A mathematical programming approach for sequential clustering of dynamic networks

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Abstract. A common analysis performed on dynamic networks is community structure detection, a challenging problem that aims to track the temporal evolution of network modules. An emerging area in this field is evolutionary clustering, where the community structure of a network snapshot is identified by taking into account both its current state as well as previous time points. Based on this concept, we have developed a mixed integer non-linear programming (MINLP) model, SeqMod, that sequentially clusters each snapshot of a dynamic network. The modularity metric is used to determine the quality of community structure of the current snapshot and the historical cost is accounted for by optimising the number of node pairs co-clustered at the previous time point that remain so in the current snapshot partition. Our method is tested on social networks of interactions among high school students, college students and members of the Brazilian Congress. We show that, for an adequate parameter setting, our algorithm detects the classes that these students belong more accurately than partitioning each time step individually or by partitioning the aggregated snapshots. Our method also detects drastic discontinuities in interaction patterns across network snapshots. Finally, we present comparative results with similar community detection methods for time-dependent networks from the literature. Overall, we illustrate the applicability of mathematical programming as a flexible, adaptable and systematic approach for these community detection problems.

1 Introduction

Complex networks exhibit several topological features that distinguish them from more simple networks such as lattices and random networks. One feature in particular is the tendency of nodes to organise themselves into a modular topology, known as community structure [1]. The detection of such communities, also known as modules, is widely accepted as means of revealing the relationship between topological and functional features of complex systems [2].

Network representations of complex systems are often static, corresponding to either a snapshot of a system at a certain point in time or an aggregation of data over multiple time points. However, in reality networks are not static; nodes and interactions can be created or cease to exist. For example, in social networks friendships are made and broken. In a business, employees retire and new members of staff are employed. In biological systems, not all interactions take place at the same time, depending upon spatial, temporal or environmental conditions [3].

A dynamic network is defined as a series of network snapshots at two or more time points, where time can represent seconds, days, years or various states of a system. The changes which occur at the node and interaction level may affect global measures and descriptions of the network e.g. community structure as modules are created, destroyed, split into multiple groups or merged together. Consequently, incorporating temporal information into network modelling frameworks may lead to more accurate representations of complex systems. It follows that a current challenge in community structure detection is the identification of modules in dynamic networks.

Community structure in relation to dynamic networks has been tackled in various ways. Consensus clustering attempts to find a partition of a system that is to some extent relevant at each time step [4–6]. Alternatively, many approaches cluster the static snapshot networks independently and employ various methods of comparison between partitions to quantify change in community structure or follow the evolution of communities [7–10]. In particular, some methods aim to detect drastic discontinuities...
in community structure which represent some form of important ‘event’ \cite{11,12}.

Where snapshots are clustered individually, historic community structure is not taken into account. It has been proposed, however, that the community structure of the network at time \( t \) should not be taken as independent of the community structure of the network at time \( t - 1 \). In other words, a network at time \( t \) is clustered with respect to a known partition of the network at \( t - 1 \). Such methods take advantage of information about the community structure of previous snapshots in order to infer the structure in the current time step, as expressed by evolutionary clustering approaches.

The first evolutionary clustering method introduced the idea of temporal smoothness, where the snapshot quality measure (e.g. modularity) is maximised at the current time point and a distance measure, the history cost (e.g. mutual information, rand index, etc.) between the current snapshot and the previous snapshot is minimised \cite{13}. A trade-off is therefore made between remaining faithful to the current data, but minimising the variation between the current partition and the previous one. Similar approaches can be seen in references \cite{14–21}. In this study, we present a mathematical model that inherits this framework and our analysis focuses on data that is sequential in nature.

Mathematical programming provides a flexible and intuitive modelling framework that has been shown to be competitive in numerous community detection algorithms \cite{6,22–29}. In our previous work we have proposed mathematical programming methods based on modularity optimisation to identify hard partitions \cite{22,24,27}, overlapping communities \cite{29} and to cluster multiplex networks \cite{6}. Here, we extend our previous work and propose a mathematical programming approach to evolutionary clustering. In particular, we report a mixed integer nonlinear programming (MINLP) model that given a series of network snapshots, returns a partition for each of the snapshots, taking into account historical information. The applicability of our method is demonstrated through its application to synthetic and real networks and through a comparative analysis with similar methods from the literature.

2 Methods

2.1 A mathematical programming model for sequentially clustering snapshots of dynamic networks

Here we report an MINLP model of evolutionary clustering that sequentially identifies the community structure of each snapshot in a dynamic network. The indices, parameters and variables associated with our model, known as SeqMod, are given below:

Indices

\( n, e \) nodes (the union from all input snapshots)
\( m \) modules

\( t \) time step

Parameters

\( \beta_{net} \) weight of link between \( n \) and \( e \) at time \( t \);
\( d_{nt} \) weighted degree (strength) of node \( n \) at time \( t \);
\( L_t \) sum of the weights in the network at time \( t \);
\( \gamma_{ne,t} \) equals to 1 if nodes \( n \) and \( e \) are in the same community at time \( t \);
\( \varepsilon \) the smoothness control, a user parameter that indicates the weights given to the preservation coefficient and modularity.

Binary Variables

\( Y_{nmt} \) equal to 1 if node \( n \) is in module \( m \) at time \( t \); 0 otherwise.

Continuous Variables

\( D_{mt} \) the sum of the weighted degree of nodes in module \( m \) at time \( t \);
\( L_{mt} \) the sum of the weights of links that are in module \( m \) at time \( t \).

Modularity expresses how well-defined the community structure of a network is reference \cite{30}. The metric provides an intuitive description of community structure and is one of the most popular methods for community structure detection. We thus employ modularity to determine the community structure of the current snapshot:

\[
Q_t = \sum_m \left( \frac{L_{mt}}{L_t} - \left( \frac{D_{mt}}{2L_t} \right)^2 \right), \quad \forall t, \tag{1}
\]

where \( L_{mt} \) is the sum of weights of links within module \( m \) at time \( t \):

\[
L_{mt} = \sum_{n,e} n > e \beta_{net} Y_{nmt} Y_{emet} \quad \forall m, t, \tag{2}
\]

and \( D_{mt} \) is the sum of weighted degrees of nodes in module \( m \) at time \( t \):

\[
D_{mt} = \sum_n d_{nt} Y_{nmt} \quad \forall m, t. \tag{3}
\]

The central idea in Evolutionary Clustering \cite{13} is to detect community structure that is consistent with current data and tracks changes smoothly over time, i.e., the community structure does not change drastically from a time step to another. This usually means that the community structure at a specific time step \( t \) should reflect the data at \( t \) as well as the community structure at the immediate previous time step. Here, however, we take a different reference time step \( t^* \) that has had the maximum modularity up to the current time step and we define a preservation coefficient, \( \Delta_t \), to measure the number of pairs of nodes that are co-clustered both at \( t^* \) and at the current time step \( t \):

\[
\Delta_t = \sum_m \sum_{n,e} n > e \gamma_{ne,t^*} Y_{nmt} Y_{emet} \quad \forall t. \tag{4}
\]
This preservation coefficient represents how similar two partitions are to each other. $\Delta t$ and $Q_t$ can be competing objectives and we define the parameter $\varepsilon$ for controlling the influence each metric has over the optimization process. This parameter is said to control the smoothness of the transitions, i.e. the influence of the historical clustering information has over the current network structure. The objective function is defined according to the definition of evolutionary clustering in reference [14]:

$$(1 - \varepsilon)Q_t + \varepsilon\Delta_t \quad \forall t.$$  \hfill (5)

Parameter $\varepsilon$ is restricted to the interval $[0, 1]$, such that when $\varepsilon = 0$, the partition at $t^*$ does not influence the clustering of the partition at the current snapshot, at $t$. If $\varepsilon = 1$, our model would simply maintain the previous partition and the modularity of the current snapshot would not be considered. This provides for a more intuitive setting of the expected smoothness of transitions.

Both $Q_t$ and $\Delta t$ are therefore normalised. $Q_t$ ranges from $-\frac{1}{2}$ to 1 [31] and $\Delta_t$ ranges from 0 (when no pairs of nodes were maintained from the previous time step to the current one) to $\Delta_{\text{max}} = \sum \gamma_{nc,t^*}$, when all pairs of co-clustered nodes from the reference time step remain together on a module at the current time step. Our objective function is therefore redefined as:

$$\frac{(1 - \varepsilon)}{3}(2Q_t + 1) + \frac{\varepsilon}{\sum \gamma_{nc,t^*}}\Delta_t.$$  \hfill (6)

Finally, SeqMod detects disjoint communities in each snapshot of the network, i.e. each node belongs to only one module at time $t$. Therefore we add the following constraint:

$$\sum_m Y_{nmt} = 1 \quad \forall n, t.$$  \hfill (7)

The complete model is formulated below:

$$\begin{align*}
\text{maximize} & \quad \frac{(1 - \varepsilon)}{3}(2Q_t + 1) + \frac{\varepsilon}{\sum \gamma_{nc,t^*}}\Delta_t \\
\text{subject to} & \quad (1, 2, 3, 4, 5) \\
& \quad L_{mt}, D_{mt} \geq 0 \quad \forall m, t \\
& \quad Y_{nmt} \in \{0, 1\} \quad \forall n, m, t.
\end{align*}$$

SeqMod is implemented in GAMS (General Algebraic Modelling System) [32] using standard branch and bound (SBB) method as the mixed integer optimisation solver and CONOPT as the NLP solver with default parameters. To ensure that we give a reasonable representation of solution space, for each clustering experiment (each with a different $\varepsilon$), the MINLP is solved iteratively 100 times, each time with a different random initial partition. After each time step is solved, the solution with the largest value of the objective function is selected. Even though an upper bound for the number of modules is provided, it is stressed that the actual number of modules in the partition is decided by the model.

### Table 1. Summary of networks used in this paper.

| Network                  | Nodes | Time steps | Classes |
|--------------------------|-------|------------|---------|
| Synthetic network        | 22    | 3          | 3       |
| High School              | 180   | 7          | 5       |
| MIT Social Evolution     | 61    | 36         | 8       |
| Brazilian Congress       | 589   | 12         | 2       |

### 2.2 Network data

SeqMod is tested on synthetic and real networks, summarised in Table 1. First, as an illustrative example, we have adapted a small synthetic network from [33]. This network comprises 22 nodes and 3 time steps. The network has 3 clearly defined communities at $t = 1$ but connections between two of them become increasingly denser at $t = 2$ and $t = 3$.

We have also tested real social networks, generated from proximity data among high school students, college students and members of the Brazilian Congress. The High School network represents the interactions among 180 high school students, over a period of 7 school days [34]. Students had to wear a device that would record any face to face contact with another student that lasted at least 20 s. The results were used to generate daily networks. The authors of the study have shown that about 91.5% of contacts made between students in this high school during the study involved students in the same class.

The third network was built from the MIT social evolution dataset [35], consisting of records of eighty MIT students who lived in the university dormitory during an academic year. Various types of interaction data (WiFi location, Bluetooth proximity, SMS and Calls exchanged) was collected using mobile phones that were given to the students. We selected Bluetooth proximity data to generate a dynamic network of weekly snapshots for 36 weeks. We restricted our network to the students who had their dormitory sector and year of studies mapped by the researchers and used only the records that had a probability of at least 20% that the involved students lived on the same dormitory floor. The final network, after applying these restrictions, had 61 nodes.

The fourth network was built from a dataset of the records of the roll call votes in the Brazilian Chamber of Deputies (the lower house of the Congress)\(^1\). To construct the networks, we first selected records in the period from 2003, the first year of the current ruling party government (PT) to 2014. Duplicated entries of each congressman were removed, results were ranked according to parties and records of the topmost four parties were selected. Two of these parties are in the opposition group (PSDB and DEM) while the other two are allied to the government (PT and PMDB). For every year in the records, we computed a weighted adjacency matrix based on [36].

\(^1\) The data was downloaded from https://github.com/estadadados/basometro
in which each cell represents the similarity of the votes of each pair of congressmen during that year. Unanimous vote sessions, i.e. cases where 95% of the present congressmen vote the same, were removed.

2.3 Alternative clustering methods

We compare our results with other algorithms for community detection through evolutionary clustering, namely Estrangement, FacetNet and DynMOGA. The Estrangement algorithm [19] solves the snapshots sequentially using modularity and a measure of dissimilarity between two partitions called estrangement, which is proportional to the number of intra-community edges that becomes inter community over time. FacetNet [16] is based on a stochastic block model for the detection of communities and a probabilistic model based on Dirichlet distribution that detects the evolution of the communities. DynMOGA is a multiobjective genetic algorithm that optimizes both modularity and normalized mutual information (with respect to the previous time step) [20].

We also compare our method to genLouvain [37], a robust algorithm with a modified version of modularity designed for multilayer networks. In order to obtain results that are comparable to an evolutionary clustering framework, we apply genLouvain sequentially at each time step, i.e. to detect the partitions for the network at \( t_1 \), we input the first snapshot; for the second time step, we provide \( t_1 \) and \( t_2 \) as input, and follow a similar procedure for all other ensuing time snapshots.

2.4 Adjusted rand index

In order to evaluate the accuracy of SeqMod in detecting the ground truth community structure, we employ the Adjusted Rand Index (ARI) metric [38], a measure of similarity between two partitions, comparing it to a null probabilistic model. ARI yields 1 for identical partitions and usually 0 for independent partitions, although it may also result in some negative values [39]. We used the mclust package implementation of ARI in R [40].

3 Results and discussion

3.1 Synthetic network

For illustrative purposes, we tested our model on a small synthetic network, shown in Figure 1. We note three clear distinct communities (one on the left and the other two on the top, and bottom right, respectively). The two rightmost communities have increasingly denser connections between them at \( t_2 \) and \( t_3 \). We investigate whether SeqMod can maintain the three communities over time and under what values of \( \epsilon \).

If we fix \( \epsilon = 0 \) for all time steps, the model does not consider historic information and each time step is solved sequentially considering only modularity. The result is shown in Figure 2. At \( t_1 \), three communities are detected, at \( t_2 \) the network is just slightly different, now the central node is placed in the red community instead of the blue. This reflects the minor changes in this snapshot, this node changed membership because at this time step it has more connections with the red community than with the others. At \( t_3 \), there are now more edges between the red and blue communities and the community structure is different from the previous time steps. The three nodes marked A, B and C on the plot either did not interact with any other node at \( t_2 \) or are not present in the network at that time step, in our model we treat such nodes as isolated nodes. Since no historic information was taken into account, these nodes are assigned each to a singleton community at that time step.

The results are different when \( \epsilon > 0 \), Figure 3 shows the partitions detected by SeqMod for the synthetic network for \( \epsilon \geq 0.10 \). Community structure is now maintained from \( t_1 \) to \( t_2 \) even though the network has changed and at \( t_3 \), SeqMod detects a merge between the red and blue communities. Note also that the absent nodes at \( t_2 \) are maintained in their original community (from \( t_1 \)). This is an advantage of the evolutionary clustering approach: it takes advantage of prior knowledge to infer the node community allocation even in the snapshot where it is not present/interacting.
Node A is a good example of a “promiscuous” node and its community membership is determined by the parameter $\varepsilon$. For example, if we fix $\varepsilon = 0.015$ the node is assigned to the green community at $t_1$ and is kept green at $t_2$, when it is absent, but on the third step it belongs to the blue community since now it has more ties with that group. The desired smoothness is controlled by the user parameter.

3.2 High school network

Our first real network is a proximity network created by aggregating information on face-to-face contacts between high school students. Students belong to one of five classes, each with a specific discipline programme: two MP classes, with a focus on mathematics and physics; two PC classes, with a focus on physics and chemistry and one PSI class, with focus on engineering. We take these classes as our ground truth communities and we investigate how closely our model detects them. Students are more likely to interact with others in similar disciplines than with the rest as previously noted by [34]. In all test cases where $\varepsilon > 0.05$ and $\varepsilon < 0.30$, this pattern occurs to some extent at these time steps; it can also be perceived from Figure 5 where for these values of $\varepsilon$, there is drastic change in interaction patterns at $t_3$ and $t_5$ (drop in ARI) that is followed by an increase in the metric for $\varepsilon > 0.30$. This seems to suggest that these classes were more involved in activities together during these days or even that these students are more likely to interact on Thursdays ($t_3$) and Fridays ($t_5$) than on Mondays ($t_1$, $t_6$), Tuesdays ($t_2$, $t_7$) and Wednesday ($t_3$).

3.3 MIT social evolution dataset

The network built from Bluetooth proximity records from the MIT Social Evolution dataset is smaller than the previous network in terms of nodes but represents a longer time period. The interactions of the students in the Social Evolution network are registered based on Bluetooth proximity data rather than face-to-face contacts, as in the High School network, so some of the records may actually correspond to “false interactions”. For example, if there is a record representing an interaction between students A and B, it may mean that they were in fact in different floors or room and not interacting at all. Therefore we restricted our network to those interactions with at least 20% of chance that they were in the same floor.

The dormitory sector where each student lived was reported to be the primary factor in defining the relationships between students [35], so we take the dormitory sectors as the ground truth communities for this network. Figure 6 shows the ARI results for various values of $\varepsilon$.

Once again, by incorporating historic information (any $0 < \varepsilon < 0.20$) the model yields results closer to the ground truth than clustering each snapshot individually. For $\varepsilon = 0.05$, the model improves in ARI over the results for $\varepsilon = 0.00$ and it still adapts to some major
changes in community structure in the network. There is a pattern of decrease in ARI during the intervals \(\{t_{12} - t_{16}, t_{19} - t_{21}, t_{26} - t_{28}\}\), the first of them correspond to the weeks of Christmas and New Year’s Eve Holidays and the network is considerably reduced during these time steps (about 80% of students were absent). This explains why the model is more capable of matching the ground truth for relatively larger \(\epsilon\) during the Holiday season: since the majority of the network consists of absent nodes, the community structure found previously is maintained.

We also note the trade-off nature of the smoothness control in the model. For this network, when \(\epsilon = 0.10\) the model yields the best result in terms of matching the true classes of the students (mean ARI = 0.371) but as it can be noticed in Figure 6, when solving with \(\epsilon = 0.05\) the model is better at detecting the changes in the network, ARI decreases during the holiday periods. One has to balance between the current state of the network, incorporating “ground truth” information sequentially, while also allowing for change. If \(\epsilon\) is set too large, for example 0.50, the model might incur “over-smoothing”, i.e. being unable to adapt to changes in the community structure on this period [42].

3.4 Brazilian congress voting dataset

The Brazilian Congress network consists of 589 nodes and 12 time steps. The ground truth used as validation for this dataset is the political alignment of the congressmen, based on their parties (either left-wing or right-wing). Figure 7 shows the ARI results found by SeqMod at each time step.
parameters were set as follows: we tested to a unique community label for comparison purposes. The tors of the same size, we have assigned each of these nodes sent nodes and ARI only compares two membership vec-

ment and FacetNet do not output information about ab-

changes in network structure, which is evident even in al-

gorithms without a smoothness parameter like FacetNet and DynMOGA. If the objective is to detect changes in the network, the smoothness parameter must be set in a way that allows for more adaptation. In SeqMod, this means that $\varepsilon$ must be set to a lower value as discussed

![Adjusted Rand Index](Image1)

Fig. 7. Brazilian congress network – the change in adjusted rand index (ARI) at each time step for SeqMod depending on the value of $\varepsilon$.

| Algorithm | High School | Social Evolution | Brazilian Congress |
|-----------|-------------|------------------|--------------------|
| SeqMod    | $0.8160$    | $0.3705$         | $0.3390$           |
| Estrangement | $0.5739$  | $0.3192$         | $0.1524$           |
| genLouvain | $0.6872$    | $0.3701$         | $0.3188$           |
| FacetNet  | $0.5907$    | $0.2429$         | $0.2434$           |
| DynMOGA   | $0.3962$    | $0.1611$         | $0.0287$           |

4 Comparative analyses

We compare SeqMod with the Estrangement, FacetNet, DynMOGA and genLouvain algorithms. Since Estrangement and FacetNet do not output information about absent nodes and ARI only compares two membership vectors of the same size, we have assigned each of these nodes to a unique community label for comparison purposes. The parameters were set as follows: we tested $\varepsilon$ in the range [0.00, 0.50], in intervals of 0.05 for SeqMod and the maximum number of communities was set to 10, 10 and 4 for the High School, MIT Social Evolution and Brazilian Congress network, respectively. For the Estrangement algorithm, the default set of parameters defined by the authors ($\delta = 0.00, 0.01, 0.025, 0.05$) were employed, plus additional parameters ($\delta = 0.1, 0.2, \ldots, 0.5$). For FacetNet a fixed number of communities should be specified, so we fixed it to the number of communities in the ground truth for each dataset. GenLouvain was tested for $\omega$ ranging from 0.00 to 0.50 in intervals of 0.05. Across all methods, deciding on which parameter to use for comparison is important, so we report the test cases that have yielded the largest average ARI (across all time steps) for each algorithm. Table 2 reports the best average ARI found by each algorithm and the parameters used to find these results.

Figure 8 shows the results on the High School network for the parameters reported in Table 2 at each time step. The algorithms seem to detect a similar pattern on the network from $t_3$ to $t_5$: they all show a decrease in ARI in this period and most algorithms show an increase in this metric during the last two time steps. SeqMod, however, provided the more accurate results, matching more closely the ground truth communities of this network through time (average ARI = 0.8160).

The best results for the MIT Social Evolution network are shown in Figure 9. SeqMod again shows the best performance overall in terms of ARI with $\varepsilon = 0.10$ followed closely by genLouvain. As discussed previously and shown in Figure 6, the results found with this parameter were the closest to the ground truth. It is noteworthy that tests with smaller $\varepsilon$ were able to show structural changes in the network during holiday periods $t_{13}$ – $t_{15}$ and around $t_{20}$ and $t_{27}$. These changes were detected by Estrangement and also by DynMOGA and FacetNet, although with less accuracy.

This fact once again highlights the existing balance between detecting the ground truth of a network and the changes in network structure, which is evident even in algorithms without a smoothness parameter like FacetNet and DynMOGA. If the objective is to detect changes in the network, the smoothness parameter must be set in a way that allows for more adaptation. In SeqMod, this means that $\varepsilon$ must be set to a lower value as discussed...
Algorithm

- Seqmod $\varepsilon = 0.15$
- Estrangement $\delta = 0.00$
- genLouvain $\omega = 0.20$
- FacetNet $M = 5$
- DynMOGA

**Fig. 8.** High school network – the change in adjusted rand index (ARI) at each time step for each algorithm.

Algorithm

- Seqmod $\varepsilon = 0.10$
- Estrangement $\delta = 0.10$
- genLouvain $\omega = 0.30$
- FacetNet $M = 8$
- DynMOGA

**Fig. 9.** MIT social evolution network – the change in adjusted rand index (ARI) at each time step for each algorithm.

before. Estrangement algorithm and SeqMod with $\varepsilon = 0.05$ exhibit the more drastic discontinuities in the network.

The tests with Brazilian Congress network show similar results (Fig. 10). SeqMod achieves the best accuracy of algorithms, followed by genLouvain. Both algorithms exhibit a sudden increase in ARI from $t_4$ to $t_5$ and from $t_8$ to $t_9$, corresponding to the years of elections and renewal of congressmen in the Chamber of Deputies in Brazil. A decrease in ARI is observed for most algorithms from $t_{11}$ to $t_{12}$ and since the ground truth is the alignment of the parties, we conjecture that this change reflects the changes in political alignment of the parties due to the recent political crisis in Brazil that started in 2014 when a major corruption scandal was revealed [43].

**5 Conclusions**

In this paper, we proposed an alternative approach to sequential clustering of evolving networks using mathematical programming. The proposed method is an extension of our previous work to consider historic information when detecting community structure of dynamic networks, under evolutionary clustering framework. We validated our model on dynamic networks with known ground truth communities and compared it to other methods who also employ both historic information and modularity optimisation.

Our model uses a different concept of similarity where a preservation coefficient allows for an intuitive understanding of the number of vertex pairs that were maintained in the network. We also introduce the concept of a “reference snapshot”, instead of using the immediate previous time step in the historical cost, we use the previous time step with the largest modularity. The reason is that we want to maximize the ability of our model to detect the ground truth of communities through time, comparing each snapshot with a previous better defined community structure.

We have shown that our model is able to detect the ground truth of dynamic networks and changes in the network community structure. We have shown with our illustrative example that our model is able to incorporate historic information to detect probable changes in the network and that it also maintains the information, shedding light on the membership of the nodes that do not interact during all time steps, i.e. that are absent or
isolated. Determining the desired smoothness is the main challenge of evolutionary clustering and consequently of our model, but from the results obtained in this study, we suggest a value around 0.10 and 0.15. Some methods automatically determine the parameter at each time step based on probabilistic models and statistical features of the network while algorithms like DynMOGA model this process as a multiobjective problem and hence do not require any parameters, but as our tests have shown, these algorithms found poor results. We intend to improve our mathematical programming model in the future to adapt to the changes in the network without the need of parameter setting, while also providing solutions of suitable quality.

Mathematical programming provides a flexible environment for modelling community detection and here we have illustrated its use for evolutionary clustering. We note that this type of modelling framework is particularly adaptable and can accommodate various user requirements, one can for example specify a module allocation for determined nodes if known, one can add constraints that change the way isolated nodes are treated or any other logical conditions necessary for a practical application. Therefore, we believe that such approaches will prove to be popular in future network analysis methodologies and applications.

Author contribution statement

JCS, LGP, LB, ST conceived and designed the experiments. JCS, LGP, LB, ST designed the computational model. JCS, LB performed the experiments. JCS gathered network data, JCS, LGP, LB, ST analysed the data. JCS, LB, ST, LGP wrote the paper.

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