  

(12)

Note that, if either \( m^+ \) or \( m^- \) are zero, then \( Q^S \) cannot be computed, thus we assume that its value is zero.

Given a graph \( G = (V, E, W) \) modeling a signed network, our objective is to find a partitioning of \( G \) in \( k \) clusters such that: (1) intra-connections are dense and most edges within clusters are positive; (2) inter-connections between clusters are sparse and most of these edges are negative.

Shi et al.\(^8\) performed an experimental study aiming at comparing different objective functions in multiobjective community detection methods, in order to choose the objectives leading to better performances. To this end, for a given network, they generated random partitions, and then computed the values of different objective functions. After that, they estimated the Pearson correlation coefficient among the objectives and observed that only negatively correlated objectives are suitable for multiobjective optimization. In fact they (1) provoke opposite effects on the number of communities, (2) avoid an algorithm to converge to trivial solutions, (3) enhance diversity and avoid premature convergence. Positively correlated objectives, instead, are equivalent to single objective methods.
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In order to verify whether the two objectives of signed modularity and frustration satisfy the property of being negatively correlated, we performed an analogous study on the synthetic networks described in Section 6. The Pearson correlation coefficient we obtained was $-0.3244$, showing, thus, a negative correlation. This result strengthens the suitability of these two objectives in discovering meaningful solutions.

4. SN-MOGA Description

In this section we give a description of the multiobjective algorithm SN-MOGA for signed networks, the representation adopted for partitioning the network, and the variation operators used.

We used the Nondominated Sorting Genetic Algorithm (NSGA-II) proposed by Srinivas and Deb in Ref. 24 and implemented in the Global Optimization Toolbox of MATLAB. SN-MOGA has been adapted with a customized population type that suitably represents a partitioning of a network and endowed with the complementary objectives.

Genetic Representation. The algorithm uses the locus-based adjacency representation proposed in Ref. 25. An individual of the population consists of $n$ genes $g_1, \ldots, g_n$, and each gene assumes a value $j$ in the range $\{1, \ldots, n\}$. Each gene corresponds to a node of the graph $G$ modeling the network. If the value of the $i$th gene is $j$, it means that there is an edge between nodes $i$ and $j$, and that both $i$ and $j$ belong to the same cluster. In this representation the number of clusters is determined by the number of connected components contained in an individual. Figure 1(a) shows a signed network of 12 nodes clusterable in the two groups $\{1, 2, 3, 4, 5, 6\}$ and $\{7, 8, 9, 10, 11, 12\}$. Dashed lines correspond to negative links, while solid lines to positive edges. The genotype corresponding to this division is

$2\ 3\ 5\ 6\ 3\ 1\ 8\ 7\ 12\ 11\ 7\ 7$
shown in Fig. 1(b) and it is interpreted as: node 1 is connected with node 2, node 2 with node 3, node 3 with node 5, and so on.

**Initialization.** A random individual is generated such that if in the $i$th position there is a value $j$, then $j$ must be one of the neighbors of $i$, i.e. the edge $(i, j)$ must exist.

**Uniform Crossover.** $SN-MOGA$ uses a standard uniform crossover operator. First a crossover mask of length $n$, i.e. the number of nodes, is randomly generated. Each value on the mask is either 0 or 1. An offspring is created by selecting from the first parent the genes where the mask is a 0, and from the second parent the genes where the mask is a 1. Uniform crossover guarantees the maintenance of the effective connections of the nodes in the network in the child individual.

**Mutation.** Analogously to initialization, fixed a position $i$, mutation randomly selects one of the neighbors of $i$ and assigns this value to the $i$th gene.

**Fitness Functions.** The two objectives to optimize are signed modularity (formula (12)) and frustration (formula (6)).

**Solution Selection.** Multiobjective optimization techniques do not return a unique solution to a problem, but a set of solutions are found through the use of Pareto optimality theory. In this context, since a vector of competing objectives must be simultaneously optimized, the goal is to obtain Pareto-optimal solutions, i.e. non-dominated solutions for which an improvement in one objective requires a degradation of another (Pareto front). Thus the Pareto front represents the compromise solutions satisfying all the objectives as best as possible. However, a solution, out of the Pareto front, should be selected. In our case we show the results when choosing the solution having the minimum frustration and that having the maximum signed modularity.

The pseudo-code of the algorithm is reported in Fig. 2. $SN-MOGA$ starts with a randomly generated population of individuals (step 1) and performs multiobjective optimization for a number of generations (steps 2–4). Then it chooses a solution from the Pareto front (step 5) and tries to improve signed modularity by moving nodes, having positive connections with nodes belonging to other clusters, to neighboring communities (steps 6–8). In the experimental result section we will show that $SN-MOGA$ is able to obtain highly accurate partitioning of the signed networks we consider.

**Computational Complexity.** $SN-MOGA$, as already described, uses the NSGA-II method. In Ref. 27 it has been proved that the run-time complexity of the NSGA-II algorithm is $O(gp \log^{b-1} p)$, where $g$ is the number of generations, $p$ is the population size, and $h$ is the number of objective functions. Since the number $h$ of objectives of $SN-MOGA$ is two, its complexity is $O(gp \log p)$. As regards genetic
**SN-MOGA Method:**

**Input:** A signed network $SN$ and the graph $G = (V, E, W)$ modeling it

**Output:** A node cluster labeling that partitions $SN$ in the optimal community structure

1. Create a population of random individuals whose length equals the number $N = |V|$ of nodes of $G$
2. While not maxGen
3. Perform a multiobjective GA with objectives
   3.1 $F(C)$ (formula (6))
   3.2 $Q_S$ (formula (12))
4. End while
5. Choose the solution $C = \{C_1, \ldots, C_k\}$ of the Pareto front having the either maximum signed modularity or minimum frustration value;
6. For each node $v_j$ of a cluster $C_i$ having at least a positive link with a node belonging to a cluster $C_l$
7. Move $v_j$ to $C_l$ provided that signed modularity $Q_S$ augments

Fig. 2. The pseudo-code of the SN-MOGA algorithm.

operators, at each generation, crossover needs $O(n)$ time, mutation $O(1)$ time, while fitness computation is composed of three terms: decoding of an individual in connected components, modularity and frustration computation. Decoding requires $O(n \log n)$ time. To compute modularity and frustration, for each node $i$ its $a_i^+$ and $a_i^-$ neighbors must be considered, then the time complexity is $O(m)$, where $m$ is the total number of edges. Fitness computation can thus be computed in $O(n \log n) + O(m) + O(m)$ time. The overall complexity of SN-MOGA is thus $O((gp \log p) \times (n \log n + m))$.

Before presenting the results, in the next section the measures used to evaluate the method are described.

5. Evaluation Measures

To validate our approach and compare it with other methods, we consider two evaluation measures: the error, as defined by Yang et al., useful when no information regarding the community structure is available, and a modified version of the well known information theory concept of normalized mutual information (NMI), applicable when the ground-truth division of the network is given.

**Error.** Yang et al. employed the frustration concept to define the error rate of a signed network partitioning $C$ as

$$error(C) = \frac{F(C)}{\sum_{i,j} |A_{i,j}|} \times 100\%$$  \hspace{1cm} (13)
As pointed out by the authors, this error function considers only the sign of the links, and completely disregards the edge density.

**Normalized Mutual Information (NMI).** When the ground-truth division of a network is known, a very popular measure to compare community structures, based on information theory, is the **Normalized Mutual Information** (NMI).

The normalized mutual information $\text{NMI}(A, B)$ of two divisions $A$ and $B$ of a network is defined as follows. Let $C$ be the confusion matrix whose element $C_{ij}$ is the number of nodes of community $i$ of the partition $A$ that are also in the community $j$ of the partition $B$.

$$\text{NMI}(A, B) = \frac{-2 \sum_{i=1}^{c_A} \sum_{j=1}^{c_B} C_{ij} \log(C_{ij} n / C_i C_j)}{\sum_{i=1}^{c_A} C_i \log(C_i / n) + \sum_{j=1}^{c_B} C_j \log(C_j / n)}$$

(14)

where $c_A$ ($c_B$) is the number of groups in the partition $A$ ($B$), $C_i$ ($C_j$) is the sum of the elements of $C$ in row $i$ (column $j$), and $n$ is the number of nodes. The denominator is a normalization factor that limits the range of values in the interval $[0, 1]$. Different types of normalizations have been proposed. We adopt the same used by Danon et al. for complex networks. If $A = B$, $\text{NMI}(A, B) = 1$, if $A$ and $B$ are completely different, $\text{NMI}(A, B) = 0$.

**Weighted Normalized Mutual Information (WNMI).** Recently, it has been proved that NMI suffers of the so called **selection bias**, i.e. the leaning to choose clustering solutions having many clusters or with fewer data points when compared with the ground-truth clustering. This provokes an unfair favorable behavior towards those methods that find a high number of clusters, independently from the true effective number. Consider, for instance, the toy example of Fig. 1(a). The division of the network reported in Fig. 3 into the three clusters $\{1, 2, 3, 6, 7\}$, $\{5, 11\}$, $\{4, 8, 9, 10, 12\}$, when compared with the ground truth division, has an NMI value of 0.1866. However, if consider the partitioning constituted by 12 singleton communities the NMI value is 0.4362, which is rather unintuitive. Thus, the importance of the NMI value for 12 singleton communities is 0.4362.
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of correcting NMI when the number of clusters is high with respect to the data size\(^{34,35,33,36}\) has been discussed, and modifications proposed.

In this paper we adopted the adjusted NMI measure proposed by Amelio and Pizzuti\(^{36}\) because, as experimentally demonstrated, it is fast to compute, differently from the high computing time required by the measure of Romano et al.,\(^{33}\) and avoids to consider very similar a predicted and the ground truth clustering when the former consists of a too few or too high number of communities with respect to the latter.

Let \(A\) and \(B\) be the ground-truth division of a network in \(c_A\) communities, and the partitioning in \(c_B\) communities obtained by a method, respectively. The weighted NMI is defined as follows:

\[
WNMI = e^{-\frac{|c_A - c_B|}{c_A}} \times \text{NMI}
\]  

(15)

The exponent of the exponential function is 0 when the predicted number \(c_B\) and the true number \(c_A\) are the same. In this case, thus, the weighted NMI and NMI values coincide. However, as the difference between \(c_A\) and \(c_B\) increases, both if either a lower or a higher number \(c_B\) of communities is obtained, the value of WNMI proportionally decreases.

In the next section we test our method and compare it with other state-of-the-art approaches by showing both the NMI and WNMI values the methods obtain.

6. Experimental Results

In this section we evaluate the capability of our approach in obtaining meaningful partitions of signed networks. As regards the parameters needed by \(SN-MOGA\), in order to set crossover and mutation rate, we executed the algorithm on the synthetic networks described in detail in the next section, by considering values between 0.1 and 0.4 for mutation, and 0.1 until 1 for crossover. Figure 4 shows the NMI and modularity values for the combinations of these values. The figure points out that there are a number of combinations that give high NMI. However, since it is known that high mutation rate could destroy good solutions, while low values do not help in escaping from local optima, we fixed it to 0.2 along with crossover fraction 0.8, which gives both high modularity and NMI values. Moreover, we set elite reproduction 10% of the population size, roulette selection function, population size 100, number of generations 100. These values have already been experimented for community detection in unsigned networks and showed to give good results. The algorithm has been executed 10 times and the average values of error rate, NMI and weighted NMI have been computed, together with standard deviation. For all the experiments, the statistical significance of the results produced has been checked by performing a t-test at the 5% significance level. The p-values returned are very small, between 2.3534e-65 and 1.1120e-30, thus the significance level is very high since the probability that a community could be obtained by chance is very low.
6.1. Evaluation on synthetic networks

In this section a more deep study on synthetic networks generated analogously to Yang et al.\textsuperscript{11} with control parameters that determine the structure of communities, is performed. In particular, we modified the benchmark proposed by Lancichinetti \textit{et al.},\textsuperscript{37} which is an extension of the classical benchmark of Girvan and Newman,\textsuperscript{38} by assigning a controlled sign to edges.

The networks consist of 128 nodes divided into four communities of 32 nodes each. Every node has an average degree of 16 and shares a fraction $\mu$ of edges with the other nodes of the network, and $1 - \mu$ of links with the nodes of its community. $\mu$ is called the \textit{mixing parameter}. When $\mu < 0.5$ the neighbors of a node inside its group are more than the neighbors belonging to the other three groups, thus a good algorithm should discover them. We generated 10 different networks for values of $\mu$ ranging from 0.1 to 0.5. In order to make the networks signed, analogously to Yang \textit{et al.},\textsuperscript{11} we used two parameters $p_-$, denoting the probability of negative links appearing within communities, and $p_+$, denoting the probability of positive links appearing between communities. Thus, for all the combinations of $p_-$ and $p_+$ values.

Fig. 4. (Color online) NMI and modularity values for mutation rate varying in the interval $[0.1, 0.2, 0.3, 0.4]$ and crossover fraction from 0.1 to 1.
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Fig. 5. (Color online) (a) NMI and (b) Weighted NMI corresponding to the maximum modularity values obtained from $SN$-$MOGA$ for all the possible $p^+$ and $p^-$ values at different values of the $\mu$ parameter.

ranging in the interval $[0, 0.1, \ldots, 1]$, we randomly assigned a negative sign to edges inside a community with probability $p^-$, and a positive sign to edges between two different communities with probability $p^+$.

Figures 5(a) and (b) depict the NMI and WNMI values obtained by running $SN$-$MOGA$ for all the combinations of parameters $\mu = [0.1, \ldots, 0.5], p^- = [0, 0.1, \ldots, 1]$, and $p^+ = [0, 0.1, \ldots, 1]$ when selecting from the Pareto front the community
structure having the highest modularity value. The figures point out that the NMI and WNMI values do not sensibly differ, meaning that the number of communities found by SN-MOGA is close to the true number, which is 4. From the figures it can be observed that, fixed a $\mu$ value, the method is not sensitive to the increase of the number of positive edges between communities. As regards $p_-$, until
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Fig. 7. (Color online) Average number of clusters obtained by SN-MOGA for $p_+$ and $p_-$ varying in the interval $[0,1]$, at different values of the $\mu$ parameter, when (a) maximum modularity and (b) minimum frustration are selected from the Pareto front.

$p_- \leq 0.4$, SN-MOGA maintains high values, however it is negatively influenced by the augmentation above 0.4 of negative links within a community.

A similar behavior can be observed in Figs. 6(a) and (b), where the solutions having the minimum frustration are now selected from the Pareto front. In this case the NMI and WNMI values obtained are lower with respect to the previous case, moreover SN-MOGA is again insensitive to the variation of negative edges for $p_- \leq 0.4$.

Figures 7(a) and (b) show the average number of clusters obtained by SN-MOGA for $p_+$ and $p_-$ varying in the interval $[0,1]$, when maximum modularity and
Fig. 8. (Color online) Error rate values obtained by SN-MOGA for $p_+$ and $p_-$ varying in the interval [0, 1] and $\mu = 0.1$.

Fig. 9. (Color online) Error rate values obtained by SN-MOGA for $p_+$ and $p_-$ varying in the interval [0, 1] and $\mu = 0.2$.

minimum frustration, respectively, are selected from the Pareto front. It can be observed that, in the former case, the number of communities is 4 for a good range of $p_-$ and $p_+$ combinations, and it almost always is not greater than 8. Solutions with minimum frustration divide the networks in more communities, particularly when $p_-$ increases. This implies a decrease in NMI and WNMI values.

Figures 8–12 show the error rate obtained by SN-MOGA, when minimum frustration solutions are chosen from the Pareto front, for increasing values of the
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Fig. 10. (Color online) Error rate values obtained by SN-MOGA for \( p_+ \) and \( p_- \) varying in the interval \([0, 1]\) and \( \mu = 0.3 \).

Fig. 11. (Color online) Error rate values obtained by SN-MOGA for \( p_+ \) and \( p_- \) varying in the interval \([0, 1]\) and \( \mu = 0.4 \).

mixing parameter \( \mu \), and combinations of \( p_- \) and \( p_+ \) values. The figures point out that the error rate is insensitive to increasing values of \( p_+ \), i.e. the augmentation of positive links between different communities does not provoke abrupt changes in the frustration value. However, the error rate increases as the percentage \( p_- \) of negative links inside the same community augments.
Fig. 12. (Color online) Error rate values obtained by \textit{SN-MOGA} for $p_+$ and $p_-$ varying in the interval $[0,1]$ and $\mu = 0.5$.

Fig. 13. (Color online) Signed modularity values obtained by \textit{SN-MOGA} for $p_+$ and $p_-$ varying in the interval $[0,1]$ and $\mu = 0.1, \ldots, 0.5$.

Finally, Fig. 13 depicts the signed modularity values obtained by \textit{SN-MOGA} for $p_+$ and $p_-$ varying in the interval $[0,1]$ and $\mu = 0.1, \ldots, 0.5$. It is worth observing that modularity values are high for $p_- \leq 0.4$, analogously to the NMI and WNMI values.
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Fig. 14. CPU time required by SN-MOGA when the number of processors doubles from 1 until 32.

6.2. Comparison with MEA_s-SN on synthetic networks

In this section we compare SN-MOGA with the method proposed by Liu et al.\textsuperscript{20} MEA_s-SN is one of the most recent evolutionary based proposals for signed networks, and it has been shown to outperform state-of-the-art methods. In order to compare the two methods, we generated an LFR benchmark, as proposed by Lancichinetti et al.\textsuperscript{37} constituted by 1000 nodes, average node degree 20, maximum node degree 50, exponent of degree distribution $-2$, community size distribution $-1$, mixing parameter $\mu$ varying as $0 \leq \mu \leq 0.5$. Also for this benchmark, in order to obtain signed networks, for all the combinations of $p_-$ and $p_+$ values ranging in the interval $[0, 0.1, \ldots, 1]$, we randomly assigned a negative sign to edges inside a community with probability $p_-$, and a positive sign to edges between two different communities with probability $p_+$. A benchmark with analogous characteristics has been used by Liu et al. to evaluate their method. We executed MEA_s-SN with the parameters suggested by the authors, i.e. number of generations 100, population size 100, crossover fraction 0.8, mutation rate 0.2, and then selected from the final population the solution having the maximum signed modularity. As regards SN-MOGA we fixed the same parameters of MEA_s-SN.

Figures 15–24 show the average values of NMI, number of communities obtained by the methods, and Weighted NMI for $p_-$ and $p_+$ ranging in the interval $[0, 1]$. For each experiment, the true number of communities is also reported. In particular, when $\mu = 0.1, 0.2, 0.3, 0.4, 0.5$ the corresponding average ground truth number of communities is 28, 28, 33, 32, and 32, respectively.

From Figs. 15–24 we can observe that MEA_s-SN has the tendency to generate a number of communities much higher than the true numbers. For instance, when $\mu = 0.1$ and $p_- \leq 0.4$ (corresponding to the first 5 rows of Fig. 15) it partitions the networks in a number of communities between 40 and 70, depending on $p_+$. When $p_- \geq 0.5$, this number sensibly increases and can reach 300. The corresponding NMI is, in these cases, unreasonably high, always above 0.8, except for $p_- = 1$. 

by Dr Clara Pizzuti on 10/04/16. For personal use only.
Fig. 15. (Color online) NMI, number of communities, and weighted NMI for all the combinations of $p^+ = \{0, \ldots, 1\}$ and $p^- = \{0, \ldots, 1\}$, when $\mu = 0.1$. The ground truth number of communities is 28. Each row corresponds to a $p^-$ value, starting from $p^- = 0$ on the first row, and $p^- = 0.4$ on the last row.
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Fig. 16. (Color online) NMI, number of communities, and weighted NMI for all the combinations of
$p_+ = \{0, \ldots, 1\}$ and $p_- = \{0, \ldots, 1\}$, when $\mu = 0$.

1. The ground truth number of communities is 28. Each row corresponds to a $p_-$ value, starting from $p_- = 0.5$ on the first row, and $p_- = 1$ on the last row.
Fig. 16. (Continued)
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Fig. 17. (Color online) NMI, number of communities, and weighted NMI for all the combinations of $p^+ = \{0, \ldots, 1\}$ and $p^- = \{0, \ldots, 1\}$, when $\mu = 0.2$. The ground truth number of communities is 28. Each row corresponds to a $p^-$ value, starting from $p^- = 0$ on the first row, and $p^- = 0.4$ on the last row.
Fig. 18. (Color online) NMI, number of communities, and weighted NMI for all the combinations of $p^+ = \{0, \ldots, 1\}$ and $p^- = \{0, \ldots, 1\}$, when $\mu = 0.2$. The ground truth number of communities is 28. Each row corresponds to a $p^-$ value, starting from $p^- = 0.5$ on the first row, and $p^- = 1$ on the last row.
Fig. 18. (Continued)
Fig. 19. (Color online) NMI, number of communities, and weighted NMI for all the combinations of $p^+ = \{0, \ldots, 1\}$ and $p^- = \{0, \ldots, 1\}$, when $\mu = 0.3$. The ground truth number of communities is 33. Each row corresponds to a $p^-$ value, starting from $p^- = 0$ on the first row, and $p^- = 0.4$ on the last row.
Fig. 20. (Color online) NMI, number of communities, and weighted NMI for all the combinations of $p^+ = \{0, \ldots, 1\}$ and $p^- = \{0, \ldots, 1\}$, when $\mu = 0.3$. The ground truth number of communities is 33. Each row corresponds to a $p^-$ value, starting from $p^- = 0.5$ on the first row, and $p^- = 1$ on the last row.
Fig. 20. (Continued)
Fig. 21. (Color online) NMI, number of communities, and weighted NMI for all the combinations of $p^+ = \{0, \ldots, 1\}$ and $p^- = \{0, \ldots, 1\}$, when $\mu = 0.4$. The ground truth number of communities is 32. Each row corresponds to a $p^-$ value, starting from $p^- = 0$ on the first row, and $p^- = 0.4$ on the last row.
Fig. 22. (Color online) NMI, number of communities, and weighted NMI for all the combinations of $p^+ = \{0, \ldots, 1\}$ and $p^- = \{0, \ldots, 1\}$, when $\mu = 0.4$. The ground truth number of communities is 32. Each row corresponds to a $p^-$ value, starting from $p^- = 0.5$ on the first row, and $p^- = 1$ on the last row.
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Fig. 22. (Continued)
Fig. 23. (Color online) NMI, number of communities, and weighted NMI for all the combinations of $p^+ = \{0, \ldots, 1\}$ and $p^- = \{0, \ldots, 1\}$, when $\mu = 0.5$. The ground truth number of communities is 32. Each row corresponds to a $p^-$ value, starting from $p^- = 0$ on the first row, and $p^- = 0.4$ on the last row.
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Fig. 24. (Color online) NMI, number of communities, and weighted NMI for all the combinations of $p^+ = \{0, \ldots, 1\}$ and $p^- = \{0, \ldots, 1\}$, when $\mu = 0.5$. The ground truth number of communities is 32. Each row corresponds to a $p^-$ value, starting from $p^- = 0.5$ on the first row, and $p^- = 1$ on the last row.
Fig. 24. (Continued)
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Table 1. Comparison between SN-MOGA and MEA<sub>s-SN</sub> on a network of 10000 nodes.

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>NMI</th>
<th>WNMI</th>
<th>NC</th>
<th>NS</th>
</tr>
</thead>
<tbody>
<tr>
<td>SN-MOGA min F(C)</td>
<td>0.6766</td>
<td>0.5376</td>
<td>123</td>
<td>0</td>
</tr>
<tr>
<td>SN-MOGA max Q&lt;sub&gt;S&lt;/sub&gt;</td>
<td>0.6766</td>
<td>0.5376</td>
<td>123</td>
<td>0</td>
</tr>
<tr>
<td>MEA&lt;sub&gt;s-SN&lt;/sub&gt; Liu et al.</td>
<td>0.9916</td>
<td>0.3797</td>
<td>196</td>
<td>64</td>
</tr>
</tbody>
</table>

In such a case the algorithm is not able to find any community structure, since it obtains almost 1000 groups, which is the number of nodes, i.e. it finds singleton communities. Nevertheless, the NMI is around 0.6. As \( \mu \) increases, the number of communities obtained by MEA<sub>s-SN</sub> increases too, while SN-MOGA is stable, finding, on average, a number of communities close to the number of the ground-truth division. Thus, while the WNMI values of MEA<sub>s-SN</sub> are drastically lower than the corresponding NMI values, the differences between the NMI and WNMI values that SN-MOGA obtains are minimal. The behavior of MEA<sub>s-SN</sub> is exacerbated as \( \mu \) increases. On the contrary, SN-MOGA continues to be rather stable as regards the average number of communities it obtains, with close NMI and WNMI values, proportionally decreasing, as expected, when \( \mu \) augments.

The figures also point out that the NMI value obtained by MEA<sub>s-SN</sub> is above 0.8 in almost all the experiments. Thus MEA<sub>s-SN</sub> outperforms SN-MOGA, even if it splits the network in many groups of small size, often singleton. When computing the weighted NMI, however, the values obtained by SN-MOGA slightly diminish with respect to NMI, and are always higher than that obtained by MEA<sub>s-SN</sub>. In fact, the WNMI values of MEA<sub>s-SN</sub> drastically reduce, due to the too high number of communities it obtains. This experiment highlights the characteristic of the weighted NMI to better discriminate solutions far from the true network division, by assigning them a lower and fairer value.

It is worth pointing out that a correlation analysis of the two objectives employed by MEA<sub>s-SN</sub> revealed a positive Pearson correlation value of 0.0517. According to the observation of Shi et al.<sup>8</sup> the multiobjective method becomes, actually, a single objective community detection method. Thus SN-MOGA effectively exploits the multiobjective approach by trying to obtain the best tradeoff between the two objective functions.

In order to more deeply investigate the differences between SN-MOGA and MEA<sub>s-SN</sub>, both methods have been executed on a synthetic network of 10000 nodes, with average node degree 64, exponent of degree distribution \(-2\), community size distribution \(-1\), mixing parameter \( \mu = 0.1, p_+ = p_- = 0.5 \). The number of clusters of the ground truth division is 100. Table 1 reports the NMI and weighted NMI (in the table denoted as WNMI), the number of clusters NC and the number of singletons NS obtained by the two methods. The behavior of MEA<sub>s-SN</sub> on this network is similar to the previous experimentation, i.e. it has the tendency of finding many clusters constituted by a single node, in this case it obtains 64
Table 2. Error obtained by SN-MOGA, MEAₜ-s-SN and Chiang’s method on the Wikipedia network, having number of nodes 7118, number of positive edges $E^+ = 83953$, and negative edges $E^- = 23118$.

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Error</th>
<th>NC</th>
</tr>
</thead>
<tbody>
<tr>
<td>SN-MOGA min $F(C)$</td>
<td>(0.0009868)</td>
<td>106</td>
</tr>
<tr>
<td></td>
<td>(0.00005728)</td>
<td></td>
</tr>
<tr>
<td>SN-MOGA max $Q_S$</td>
<td>0.0016</td>
<td>115</td>
</tr>
<tr>
<td></td>
<td>(0.0001142)</td>
<td></td>
</tr>
<tr>
<td>MEAₜ-s-SN Liu et al.²⁰</td>
<td>0.0020</td>
<td>3341</td>
</tr>
<tr>
<td></td>
<td>(0.0001609)</td>
<td></td>
</tr>
<tr>
<td>k-way Chiang et al.¹⁶</td>
<td>0.2186</td>
<td>3-30</td>
</tr>
</tbody>
</table>

Note: The error has been computed as Chiang et al.¹⁶

6.3. Comparison on Wikipedia network

In this section we consider a real life signed network, namely the English Wikipedia network for admin elections, studied by Leskovec et al.¹⁹ downloadable from http://konect.uni-koblenz.de/networks/elec. The network is constituted by 7118 nodes, and has number of positive edges $E^+ = 83953$, and negative edges $E^- = 23118$. This network has also been tested by Chiang et al.¹⁶ by applying their k-way multilevel algorithm. In Ref. 16 the authors reported the error they computed by applying formula (13), where the denominator, however, is substituted by $n^2$, i.e. the square of the number of nodes. We executed 10 times both SN-MOGA and MEAₜ-s-SN on this network and in Table 2 the error, computed like Chiang et al., obtained by the two methods, with the standard deviation in parenthesis, and that obtained by Chiang et al.,¹⁶ are reported. Moreover, also the average number $NC$ of obtained clusters is shown. For SN-MOGA two results are shown: when the solution having minimum frustration $F(C)$ and maximum modularity $Q_S$ are chosen from the Pareto front. The table points out that SN-MOGA obtains lower errors
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Fig. 25. (Color online) Synthetic networks reported in Yang et al. For each network, colors correspond to the true partitioning, red dotted lines to the negative edges and black lines to the positive edges.

and standard deviations, both when minimum frustration and maximum modularity are chosen. In the former case the error value is the lowest, being 0.0009868. MEA_{SN} on this network is not able to detect meaningful groups of nodes. In fact, it finds 3341 communities, where 3303 are singleton nodes. Thus it does not assign almost half of the nodes to any community. The number of clusters found by SN-MOGA, instead, has been, on average, 106 for minimum frustration, and 115 for maximum modularity.

As regards the method of Chiang et al., since the number of clusters must be given as input parameter, the authors computed the empirical error for values of $k$ ranging from 3 to 30. The value they reported is 0.2186, which is much higher than that obtained by SN-MOGA. They observed that, for each $k$, the errors are very close. Since SN-MOGA finds around 100 clusters, the range of values used by Chiang et al. was perhaps insufficient to obtain a reasonable partitioning of the Wikipedia network. This result confirms the advantage of applying SN-MOGA, which is capable of finding meaningful divisions with small frustration values, without any knowledge on the network structure and no need of fixing the number of communities in advance.

7. Comparison with Particle Swarm Optimization

In this section a comparison between SN-MOGA and the Particle Swarm Optimization based method of Gong et al., on two artificial signed networks and two real-life networks analyzed by Yang et al., is presented.

The two artificial signed networks, illustrated in Fig. 25, show the difference between balanced and partitionable networks. The network Network 1, consisting of 28 nodes, 30 positive edges and 12 negative edges and displayed in Fig. 25(a), is partitionable and can be divided into the three groups \{4, 5, 6, 7, 22, 23, 24, 25, 13, 14, 15, 16\}, \{8, 9, 26, 27, 17, 18\}, and \{20, 21, 10, 11, 12, 1, 2, 3, 19, 28\}. Network 2, having 28 nodes, 30 positive edges and 19 negative edges (Fig. 25(b)),
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Fig. 26. (Color online) Networks representing the political alliances and oppositions among 16 Gahuku-Gama subtribes (left) and the Slovene Parliamentary Party (right). Colors correspond to the true partitioning, red dotted lines to negative edges and black solid lines to positive edges.

Table 3. Comparison between \textit{SN-MOGA} and the \textit{MODPSO} method of Gong et al.

<table>
<thead>
<tr>
<th>Network</th>
<th>Algorithm</th>
<th>Modularity</th>
<th>NMI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Network 1</td>
<td>\textit{SN-MOGA} min $F(C)$</td>
<td>0.5612 (0.5612)</td>
<td>1 (1)</td>
</tr>
<tr>
<td></td>
<td>\textit{SN-MOGA} max $Q_S$</td>
<td>0.5612 (0.5612)</td>
<td>1 (1)</td>
</tr>
<tr>
<td></td>
<td>\textit{MODPSO}</td>
<td>0.5213 (0.5112)</td>
<td>1 (0.9742)</td>
</tr>
<tr>
<td>Network 2</td>
<td>\textit{SN-MOGA} min $F(C)$</td>
<td>0.5257 (0.5257)</td>
<td>1 (1)</td>
</tr>
<tr>
<td></td>
<td>\textit{SN-MOGA} max $Q_S$</td>
<td>0.5257 (0.5257)</td>
<td>1 (1)</td>
</tr>
<tr>
<td></td>
<td>\textit{MODPSO}</td>
<td>0.5643 (0.5634)</td>
<td>1 (0.9959)</td>
</tr>
<tr>
<td>GGS</td>
<td>\textit{SN-MOGA} min $F(C)$</td>
<td>0.4310 (0.4310)</td>
<td>1 (1)</td>
</tr>
<tr>
<td></td>
<td>\textit{SN-MOGA} max $Q_S$</td>
<td>0.4310 (0.4310)</td>
<td>1 (1)</td>
</tr>
<tr>
<td></td>
<td>\textit{MODPSO}</td>
<td>0.4310 (0.4310)</td>
<td>1 (1)</td>
</tr>
<tr>
<td>SPP</td>
<td>\textit{SN-MOGA} min $F(C)$</td>
<td>0.4556 (0.4556)</td>
<td>1 (1)</td>
</tr>
<tr>
<td></td>
<td>\textit{SN-MOGA} max $Q_S$</td>
<td>0.4556 (0.4556)</td>
<td>1 (1)</td>
</tr>
<tr>
<td></td>
<td>\textit{MODPSO}</td>
<td>0.4547 (0.4532)</td>
<td>1 (0.9949)</td>
</tr>
</tbody>
</table>

is also partitionable in the same three groups of \textit{Network 1}. The main difference between these two networks is that the former is also balanced since it has a two-way partitioning constituted by the first group and the union of the other two groups, while the latter is not balanced.

The \textit{Gahuku-Gama Subtribes} (GGS) social network (Fig. 26(a)) describes the political alliances (29 positive edges) and oppositions (29 negative edges) among 16 sub-tribes. The Slovene Parliamentary Party (SPP) network (Figure 26(b)) shows the relation among 10 parties of the Slovene Parliament in 1994. It has 18 positive edges and 27 negative edges. Table 3, for each network, shows the maximum modularity and NMI values obtained by \textit{MODPSO}, as reported in Gong et al.,\textsuperscript{9} while for \textit{SN-MOGA} the modularity and NMI values when the solution having minimum frustration is chosen from the Pareto front, and the solution with maximum modularity is chosen from the Pareto front with the corresponding NMI value. Average
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values for both the methods are in parenthesis. The table points out that both methods obtain the ground truth division for all the networks. However, MODPSO does not find the best solution for all the executions, as average values in parenthesis show. Thus SN-MOGA has a more stable behavior than MODPSO. Moreover, the modularity values obtained by SN-MOGA are higher on Network 1 and SPP, while lower on Network 2 and the same on GGS. Thus we can observe that both methods are able to properly divide these networks.

8. Running Time Analysis

One of the main criticisms to evolutionary based methods is the high execution time required to obtain a solution. However, it is known that genetic algorithms are naturally parallelizable. Since SN-MOGA has been written in Matlab, we could exploit the Parallel Computing Toolbox implemented in Matlab to allow multicore processing. We executed SN-MOGA on a computer cluster of 32 nodes, with 4 Gbyte of RAM and a 16-core Intel Xeon CPU at 2.6 GHz each.

In order to show the drastic reduction of execution times that can be obtained when the network size is large, Fig. 14 shows the time in seconds required by SN-MOGA to find a solution for the Wikipedia network (recall it has 7118 nodes, 83953 positive edges, and 23118 negative edges), when the number of cores used varies as 1, 2, 4, 8, 16, and 32. Population size and number of generations have been fixed to 100.

The figure points out that SN-MOGA presents a superlinear speedup when using two cores instead of one. In fact the running time reduces from 5 hours and a half to almost 2 hours. Moreover, the speedup is linear from 2 to 16 cores, since doubling the number of cores, the time required to execute the method becomes the half, and almost linear for 32 cores. In this latter case 20 minutes required with 16 cores reduce to 14 minutes on 32 cores. This experiment shows that SN-MOGA has a very good scalability. Thus, having at disposal sufficient computational resources, the method is able to deal with networks of very large size.

9. Conclusions

The paper proposed a multiobjective approach to detect communities in signed networks. The method optimizes two objectives in order to find network divisions such that intra-connections are dense and most edges within clusters are positive, and inter-connections between clusters are sparse, and most of these edges are negative. In order to evaluate the method, selection bias of the most used evaluation measures, namely the normalized mutual information, has been pointed out, and a corrected measure, the Weighted NMI, that avoids this bias adopted. An extensive experimental evaluation on randomly generated networks for which the ground-truth division is known, proved the ability of the method to find solutions having low frustration and high NMI and WNMI values. Furthermore, community structure found on the real life network Wikipedia showed that the error obtained by
SN-MOGA is lower than that obtained by MEA_{s-SN} and the k-way method of Chiang et al. A comparison with the MODPSO method, based on Particle Swarm Optimization, showed that the two methods are comparable, though SN-MOGA has a more stable behavior than MODPSO. Because of the genetic representation, the method cannot assign a node to multiple communities, thus generating overlapped community structures. Future work will investigate extensions to locus-based representation to obtain overlapping communities.

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