A Genetic Algorithm for Identifying Overlapping Communities in Social Networks Using an Optimized Search Space

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ABSTRACT

There are currently many approaches to identify the community structure of a network, but relatively few specific to detect overlapping community structures. Likewise, there are few networks with ground truth overlapping nodes. For this reason, we introduce a new network, Pilgrim, with known overlapping nodes, and a new genetic algorithm for detecting such nodes. Pilgrim is comprised of a variety of structures including two communities with dense overlap, which is common in real social structures. This study initially explores the potential of the community detection algorithm LabelRank for consistent overlap detection; however, the deterministic nature of this algorithm restricts it to very few candidate solutions. Therefore, we propose a genetic algorithm using a restricted edge-based clustering technique to detect overlapping communities by maximizing an efficient overlapping modularity function. The proposed restriction to the edge-based representation precludes the possibility of disjoint communities, thereby, dramatically reducing the search space and decreasing the number of generations required to produce an optimal solution. A tunable parameter $r$ allows the strictness of the definition of overlap to be adjusted allowing for refinement in the number of identified overlapping nodes. Our method, tested on several real social networks, yields results comparable to the most effective overlapping community detection algorithms to date.

Keywords: Overlapping Community Detection; Genetic Algorithm; Social Networks

1. Introduction

Many complex systems, such as social networks, autonomous systems, and html structures, can be usefully represented as graphs of edges and nodes. Communities, defined as subsets of nodes with more internal edges than external edges, reveal interrelated subgroups within network structures. Information on these community structures can provide useful information about systems themselves. A variety of methods for detecting these communities have been proposed. Unfortunately, most community detection algorithms treat communities as discrete sets of nodes, not allowing for overlap. However, this classification does not represent the actual structure of many complex systems; in social networks for example, many people belong to more than one social group.

Some community detection algorithms have been adapted to find overlapping community structures. Label Propagation Algorithm, or LPA, for example was modified into the Speaker-Listener Label Propagation Algorithm, or SLPA to increase accuracy and enable overlap detection. By assigning communities based on a belonging threshold, instead of by strongest belonging, nodes can be part of more than one community [1]. Other algorithms, such as Community Overlap Propagation Algorithm (COPRA) [2], Greedy Clique Expansion (GCE) [3], and Order Statistics Local Optimization Method (OSLOM) [4] use various methods to determine overlapping memberships.

COPRA uses LPA as a basis for its overlapping detection. Instead of assigning each node a single label based on its neighbor’s labels, COPRA assigns each node a belonging coefficient for each community. Whenever a coefficient falls below a threshold it is removed and the remaining coefficients are normalized. If the node does not have any memberships greater than this threshold, it maintains only the coefficient with the highest belonging. The nodes that retain multiple labels at the termination of the algorithm are overlapping nodes.

GCE uses maximal cliques as seeds for detecting
community structure. A maximal clique is defined as a group of nodes that is entirely connected, and is not part of a larger set of entirely connected nodes. The algorithm begins with as many communities as there are maximal cliques that are larger than the parameter k and greedily expands each, absorbing nodes that maximize a local fitness function. All communities that are too similar to one another are discarded. Because each community expands independently they can absorb the same nodes; these nodes are the overlap between two or more communities.

OSLOM works by finding the smallest significant communities in the network and building up a hierarchical structure. In each iteration, the smallest significant communities are detected. In the following iteration each of these communities is treated as a node, with the connections between communities refactored into weighted edges. The process is repeated until the entire network is joined together. Again, like in GCE, overlapping nodes may be identified because each community is detected independently, and the optimal structures for two communities might both contain a single node.

These algorithms demonstrate the variety of methods employed to detect overlapping community structure. Unfortunately there are only a few networks with known overlapping structure on which these algorithms may be compared. As a substitute, social networks with unknown overlapping structure may be used along with overlapping fitness functions to make a rough comparison in the same way that networks with unknown ground truths can still be analyzed using modularity. Our study introduces a new community detection technique based on genetic algorithms, and evaluates it on networks with and without known overlapping structures, producing both high modularity scores and accurate detection of known overlapping nodes.

2. Networks

This study makes use of a total of seven real-world social networks: karate [5], lesmis [6], dolphins [7], books, football [8], jazz [9], and Pilgrim. Karate is a well-known friendship network of a karate club in the 1970’s. Lesmis represents the co-appearance of characters in Victor Hugo’s Les Miserables. Dolphins are a collection of the associations of dolphins studied off the coast of New Zealand. Books represent the co-purchasing of political books on Amazon at the time of the 2004 US presidential election. Football is composed of the games played by Division IA college football teams in 2000. Jazz represents the collaboration network of a selection of Jazz musicians. Finally Pilgrim is a newly introduced network built on the social interactions of a high school class.

The Pilgrim high school network is composed of 34 nodes and 128 edges based on the friendship structure of a single class. It was created by two members of the class, Dickinson and Lieb, from experience in different communities within the class. The density of the graph is not uniform, resulting in a variety of different community structures. Some portions of the graph are very sparsely connected, while others form cliques. There is also a heavy amount of overlap between two of the larger communities presenting an interesting problem for algorithms designed to produce overlapping partitions. The basic ground truth community structure of this network is known, as is the overlapping community structure. Because it is a relatively small network, the overlap may be easily represented and studied as in Figure 1.

3. Modularity

Probably the most widely used measure for evaluating the goodness of a community structure without a ground truth is Newman’s modularity function [10]. Modularity is a metric of the difference in the internal density of the community structures before and after a random rearrangement of the edges of the graph (1). The full definition of the function has been omitted because of its common use. This function is only compatible with disjoint communities in a graph. There are, however, two variations of modularity designed specifically for evaluating overlapping community structures.

\[
Q = \frac{1}{2m} \sum_{i,j} \delta(c_i,c_j) \left[ A_{ij} - \frac{k_i k_j}{2m} \right]
\]

In [11], Shen introduced \( EQ \), an adaptation of Newman’s modularity function designed to support overlapping communities. The equation for \( EQ \) strongly resem-
bles the original modularity function (2) (1). In this equation, \( m \) is defined as the number of edges in the graph. \( C \) is the set of communities, and \( O_i \) is the number of communities to which the node \( v \) belongs. The presence of an edge between two nodes \( v \) and \( w \) is represented as the value in the corresponding position of the adjacency matrix \( A_{vw} \). This equation reduces to the same value as modularity when each node is a member of only one community.

\[
EQ = \frac{1}{2m} \sum_{i \in C} \sum_{j \in C} \frac{1}{O_i O_j} \left[ A_{ij} - \frac{k_i k_j}{2m} \right]
\]  

Another metric proposed in [12] by Nicosia provides a more complex and potentially more accurate evaluation of the goodness of an overlapping community structure (3). This measure makes use of several additional functions defined in [12]. For its arbitrary function \( f = 2px - p \), the recommended \( p \) value of 30 has been adopted. As in \( EQ \), \( C \) is the set of communities, and \( V \) is the set of nodes. In addition to this, \( r_{ijc} \) and \( s_{ijc} \) are weights based on the proportion of belonging to a community. A full explanation of them is available in [12]. The time complexity of this fitness function is almost \( O(n^3) \) making it unsuitable for use inside of an algorithm.

\[
Q_{ov} = \frac{1}{m} \sum_{c \in C} \sum_{j \in V} \left[ r_{ijc} A_{ij} - s_{ijc} \frac{k_i^{ov} k_j^{ov}}{m} \right]
\]  

4. Algorithms

In this section, we present two different approaches to overlapping community detection. The first is LabelRank, a deterministic variation of SLPA based on the properties of matrix multiplication. The second is a variation of genetic algorithms designed to provide a large variety of potential solutions. The first approach is more efficient, while the second is more thorough.

4.1. LabelRank

LabelRank was proposed by Xie as a stabilization of SLPA [13]. Its results compare favorably with SLPA with the additional benefit of being deterministic. LabelRank works in four phases: propagation, inflation, cutoff, and conditional update. The first phase, propagation, generates a new probability matrix, \( P' \), by multiplying the current probability matrix by the adjacency matrix of the graph. In the initialization of \( P' \), each node has an equal probability of belonging to each of its immediate neighbors. The next stage, inflation, accelerates the process of gravitating nodes toward communities by increasing high probabilities while simultaneously decreasing low ones. This function takes a value of \( P' \) and sets it equal to \( \frac{P'}{P_i} \) where \( \alpha \) is the chosen inflation parameter, and \( P_i \) is a row in \( P \). Once the values of \( P' \) have been inflated, the cutoff phase sets all values smaller than some threshold \( r \in [0, 1] \) equal to 0. This reduces the amount of memory necessary to maintain the matrix without having an effect on the results of the algorithm. The final phase is the conditional update in which only nodes meeting certain conditions have their rows of the probability matrix altered. This function requires that the set of most probable labels for node \( i \) be a subset of the most probable label sets of a certain threshold \( q \) of its neighbors. A conditional update is used to prevent the quality of the partition from decreasing after it reaches its peak. The algorithm terminates when the number of nodes changing their labels stabilizes.

Because LabelRank produced similar, and often better, performance when compared to SLPA, it showed promise for being adapted to detect overlapping communities. In fact, the only necessary modification was the introduction of a proportion of membership threshold \( a \). After termination instead of assigning each node its maximum probability label, each node is assigned every label for which it has a probability greater than \( a \). If no probabilities are greater than \( a \), then the maximal probability label is allotted. This simple modification was designed to enable LabelRank to detect overlap in the same way that SLPA does.

4.2. Overlap-Detection Genetic Algorithm

LabelRank has the distinct property of being deterministic. Because our initial testing showed that in many cases this could be a disadvantage, a new algorithm for detecting overlapping communities was developed based on previous genetic algorithms. The genetic algorithm for overlap detection, abbreviated OGA, unlike LabelRank provides a variety of potential solutions. It requires several parameters defined as follows: a specified number of iterations \( iter \), a proportion of the population to maintain as the fittest \( f \), a crossover rate \( q \), a mutation rate \( mut \), and a membership threshold \( r \). This algorithm uses an edge-based representation for clustering in order to detect overlapping communities of nodes [14]. In order to reduce the search space to likely partitions, a similar representation to that of [15] is used. The gene is the same length as the edge set of the graph. Each index represents an edge, and the corresponding value also represents an edge. The pairing of two edges in the gene indicates that they share a community. The pseudo code for OGA is presented in Algorithm 1.

Algorithm 1: OGA

Input: graph, popSize, iter, f, q, mut, r
# n is the number of nodes in the graph
# A is the adjacency matrix of the graph

main()
population \leftarrow \text{matrix}(n, \text{popSize})

\textbf{for each} row in population:
\hspace{1em}row = \text{generate\_individual}()
\textbf{end for}
\textbf{for} i \text{ in } 1 \text{ to } \text{iter}:
\hspace{1em}fittest \leftarrow \text{evaluate(population, } f)\hspace{1em}
\hspace{1.5em}\text{population} \leftarrow \text{crossover(fittest, } q)\hspace{1em}
\hspace{1.5em}\text{population} \leftarrow \text{mutate(population, } mut)\hspace{1em}
\textbf{end for}

\text{generate\_individual}()
\hspace{1em}\textbf{for} i \text{ in } 1 \text{ to } n:
\hspace{2em}\text{individual}[i] \leftarrow \text{random(neighbors(graph, } i))\hspace{2em}
\textbf{end for}
\textbf{return} \text{individual}

\text{evaluate(population, } f)
\hspace{1em}\text{population} \leftarrow \text{sort(population, fitness)}\hspace{1em}
\hspace{1em}\text{fittest} \leftarrow \text{list()}
\hspace{1em}\textbf{for} i \text{ in } 1 \text{ to } \text{popSize} \times f:
\hspace{2em}\text{fittest.add(population}[i])\hspace{2em}
\textbf{end for}
\textbf{return} \text{fittest}

\text{crossover(fittest, } q)
\hspace{1em}\textbf{for each} individual in population:
\hspace{2em}\text{father} \leftarrow \text{random(fittest)}\hspace{2em}
\hspace{2em}\text{mother} \leftarrow \text{random(fittest)}\hspace{2em}
\hspace{1em}\textbf{for} i \text{ in } 1 \text{ to } n:
\hspace{2em}\text{if} \text{random(0 to 1)} > q:
\hspace{3em}\text{individual}[i] \leftarrow \text{father}[i]\hspace{3em}
\hspace{2em}\text{else}
\hspace{3em}\text{individual}[i] \leftarrow \text{mother}[i]\hspace{3em}
\hspace{2em}\textbf{end if}
\hspace{2em}\textbf{end for}
\hspace{1em}\textbf{return} \text{population}

\text{mutate(population, } mut)
\hspace{1em}\textbf{for each} individual in population:
\hspace{2em}\textbf{for} i \text{ in } 1 \text{ to } n:
\hspace{3em}\text{if} \text{random(0 to 1)} < mut:
\hspace{4em}\text{individual}[i] \leftarrow \text{random(neighbors(graph, } i))
\hspace{2em}\textbf{end if}
\hspace{2em}\textbf{end for}
\hspace{1em}\textbf{return} \text{population}

The first step of OGA is to generate a random, safe population. In this context, safe means that an edge pairing can only take place between adjacent edges in the network. In generating the random safe population the value at each position of the gene is selected randomly from the edge sets of the two nodes connected to that edge. In doing so, the total search space for the algorithm is greatly reduced without removing possibly-maximal solutions. An example of this structure in relation to a graph is given in Figure 2.

The next several operations of the algorithm are repeated for the predefined number of iterations, iter. First the fittest members of the population are selected out of the population. This requires a fitness function through which to compare genes, and a percentage threshold to determine the size of the fittest. The chosen fitness function is the faster of the two overlapping modularity functions, $\text{EQ}$. Results should be improved by maximizing one of the measures of the goodness of overlapping partitions.

In order for this function to be useable, however, the membership must be converted to a node-based format. The transformation is done using a recursive tracing function. Because each edge in the representation is associated only with one other edge, these connections must be traced outward from a starting edge to discover the entire community. If the edges were treated as nodes, and their associations taken from the gene as edges, the edge communities would be the disjoint sub graphs of the network as seen in Figure 3. This provides a non-overlapping community partition of the edges. In order to determine the required node membership, a simple percentage is calculated. The membership to each community is the number of labels matching that community in the edge set of the node divided by the degree of that node. If this value is higher than some threshold $r$ then the node belongs to that community. Nodes without any memberships greater than $r$ are assigned to the community to which they have the highest membership. For example, in the sample graph node four would belong to

\begin{figure}
\centering
\includegraphics[width=0.8\textwidth]{example_graph.png}
\caption{(a) Example graph; (b) A safe gene for the network in (a) for example 3 implies a pairing of edges 3 and 1 and places them in the same community. A quick check verifies that they are adjacent via node 2 so this is a safe connection. The same is true of every index-value pair in this gene since it is a safe gene.}
\end{figure}
both communities because half of its edges are in each community Figure 2.

The next step of OGA is to generate a new population from the fittest by selecting two parents and crossing them over. The crossover is performed using a crossover rate $q$. For each position in the gene, the child has a $q$ probability of receiving the value from the first parent, and a $1 - q$ probability of receiving it from the second. Because both parents were initialized with safe genes, the child is necessarily safe. If the value exists at a position in the parent, then the two edges must share a node.

Crossover is followed by a repaired mutation [15]. A normal mutation, in which random values are assigned to randomly selected positions in genes, would explore unlikely solutions and make genes unsafe. Instead, the value at any edge randomly selected according to the mutation rate is set to the index of an adjacent edge. In doing this, the population remains safe, and the search space does not expand unnecessarily. Following this step, the process is repeated for the specified number of iterations, after which the fittest gene of the entire process is returned.

The approximate time complexity of the initialization, crossover, and mutation functions is $O(m*popSize)$. Because the search space is greatly reduced by the safe representation used, the number of iterations required scales linearly as well, keeping the time complexity below $O(n^2)$. The only remaining bottleneck on the runtime is the fitness function itself, which in this case has a time complexity of $O(n^2)$. Therefore the overall runtime of OGA is $O(n^2)$.

5. Results

In this section, we first experiment with our adaptation of LabelRank as an overlapping community detection algorithm before concluding that the efficiency gains due to its deterministic nature were less important than the maximal accuracy which was lost. In response to this, we present the results of our proposed overlapping community detection algorithm alongside SLPA across several real world social networks using $EQ$ and $Qov$ modularity scores as our evaluation criteria. To further demonstrate the performance of our algorithm, and the flexibility provided by its $r$ parameter, we compare our identified overlapping communities to those of the ground truth on the Pilgrim network.

5.1. LabelRank on Overlapping Communities

The modified LabelRank algorithm was tested on our selected networks against SLPA. SLPA was chosen as a baseline for comparison because of its affinity with LabelRank, and its excellent performance in comparison with other algorithms for detecting overlapping community structures [16]. Both algorithms were tuned on a full range of parameters. LabelRank used the following parameter ranges: $in \in \{1,5,12\}$, $q \in \{0.5,0.6\}$, $r \in \{0.1\}$, and $\alpha \in \{0.1,0.15,0.2,0.25,0.3\}$ . SLPA was run with varying numbers of iterations $in \in \{25,50,75,100\}$, and various thresholds $\alpha \in \{0.1,0.15,0.2,0.25,0.3\}$ . During the tuning phase, SLPA was run ten times with each possible combination of its parameters on every graph. The parameter set that resulted in the highest overlapping modularity values for each graph was then used to produce a final result. Using these parameters, SLPA was run nine more times recording the minimum, median, and maximum modularity scores for each modularity function. LabelRank too was run using its optimal parameters to produce the modularity scores recorded in Tables 1-3.

From these results, it appears that LabelRank generally performs close to the median modularity of SLPA. These results are illustrated in Figures 4 and 5. When LabelRank is optimized to perform its function with as little unnecessary overhead as possible, its runtime is linear [13]. This is approximately the same runtime as SLPA [1]. Since both algorithms have approximately the same

| Table 1. LabelRank and SLPA modularity. |
|-----------------|-----------------|-----------------|-----------------|
| LabelRank       | SLPA min         | SLPA median     | SLPA max        |
| karate          | 0.371466         | 0.28709         | 0.37147         |
| Pilgrim         | 0.437805         | 0.08859         | 0.36905         |
| lesmis          | 0.444812         | 0.23090         | 0.526688        |
| dolphin         | 0.491515         | 0.33794         | 0.47076         |
| books           | 0.492297         | 0.44860         | 0.49359         |
| football        | 0.603070         | 0.58062         | 0.59874         |
| jazz            | 0.281997         | 0.28100         | 0.33084         |
runtime LabelRank is better suited for detecting overlapping partitions on large networks. This is because the result of LabelRank is deterministically approximately that of the median SLPA score. It would likely take several runs of SLPA to produce a higher score, and doing so would take several times as much processing time and power.

This does not change the fact that SLPA can produce higher modularity scores given more opportunities to run. Since LabelRank simulates the spread of labels strictly according to probabilities it is unsurprising that it results in scores near the median SLPA scores. Because SLPA allows these probabilities to act on the spread of labels randomly it produces some variation from that median point. Many of these variations will influence the quality of the partition negatively, but some will have a positive influence on the result allowing SLPA to produce higher maximal scores. SLPA is able to perform better over multiple runs because it has an enlarged search space which generates a pool of candidates from which to choose the preferred solution.

### 5.2. OGA in Overlapping Community Detection

Because the larger selection of candidate solutions in SLPA produced higher optimal results, we implemented OGA, a genetic algorithm which maximizes $EQ$. By enlarging the search space to include all possibilities in which communities are not disjoint, we hoped to create better overlapping partitions. OGA uses several tunable parameters, the most important of which we discovered was $r$, the percentage of edges belonging to a given community a node needs to be considered part of that community. Other parameters such as population size, proportion considered the fittest, and number of iterations remained relatively constant. The optimal value for $r$ for every graph tested was 0.2. Higher values tended to produce too little, if any, overlap, while lower values resulted in the detection of too many overlapping nodes. The results of our genetic algorithm are comparable to

**Table 2. LabelRank and SLPA $EQ$**

<table>
<thead>
<tr>
<th>Graph</th>
<th>LabelRank</th>
<th>SLPA min</th>
<th>SLPA median</th>
<th>SLPA max</th>
</tr>
</thead>
<tbody>
<tr>
<td>karate</td>
<td>0.415516</td>
<td>0.340155</td>
<td>0.410092</td>
<td>0.450362</td>
</tr>
<tr>
<td>Pilgrim</td>
<td>0.391510</td>
<td>0.246385</td>
<td>0.543144</td>
<td>0.570444</td>
</tr>
<tr>
<td>lesmis</td>
<td>0.451008</td>
<td>0.369536</td>
<td>0.504252</td>
<td></td>
</tr>
<tr>
<td>dolphins</td>
<td>0.421700</td>
<td>0.494279</td>
<td>0.61182</td>
<td></td>
</tr>
<tr>
<td>books</td>
<td>0.489592</td>
<td>0.500836</td>
<td>0.508527</td>
<td></td>
</tr>
<tr>
<td>football</td>
<td>0.607479</td>
<td>0.594251</td>
<td>0.61182</td>
<td></td>
</tr>
<tr>
<td>jazz</td>
<td>0.290611</td>
<td>0.286760</td>
<td>0.300419</td>
<td>0.446170</td>
</tr>
</tbody>
</table>

**Table 3. LabelRank and SLPA $Q_{ov}$**

<table>
<thead>
<tr>
<th>Graph</th>
<th>LabelRank</th>
<th>SLPA min</th>
<th>SLPA median</th>
<th>SLPA max</th>
</tr>
</thead>
<tbody>
<tr>
<td>karate</td>
<td>0.634439</td>
<td>0.503739</td>
<td>0.632735</td>
<td>0.667816</td>
</tr>
<tr>
<td>Pilgrim</td>
<td>0.648807</td>
<td>0.257710</td>
<td>0.604430</td>
<td>0.731975</td>
</tr>
<tr>
<td>lesmis</td>
<td>0.646621</td>
<td>0.131625</td>
<td>0.748242</td>
<td>0.768513</td>
</tr>
<tr>
<td>dolphins</td>
<td>0.613071</td>
<td>0.470771</td>
<td>0.697240</td>
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</tr>
<tr>
<td>books</td>
<td>0.692267</td>
<td>0.693832</td>
<td>0.741700</td>
<td>0.756844</td>
</tr>
<tr>
<td>football</td>
<td>0.692091</td>
<td>0.595320</td>
<td>0.683301</td>
<td>0.715176</td>
</tr>
<tr>
<td>jazz</td>
<td>0.332284</td>
<td>0.345375</td>
<td>0.496795</td>
<td>0.712619</td>
</tr>
</tbody>
</table>

![Figure 4. LabelRank and SLPA modularity scores on karate.](image1)

![Figure 5. LabelRank and SLPA modularity scores on football.](image2)
the highest results of SLPA after 9 runs as can be seen in 

*Table 4* and *Figures 6* and 7.

The overlapping modularity scores of OGA are very close to those of SLPA on every network. This implies a particularly good partitioning considering [16] determined that SLPA is one of the most consistently accurate algorithms for detecting overlapping community structure. Of the fourteen algorithms tested in [16] SLPA was among the top performers on each network.

Analyzing the identified overlap in the Pilgrim network yields interesting results. The only detected overlapping node in the optimal modularity solution for Pilgrim was 21. In the next run, with a slightly lowered \( r \) value OGA produced a slightly lower modularity result, but a much more accurate list of overlapping nodes. This second list included the following nodes: 5, 15, 24, 21, and 22. Of these 5 nodes, 3 are overlapping in the ground truth, while the other 2 are very close to the community borders. When the \( r \) parameter was reduced slightly further, it produced a larger list of overlapping nodes, including only 2 additional nodes that are not quite overlapping. All of these results are displayed in *Figure 8*. The colors, red, orange, and yellow, correspond to the lowest \( r \) value 0.2, 0.1, and 0.08, under which a given nodes was categorized. Outline colors show the ground truth of the network.

Further analysis of these results was done using the F-score to evaluate the detection of overlapping nodes. A correctly identified overlapping node is a true positive, or \( TP \). Similarly a node incorrectly identified as overlapping is a false positive or \( FP \). Inversely nodes not identified as overlapping which in fact are constitute false negatives, \( FN \), and nodes correctly identified as not overlapping are true negatives, \( TN \). These counts are used to calculate precision and recall which are \( \frac{TP}{TP + FP} \) and \( \frac{TP}{TP + FN} \), respectively. Precision then represents the percentage of nodes classified as overlapping which are in fact overlapping, and recall is the percentage of ground truth overlapping nodes identified. These two metrics may be combined into an F-score,

\[
2 \times \frac{\text{precision} \times \text{recall}}{\text{precision} + \text{recall}}
\]

producing scores between 0 and 1 where 1 is the optimal result. This metric balances the optimization between two goals, discovering all overlapping nodes and only discovering truly overlapping nodes. The relationship

<table>
<thead>
<tr>
<th>Modularity</th>
<th>Algorithm</th>
<th>karate</th>
<th>Pilgrim</th>
<th>lesmis</th>
<th>dolphins</th>
<th>football</th>
</tr>
</thead>
<tbody>
<tr>
<td>( EQ )</td>
<td>OGA</td>
<td>0.445225</td>
<td>0.472687</td>
<td>0.578773</td>
<td>0.540742</td>
<td>0.603323</td>
</tr>
<tr>
<td></td>
<td>SLPA</td>
<td>0.450362</td>
<td>0.474091</td>
<td>0.570444</td>
<td>0.504252</td>
<td>0.611182</td>
</tr>
<tr>
<td>( Q_{ov} )</td>
<td>OGA</td>
<td>0.637640</td>
<td>0.735629</td>
<td>0.746658</td>
<td>0.752494</td>
<td>0.712163</td>
</tr>
<tr>
<td></td>
<td>SLPA</td>
<td>0.667816</td>
<td>0.731975</td>
<td>0.768513</td>
<td>0.744916</td>
<td>0.715176</td>
</tr>
</tbody>
</table>

*Figure 6. OGA and SLPA \( EQ \) on several real social networks.*

*Figure 7. OGA and SLPA \( Q_{ov} \) on several real social networks.*
6. Conclusion

This paper has presented a genetic algorithm for overlapping community detection, OGA, comparable to SLPA, one of the best existing algorithms, in performance. Using a restricted representation to limit the number of possible partitions, the time complexity is reduced to $O(n^2)$. In addition to this the genetic structure of OGA provides flexibility to introduce new fitness functions. The $EQ$ and $Q_{ov}$ scores of this algorithm are on par with SLPA, one of the premier algorithms for detecting overlapping community structure in social networks. Comparing this discovered overlap with the ground truth overlapping structure of Pilgrim reveals some disparity between optimal modularity and ground truth community structures. This demonstrates the usefulness of networks with ground truth overlapping structure for the evaluation of community detection algorithms.

7. Acknowledgements

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