Graph-Aware Evolutionary Algorithms for Influence Maximization

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ABSTRACT
Social networks represent nowadays in many contexts the main source of information transmission and the way opinions and actions are influenced. For instance, generic advertisements are way less powerful than suggestions from our contacts. However, this process hugely depends on the influence of people who disseminate these suggestions. Therefore modern marketing often involves paying some targeted users, or influencers, for advertising products or ideas. Finding the set of nodes in a social network that lead to the highest information spread—the so-called Influence Maximization (IM) problem—is therefore a pressing question and as such it has recently attracted a great research interest. In particular, several approaches based on Evolutionary Algorithms (EAs) have been proposed, although they are known to scale poorly with the graph size. In this paper, we tackle this limitation in two ways. Firstly, we use approximate fitness functions to speed up the EA. Secondly, we include into the EA various graph-aware mechanisms, such as smart initialization, custom mutations and node filtering, to facilitate the EA convergence. Our experiments show that the proposed modifications allow to obtain a relevant runtime gain and also improve, in some cases, the spread results.

CCS CONCEPTS
• Human-centered computing → Social networks; Social network analysis; • Theory of computation → Evolutionary algorithms; • Mathematics of computing → Graph algorithms.

KEYWORDS
Social Network, Influence Maximization, Evolutionary Algorithm, Graph Theory, Combinatorial Optimization.

1 INTRODUCTION
Online social networks have become a relevant part in the daily life of the majority of people. Social networks are used every day to maintain contact with other people, read news, search for a job, watch movies, or do shopping. All these things are usually done based on what other people say about products, jobs, events, and so on. More formally, when the activity of a user $v$ implies an action of the user $u$, there is an edge $u \rightarrow v$ in the graph representing the social network. This edge might indicate, for example, a voter supporting a candidate for an election, or the fact that if a customer $u$ buys a product, the same product will be bought by the user $v$. In general, the notation $u \rightarrow v$ indicates that information spreads from $u$ to $v$ with a certain probability. In case of success, $v$ is said to be activated by $u$. As we will see later, different probabilistic models, called diffusion models, are now well-established tools to study how influence propagates over social networks.

2 BACKGROUND
As said earlier, the IM problem consists in finding the set of nodes that will lead to the maximal number of node activations according to a given influence spread model. More specifically, the problem can be formulated as follows: given a network, represented with a graph $G$, a diffusion model $m$, and a budget $k$, find a set of $k$ initially active nodes, such that the influence spread is maximized. Here, we use the Independent Cascade (IC) and Weighted Cascade (WC) models [20]. In IC, at each timestep $t$ a node can be either active or inactive. A node can transit from inactive to active, but it cannot become inactive again. At the beginning, only the nodes in the seed set $S$ are in the active state. Then, at each timestep each active node $u$ has a chance of activating its neighboring nodes $v$ (if the edge $u \rightarrow v$ exists) with a probability $p$, which is the same for all the nodes in the graph and is a given parameter. The simulation stops
once a convergence condition is met, see Algorithm 1 in Section 4.1. Influence spread is thus given by the number of active nodes at the end of the simulation. In WC, influence spread is calculated in the same way as in IC, the only difference being the probability of node activation: given a node \( v \), the probability of being activated by one of its neighboring nodes \( u \) is \( 1/(\text{indegree}(v)) \).

3 RELATED WORKS

The existing approaches for solving the IM problem can be divided into two main categories: approximation algorithms with provable guarantees, and metaheuristics. The former provide good results, but their long runtime make them impractical on large-scale networks. The latter compromise runtime with approximate solutions, although they suffer from curse of dimensionality. We provide below an overview of the related works from both categories.

3.1 Approximation algorithms

The IM problem was formulated for the first time as a combinatorial optimization problem in [20], and proven to be NP-hard. The authors also proposed a greedy algorithm that yields an \( (1 - 1/e - \epsilon) \) approximation. The submodular property of the objective function was later used in the CELF algorithm [23], producing a 700x speedup w.r.t. a greedy approach. This property was further explored in CELF++ [13], with an additional 35-55% gain in runtime.

More recent algorithms, namely TIM and TIM+ [27], and its improved version IMM [26], use a different approach to calculate the nodes with the maximum spread, based on Reverse Reachable (RR) sets. Each RR set is obtained by taking the set of nodes reached by the spread from one node (one for each RR set) and by putting them in an hypergraph as an hyperedge. The solution is then calculated iteratively by selecting the node with the highest degree in the hypergraph and removing then that node with all its incident edges. The difference between the three algorithms consists in the ways of calculating the sufficient number of RR sets and the method used to sample nodes for their generation. All three algorithms outperform both CELF and CELF++, while IMM outperforms both TIM and TIM+. A further improvement of IMM is BCT [16], specifically designed for cost-aware targeted viral marketing, where each node has a benefit and a cost associated to it. BCT uses this information in order to first sample nodes with higher benefit for creation of the RR sets, resulting in an algorithm significantly faster than IMM.

3.2 Metaheuristics

The first application of EAs to the IM problem was proposed in [3], in which a basic EA without any domain knowledge outperformed the greedy algorithm [20] in terms of runtime while obtaining comparable influence spread results. Since that, a plethora of improvements were done in this direction. In [29], for instance, a GPU-parallelized EA outperformed the greedy algorithm given the same runtime, taking up to 34 times less time to archive the same result. The impact of the EA parameters on the IM problem was studied instead in [30]. Another recent study [10] investigated the effect of two selection schemes used for solving the IM problem with EAs, and found that the \((\mu, \lambda)\) scheme performs better than \((\mu + \lambda)\) when the size of the seed set increases.

Other works have investigated the use of smart initialization of the initial EA population. In [31], different initialization strategies were compared, with PageRank-based initialization [25] giving the worst results. A smart initialization approach was also adopted in [11], that unlike standard high-degree initialization can guarantee a higher diversity of the solutions. In the same work, the authors used 2-hop spread [22] as approximated fitness function, and a similarity-based local search. Their results were quite promising, with a 10x speedup w.r.t. CELF and comparable influence results. In another work, the same authors proposed a discrete Particle Swarm Optimization algorithm (DPSO) [12], which made use of 2-hop spread approximation and a smart initialization based on degree discount. DPSO outperformed CELF++ in terms of runtime, while the influence results were similar. Smart initialization techniques were also studied in [21], where each node has a probability to be inserted into the initial population proportional to its degree, and [9], producing better results than the basic EA.

Custom genetic operators were used in [24], where an informed mutation operator was introduced, based on a neural network predicting which nodes to change in a solution using centrality metrics as inputs. In [32], the EA premature convergence problem was handled using multi-population competition with specific crossover and mutation operations among populations. In other studies the IM problem was formulated instead as a multi-objective problem. In [15] a multi-objective differential evolution was used to maximize influence while minimizing the information delivery cost. Similarly, a multi-objective EA was used in [4] to maximize influence while minimizing the size of the seed set.

4 METHODS

We now explain the main modifications we applied to the basic EA proposed in [3]. This algorithm uses a direct encoding in which each individual genotype is simply a vector of IDs of nodes in the social network graph, i.e., a seed set. Initialization is done by randomly selecting a seed set of \( k \) nodes from the graph. At each generation, parents are selected by performing tournament selection, which then reproduce by means of one-point crossover operator. Differently from [3], the crossover operator adopted here has two additional constraints: 1) to produce solutions without node repetitions, and 2) to produce solutions different from those given in input, in order to improve diversity. To satisfy the former condition, we swap only nodes which are not in common between the two parents, while maintaining the other nodes in common. Regarding the latter property, we simply force a mutation of at least one random node in each child produced by crossover. This is then followed by a random mutation that simply resets each node, with a given probability, to one of the other possible node IDs (excluding those already present in the seed set). The new population is then obtained with generational replacement with elitism. The evolution is continued until a maximum number of generations is reached, or stopped earlier if there is not any improvement for \( \sigma \) consecutive generations, where \( \sigma \) was set to 10% of the maximum number of generations. In our experiments, we set \( \text{population\_size}=100, \text{generational\_budget}=100 \) generations, \( \text{crossover\_rate}=1.0, \text{mutation\_rate}=0.1, \text{tournament\_size}=5 \), and \( \text{num\_elites}=1 \). This parameterization follows the one used in [3],
to obtain a fair comparison with the basic EA. A specific parameter
tuning, as done in [30], is instead out of the scope of this work.

In order to improve the basic EA, we collected the best ideas
from the literature, discussed in the previous Section, and put them
in practice by adapting them to the algorithm. As we will see in the
next Section, some of them were more successful, while some others
hardly produced any improvement. Nevertheless, we report all of
our results in order to provide valuable knowledge on what are
the most promising directions. In a nutshell, the objectives of our
work are to: 1) reduce the runtime of the fitness function, the main
bottleneck of the EA, by using fitness function approximations; 2)
boost the EA convergence with smart initialization; 3) improve the
search efficiency by using graph-aware mutations, also including a
mechanism for dynamic selection of multiple mutations; 4) reduce
the EA search space by filtering the most promising nodes before
starting the search. Next, we discuss these four elements in detail.

4.1 Fitness function approximations

The influence spread computation under the IC and WC models
was proven to be #P-complete in [13], and as such it is one of
the main bottlenecks for the IM algorithms. This is usually calculated
by using Monte Carlo (MC) sampling. According to this procedure,
the influence spread is calculated by simulating the information
spread in the graph, using the specified diffusion model, a given
number of times (usually very large, > 10,000). The influence spread
is then approximated by the mean spread across the simulations.

As seen earlier, an alternative to the expensive MC sampling
is the use of RR sets [8]. A preliminary study of surrogate models
was instead conducted in [5], but the results were contrasting.
More promising solutions make use of spread function approxima-
tions, which are faster than MC sampling. This is the idea we have
decided to adopt in this work. In particular, we use two different
approximations. The first one is the 2-hop spread approximation
function introduced in [22], which as discussed earlier was used in
[11] because of its efficiency. According to this approximation, the
influence spread is given by:

$$
\hat{\sigma}_S = \hat{\sigma}_S \left( \sum_{s \in S} \sum_{c \in C_u \cap S} \sum_{d \in \text{neighbors}(c) \cap S} p(s, c) p(c, d) \right) - \chi,
$$

where $\chi = \sum_{s \in S} \sum_{c \in C_u \cap S} \sum_{d \in \text{neighbors}(c) \cap S} p(s, c) p(c, d)$, $\sigma_u^1$ is the one-
hop influence spread of node $u$, defined as $\sigma_u^1 = 1 + \sum_{c \in C_u} p(u, c)$,
and $C_u$ denotes the set of neighbors of node $u$.

The second approximation is a variation of MC sampling, called
MC max-hop. In this case, influence spread is propagated up to a maxi-
mum number of hops $\text{max}_\text{hop}$. If e.g. $\text{max}_\text{hop} = 2$, the influence
is propagated up to the neighbors of the neighbors of the seed set $S$.
The pseudo-code of the algorithm is shown in Algorithm 1, where
setting $\text{max}_\text{hop}$ to $\text{Inf}$ produces the original MC sampling.

4.2 Smart initialization

Introducing custom-built solutions into the initial population is a
common practice in Evolutionary Computation [6, 7, 18]. As we
have seen earlier, several works [9, 11, 12, 21, 31] have reported a
positive effect of smart initialization on the IM problem. Here we
consider different initialization techniques, described below.

Single smart solution: A single “smart” solution is inserted into
the initial population, that is generated by taking the first $k$ nodes
obtaining the highest centrality scores. We tested the following
centrality metrics:

- betweenness: measures how often the node is on the shortest
  path among any two nodes in the graph;
- closeness: corresponds to the mean length of the shortest
  path between the node and all the other nodes in the graph;
- degree: the number of out-going links of the node;
- eigenvector: nodes neighboring with few highly-connected nodes
  score higher;
- katz: a variant of the eigenvector metric, where the distance
  between the node and all the other nodes in the graph;

Multiple smart solutions: A percentage of the initial population
is initialized according to one of the following strategies:

- degree random: the probability of a node to be inserted into a
  solution is proportional to its degree;

\begin{algorithm}
   \KwResult{Monte Carlo max-hop simulation.}
   \textbf{input:} seed set $S$, seed set size $k$, graph $G$, social model $\text{model}$, number of simulations $\text{no}_\text{simulations}$, maximum number of hops $\text{max}_\text{hop}$
   \textbf{output:} spread mean value $\text{avg}$, spread standard deviation $\text{std}$
   \begin{enumerate}
      \item \textbf{procedure} \text{MONTECARLOMAXHOPSIMULATION} \\
         \hspace{1em} sample $\leftarrow$ initialize\_array($k$) \\
         \hspace{1em} $i \leftarrow 1$
      \item for $i \leq \text{no}_\text{simulations}$ do
      \hspace{1em} $A \leftarrow S$
      \hspace{1em} $B \leftarrow S$
      \hspace{1em} converged $\leftarrow$ False
      \hspace{1em} $j \leftarrow \text{max}_\text{hop}$
      \hspace{1em} while not converged and $j \leq \text{max}_\text{hop}$ do
      \hspace{1em} \hspace{1em} for $n$ in $B$ do
      \hspace{1em} \hspace{1em} \hspace{1em} neighbors $\leftarrow$ get\_neighbors($n$, $G$)
      \hspace{1em} \hspace{1em} \hspace{1em} not\_activated $\leftarrow$ set\_difference(neighbors, $A$)
      \hspace{1em} \hspace{1em} \hspace{1em} for $m$ in not\_activated do
      \hspace{1em} \hspace{1em} \hspace{1em} \hspace{1em} spread\_prop $\leftarrow$ get\_influence($n$, $m$, $\text{model}$)
      \hspace{1em} \hspace{1em} \hspace{1em} \hspace{1em} $p \leftarrow$ random\_real($0$, $1$)
      \hspace{1em} \hspace{1em} \hspace{1em} \hspace{1em} if $p >$ spread\_prop then
      \hspace{1em} \hspace{1em} \hspace{1em} \hspace{1em} \hspace{1em} add\_element\_to\_set($m$, $\text{nextB}$)
      \hspace{1em} \hspace{1em} \hspace{1em} $j \leftarrow j + 1$
      \hspace{1em} \hspace{1em} $A \leftarrow \text{union}(A, B)$
      \hspace{1em} \hspace{1em} $j \leftarrow j + 1$
      \hspace{1em} \hspace{1em} $\text{avg} \leftarrow \text{mean}(\text{sample})$
      \hspace{1em} \hspace{1em} $\text{std} \leftarrow \text{standard\_deviation}(\text{sample})$
      \item converged $\leftarrow$ True
   \end{enumerate}
\end{algorithm}
• degree random ranked: the probability of a node to be inserted into a solution is proportional to its ranking position w.r.t. the nodes’ degree;
• community degree: the general idea is that the network might have community structure and a good solution should contain the most influential nodes from the largest communities. Here, we use the Louvain algorithm [28] to perform community detection. The solutions are then built as follows: first, we select a community, with probability proportional to its size. Then, we select the node within the selected community, with probability proportional to its degree.

When using the initialization with both single and multiple smart solutions, the rest of the population is initialized randomly.

4.3 Graph-aware mutations
We attempt to accelerate the search process by introducing some knowledge about the node properties and graph structure into the mutation operator. In particular, we considered four different mutation schemes, described below.

Global mutation methods: We mutate the selected node to another node randomly chosen from the graph. Table 1 recapitulates the node selection techniques we use with the global mutation methods.

Local mutation methods: The idea is to perform a local search on the graph, i.e., to apply a mutation that preserves a certain proximity to the current solution (i.e., the seed set). In this case the node to be mutated is selected randomly, while the new node is chosen according to one of the criteria listed in Table 2.

Combination of multiple mutations: We have observed that some mutations are more effective than others when used in different evolution phases and when applied to networks with different degree distributions. Here, we model the dynamic selection of which mutation to use within a pool of available mutations as a non-stationary Multi Armed Bandit (MAB) problem. To solve the MAB problem, we adopt the Upper Confidence Bound (UCB1) algorithm [1], which promotes actions (in our case, mutations) with larger reward uncertainty. More precisely, to select the next action, the UCB1 algorithm maximizes the following expression:

\[ Q(a) + \text{exploration weight} \cdot \sqrt{\frac{2 \log(t)}{N_t(a)}} \]

where \( Q(a) \) is the cumulative reward of the last \( n \) times the action \( a \) was selected (\( n \) is a sliding window size), \( N_t(a) \) corresponds to the number of times \( a \) was chosen, and \( t \) is the counter of the total action selections. To shrink the learning phase, we adopted an exponentially decaying exploration weight \( \text{exploration weight}(g) = g^{-3} \), where \( g \) is the generation counter. Note that the sliding window is used since the problem is non-stationary, i.e., the reward of a certain mutation may change during the evolutionary process. This mechanism should then produce a fair trade-off between exploration and exploitation along the search.

4.4 Node filtering
The search space of the IM problem is combinatorial, therefore most existing algorithms scale poorly with the graph size. We investigate two ways to reduce the search space by filtering the nodes in the graph before applying the EA.

Min degree nodes: Nodes with low degree have a very little chance of influencing the others, so it is very unlikely that they would be part of the optimal solution. Here, we filter the \( n \) nodes with degree larger than a given threshold.

Best spread nodes: When considering large-scale networks, it is very likely that the optimal seed set would be made of nodes placed far enough from each other, to avoid activating the same nodes. According to this assumption, it is highly probable that the nodes in the optimal solution are isolated among them. In other words, their activations do not overlap and the resulting spread is very close to the sum of the spreads of each of the nodes calculated separately. To test this assumption, we first compute the spread generated by each node, separately, and then filter the best \( n \) nodes in terms of mean spread. This technique requires the spread measures of any two nodes to be comparable with each other, in order to determine which one is bigger. Furthermore, this technique is computationally expensive since it requires to perform MC simulations for all the nodes in the network, before running the EA. However, deciding a priori the required number of MC simulations needed to have a statistically significant comparison of the spread produced by all nodes (averaged across multiple MC simulations, as seen earlier) is not possible. Therefore, we use the following procedure that increases iteratively the number of MC simulations, until a needed precision is reached:

1. Initialize a node set \( N \) with all the nodes in the graph, and set the maximum error rate \( \text{max_error_rate} \) to a given value.

| Mutation | Description |
|-----------|-------------|
| Global random | The node to be mutated is selected randomly. |
| Global low degree | The mutation probability of a node is inversely proportional to its degree. |
| Global low spread | The mutation probability of a node is inversely proportional to its influence spread value. |
| Global low additional spread | The mutation probability of a node is inversely proportional to the increase in influence spread it produces when added to the seed set excluding it. |

| Mutation | Description |
|-----------|-------------|
| Local neighbors random | The new node is selected randomly among the neighbors of the node to be mutated. |
| Local neighbors second degree | The new node is selected among the neighbors of the node to be mutated, with probability proportional to its degree. |
| Local neighbors approximated spread | The new node is selected among the neighbors of the node to be mutated, with probability proportional to its approximated spread. |
| Local embeddings random | The new node is selected randomly among the nodes closest to the node to be mutated, according to its corresponding node2vec [14] embeddings (i.e., a continuous feature representation of the graph’s nodes, trained to predict the probability of nodes being neighbors). |
(2) Perform MC sampling of the spread of each node in $\mathcal{N}$ with a number of MC simulations (which might be different for each node) needed to ensure a maximum error rate $\text{max\_error\_rate}$ on the mean spread. The error of the mean is estimated by calculating the confidence interval of the Student’s $T$ distribution.

(3) Set $\mathcal{N}$ to the best $n$ nodes in terms of mean spread, plus all the nodes in the graph which are not statistically comparable with the node with the lowest spread among the best $n$ nodes. Decrease $\text{max\_error\_rate}$ and go to step 2.

This loop is repeated until all the best $n$ nodes are comparable with the others. However, the sample size required to compare any pair of nodes would be too costly, thus we stop the loop as soon as the number of best nodes falls in a given range $[l, u]$. More specifically, as soon as the algorithm finds a number of best nodes $l$, and the number of nodes incomparable with those $l$ nodes is lower than $u - l$, it stops. When the algorithm terminates, it returns the $l$ best nodes, plus all the nodes which cannot be compared with them.

| Table 3: Real-world datasets specifications. |
|---------------------------------------------|
| Dataset | Graph size | Node degree |
|---------|-------------|-------------|
| Wiki-vote | 7,115 nodes 103,689 edges | Avg 14.57 Std 42.28 Min 0 Max 893 Median 2 |
| Amazon | 262,111 nodes 1,234,877 edges | Avg 4.71 Std 0.95 Min 0 Max 5 Median 5 |
| CA-GrQc | 5,242 nodes 14,496 edges | Avg 5.52 Std 7.92 Min 1 Max 81 Median 3 |

| Table 4: Synthetic Barabási-Albert datasets specifications. The $n_{\text{edges}}$ parameter indicates the number of edges added from a new node to the existing nodes. |
|---------------------------------------------|
| Graph size | Node degree |
|-------------|-------------|
| 1,000 nodes $n_{\text{edges}}=1$ | Avg 1.99 Std 3.51 Min 1 Max 75 Median 1 |
| 1,000 nodes $n_{\text{edges}}=3$ | Avg 5.98 Std 7.37 Min 3 Max 99 Median 4 |
| 1,000 nodes $n_{\text{edges}}=5$ | Avg 9.95 Std 10.57 Min 2 Max 114 Median 7 |
| 1,000 nodes $n_{\text{edges}}=7$ | Avg 13.90 Std 12.80 Min 7 Max 137 Median 10 |
| 1,000 nodes $n_{\text{edges}}=9$ | Avg 17.84 Std 15.65 Min 9 Max 148 Median 12 |
| 1,000 nodes $n_{\text{edges}}=11$ | Avg 21.76 Std 19.25 Min 11 Max 211 Median 15 |
| 10,000 nodes $n_{\text{edges}}=1$ | Avg 1.99 Std 4.13 Min 1 Max 243 Median 1 |
| 10,000 nodes $n_{\text{edges}}=3$ | Avg 5.99 Std 8.96 Min 3 Max 293 Median 4 |
| 10,000 nodes $n_{\text{edges}}=5$ | Avg 9.99 Std 13.45 Min 2 Max 365 Median 7 |
| 10,000 nodes $n_{\text{edges}}=7$ | Avg 13.99 Std 17.00 Min 7 Max 446 Median 10 |
| 10,000 nodes $n_{\text{edges}}=9$ | Avg 17.98 Std 21.25 Min 9 Max 470 Median 12 |
| 10,000 nodes $n_{\text{edges}}=11$ | Avg 21.98 Std 25.97 Min 11 Max 695 Median 15 |

5 EXPERIMENTS

For the experiments, we used three real-world graphs taken from the SNAP repository [19], namely Wiki-Vote, Amazon and CA-GrQc. In Wiki-Vote the edges represent who-voted-whom information in elections for promoting adminship. The Amazon dataset contains co-purchased products relations. The smallest dataset, CA-GrQc, is a network of collaboration in the General Relativity and Quantum Cosmology fields. Table 3 contains a detailed description of each of the aforementioned graphs. For testing the function approximations, we also used 6 synthetic datasets generated by the Barabási-Albert model [2], detailed in Table 4. These were created using the networkx library [17] with a random seed parameter set to 0. Except for the function approximations experiments, where we set $p = 0.1$ in the IC model and max_hop = 2, in all the other experiments we set $p = 0.01$ and max_hop = 3. Each of the four proposed modifications presented above (fitness function approximations, smart initialization, graph-aware mutations, and node filtering) was tested separately. Every experimental condition was repeated for 10 independent runs (in the next figures, we provide mean values ± std. dev.)

5.1 Fitness function approximations

The first part of the experimentation was aimed at verifying which fitness function approximation works best in terms of runtime and quality of results. To do that, we compared the runtime and spread (mean and std. dev. across multiple simulations) computed by MC sampling and the two approximations, i.e., 2-hop spread and MC max-hop ($\text{max\_hop} = 2$), on the different datasets above, under both the IC and WC models. For each dataset and model, the three methods were executed on 100 randomly generated seed sets, the same for all the methods. The Pearson correlation on the spread values calculated by the proposed approximations and MC sampling was then calculated.

Generally, we observed that both approximations in some cases had an important gain in runtime and high spread correlations w.r.t. MC sampling. The correlation of the spread values is high for both approximations ($>0.9$) for the WC model, and for the IC model with low $p$ values ($p \leq 0.1$). On the other hand, we observed a dependency of the runtime on the size and connectivity of the graph, the size of the seed set $k$, and, of course, the number of simulations used for the MC methods. Figure 1 shows the runtime of the three methods on six selected datasets as the number of MC simulations varies (WC model, seed set size $k = 5$). Here we can observe the linear growth of the runtime of the MC methods w.r.t. the number of simulations. We can also notice that in some cases the 2-hop spread is slower not only when the number of MC simulations is low (somehow an obvious result), but also when the dataset size grows. Another thing which can be noticed is that MC max-hop is particularly useful as the dataset connectivity increases. See the Barabási-Albert graphs in the figure, where MC max-hop obtains a lower performance gain in the top row (lower connectivity) than in the bottom row (higher connectivity). We observed this trend in all our experiments: more connected graphs yield a longer influence spread process since usually more nodes get influenced, so a truncation of this process up to some level (as in MC max-hop) can lead to larger runtime gains.

We also observed that the runtime has, obviously, a linear relationship with the seed set size $k$. Figure 2 reports the runtime as a function of $k$ (WC model, number of MC simulations set to 100). Here we can observe that the 2-hop spread may be convenient only if the graph size is kept small, while for the real-world cases it might be the slowest spread evaluation method.

\footnote{Our code is available at: https://github.com/katerynak/Influence-Maximization.}
5.2 Smart initialization

We tested smart initialization only on the three real-world datasets. Overall, the results indicate that smart initialization is most useful in the case of datasets that contain single, highly connected nodes.
Single smart solution: We were not able to run complete experiments for the Amazon dataset, given their cost in memory and/or runtime (we did not consider centrality metrics with computation time > 12 hours). As shown in Figure 3, the degree metric is a clear winner for the Wiki-Vote dataset, which is reasonable considering its degree distribution. CA-GrQc experiments did not present a clear difference between various metrics under the IC model, while when considering the WC model the betweenness metric obtained slightly better results w.r.t. the other metrics. The only metric we could compute for the Amazon dataset was the degree metric, but this did not bring any improvement to the basic EA (results not shown for brevity).

Multiple smart solutions: Due to computing resource limitations, we could not run these experiments for different values of k. We used instead a fixed seed set size, k = 10, and a percentage of smart solutions of 50%. From Table 5, it can be seen that there is not a clear winner. On the other hand the degree random strategies are to be preferred because of the much lower computation time requirement w.r.t. the community degree strategy.

Table 5: Multiple smart solutions initialization techniques compared (for k = 10).

| Dataset   | Model | Best fitness (averaged across 10 runs) |
|-----------|-------|----------------------------------------|
|           |       | Degree | Comm. degree | Degree | No smart. init. |
|           |       | random | ranked | random | init. |
| Wiki-Vote | WC    | 226.24 | 223.35 | 230.65 | 220.75 |
| Wiki-Vote | IC    | 93.7   | 93.21  | 94.04  | 87.92  |
| Ca-GrQc   | WC    | 147.38 | 147.36 | 146.91 | 146.13 |
| Ca-GrQc   | IC    | 18.64  | 18.59  | 18.59  | 18.67  |
| Amazon    | WC    | 66.55  | 65.99  | 67.97  | 67.87  |
| Amazon    | IC    | 10.79  | 10.83  | 10.80  | 10.82  |

5.3 Graph-aware mutations

Also in this case we tested only the Wiki-vote, CA-GrQc and Amazon graphs. We omit for brevity the detailed results, but we report below our main findings.

Global mutations: The global low degree mutation obtained slightly better results on the graphs presenting a low number of large degree nodes, such as Wiki-Vote and CA-GrQc, while for the Amazon dataset it led to worse results. The runtime of these methods were in general comparable, except the global low additional spread method whose runtime grows linearly with the seed set size.

Local mutations: We report the results of the local mutations in Figure 4. For this specific experiment, we trained node2vec on the Amazon dataset and fine-tuned its parameters. As expected then, the local embeddings random mutation performed better on the Amazon dataset, while its results for CA-GrQc and Wiki-Vote were way worse than the other mutations: this shows that local embeddings may work well, but at a cost of an expensive and time-consuming tuning process. On the other hand, the local neighbors second degree mutation was at least as good as the global random mutation for all the datasets, while the local neighbors approximated spread and the local neighbors random mutation worked well on CA-GrQc and Wiki-Vote, but not on the Amazon dataset.

Combination of multiple mutations: We tested the UCB1 algorithm with all the mutations in Tables 1-2 and a sliding window size of 100. We noted that while the improvement w.r.t. the basic EA was not very significant, it was however positive for all graphs, which confirms that the dynamic selection of mutations adapts well to different cases (results not shown for brevity).

5.4 Node filtering

Filtering was tested only on the real-world graphs. Below we report our findings.

Min degree nodes: Min-degree values of 1, 2, 3 and 4 were tested. We obtained a visible improvement w.r.t. the basic EA only on the Wiki-Vote dataset, while in the other cases the high std. dev. did not permit to define a clear winner.

Best spread nodes: We used MC max-hop with max_hop = 2 to approximate the spread of each single node; max_error_rate was set initially to 0.8 and subsequently decreased of 0.1 at each iteration. We set l = 10^2 and u = 10^{11}, in order to have the same search space size regardless of k. From Figure 5, we can see that this method significantly improved the basic EA (up to 40%).

6 CONCLUSIONS

We improved a basic EA applied in previous research to the IM problem with various graph-aware enhancements aimed at reducing the algorithm runtime. Using MC max-hop instead of MC sampling to evaluate the influence spread permitted a significant computational time saving, and allowed the possibility to conduct a variety of experiments. The most important progress was achieved by limiting the search space of the EA by means of node filtering, a method which selected the most promising nodes in terms of information spread before running the evolutionary search: the limited number of node combinations allowed in fact the EA to better scale with the increasing graph size. Moreover, the combination of several local and global graph-aware mutations permitted the EA to adapt to graphs with different structure or connectivity.

There are, on the other hand, some limitations in the proposed mechanisms. For instance, MC max-hop may be not a good choice in case of networks with scarce connectivity or under the IC model with high probability values. Furthermore, node filtering is valid only under the assumption that the most influential nodes are distant, in the sense that they do not influence the same nodes. So this method may perform worse on small-scale networks where the optimal seed set may include nodes with common spread influence. Additionally, the runtime of the metrics which involve the computation of the influence spread, as well as the degree-based mutations, is proportional to the fitness of the solution. So, the runtime of the algorithm is proportional to the graph connectivity and a large number of fitter candidates found during the search may imply an increasingly high computation cost.

There is still further research to do on graph-aware mutations, which would probably work better with a small number of pre-filtered nodes. In particular, the success of the node2vec embeddings mutation (although with some hyper-parameter search) suggests
that this might be a promising direction to explore more in depth, also because differently from the other tested mutations this kind of mutation does not need node-specific expensive computations.
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