Abstract—Communication networks are prone to virus and worms spreading and cascading failures. Recently, a number of social networking worms have spread over public Web sites. Another example is error propagation in routing tables, such as in BGP tables. The immunization and error curing applied to these scenarios are not fast enough. There have been studies on the effect of isolating and curing network elements, however, the proposed strategies are limited to node removals.

This paper proposes a link isolation strategy based on the quarantining of susceptible clusters in the network. This strategy aims to maximize the epidemic control while minimizing the impact on the clusters performance. We empirically study the influence of clustering on robustness against epidemics in several real-world and artificial networks. Our results show an average curing rate improvement above 50% for the studied real-world networks under analysis.

I. INTRODUCTION

Epidemics on networks, from worm epidemics in computer networks to information spread in P2P and ad-hoc networks [13], [28] have recently attracted a lot of attention.

After the scanning worms, a new challenge for network security is posed by the strain of worms that use social networking Websites to spread. Web applications for exchange of information and data introduced new vectors of spread. Many social network worms use AJAX\(^1\) scripts like Samy [19], Yamanner [7] and Mikey [20]. Worm spreading usually involves user interaction in order to download worm payload on the local machine as Koobface [4], but recently Web clients are infected simply by visiting a Web page; no user interaction is necessary [19]. The infection risk increases since social networks are not restricted only to Facebook and Twitter, but are becoming embedded in other not strictly social websites like Digg and Youtube. Additionally, social networks have power law network structure which makes them prone to epidemic spreading [28], [3], [17] and [26].

The epidemic algorithms for information dissemination in unreliable distributed networks such as P2P and ad-hoc networks show similar epidemic dynamics on networks [13], [5]. Finally, the propagation of faults and failures can be modeled as an epidemic. Coffman et al. [18] models cascading BGP failures on a fully connected topology. We concentrate on protection against worms and error propagation in communication networks.

The protection of important networks in the above mentioned cases is in practice not fast enough, and the infection easily reaches all the segments of the network. This paper proposes and analyzes a fast method to stop or reduce epidemic spreading on networks. When an epidemic is detected, a network cut is performed by removing links leading to several disconnected clusters of nodes. This clustering allows limited intercommunity communication between nodes to continue, while possibly quarantining the rest of the network. Many real-world networks from on-line social networks to airline transport networks and Internet ASes network typically show a strong community structure [15], [21]. Depending on the speed of the epidemic reaction, it is possible to totally prevent any risk of infection for a number of disconnected clusters. Even with very delayed reaction, the amount of protection, that has to be applied in the network in order to stop the spreading, can be reduced. Thus, clustering can be used in addition to other protection methods.

The removal of links as protection against epidemics was proposed in mathematical epidemiology. The Equal Graph Partitioning (EGP) method uses immunization to remove specific nodes that cut the graph into clusters [6]. However, the immunization takes time, while individual nodes can stop communicating with other nodes immediately after receiving the news about the epidemic. Several authors have studied the reduction of disease spreading using air line restrictions. Goedecke et al. [16] and Epstein et al. [22] used the Susceptible Exposed Infected Recovered (SIER) model and dynamic time travel restrictions. Marcelino et al. [21] used the Susceptible Infected (SI) model together with edge betweenness and the Jaccard coefficient to increase the spreading time [21] by 81% by removing 25% of the links. Due to the multicommunity structure of the network with most connected nodes not being the most central, the optimal strategy for flight cancellation is not the removal of nodes (cities), but the removal of intercommunity flights, which introduced an increase in spreading time [21]. We are interested in specific link removal such that intra-community communication is preserved. We are not interested in optimizing of clustering algorithm, but instead in the general improvement of protection that is possible by using a well-

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\(^1\)Asynchronous JavaScript and XML
defined clustering algorithm.

Several algorithms have been proposed to find network communities. Modularity maximization is the most popular method. The modularity $Q$ is a quantitative criterion to evaluate how good a graph partition is [25]. It maximizes links within communities, while minimizing the links between them. Modularity maximization is an NP problem, given the exponential number of possible partitions. In this paper, we use a greedy heuristic proposed by Clauset et al. [8] to find an optimal modularity clustering.

In order to quantify the improvements of the network clustering in terms of epidemics, we use the epidemic threshold concept and the $N$-intertwined Susceptible Infected Susceptible ($SIS$) epidemic model [24] on a large set of networks. In a $SIS$ epidemic model, the epidemic can be stopped, provided the network protection functionalities against the virus perform faster than the reproduction of the virus. The epidemic thus exhibits threshold behavior.

In section II, we explain the protection algorithm and describe the networks that we examine. The epidemic theory used to estimate the protection is explained in section III. Results are presented in section IV, with comparison of random link removal and the modularity algorithm in section IV-C.

II. QUARANTINE MODEL AND NETWORKS

The protection method of dividing the network into clusters by removing links will be referred to as clustering or quarantining. The moment when a network is quarantined determines how many nodes are completely protected, since the virus is not able to infect nodes outside its cluster. In the first case, if we are able to quarantine a network into clusters faster than the virus is spreading, only a single cluster will contain infected nodes. On the other hand, if the virus infects all the clusters before a quarantine takes place there are still benefits, which are discussed in more details in section IV-B. Usually, the effective speed of clustering the network will be somewhere in between.

We discuss the two boundary cases separately. In the first case, we determine the size of the clusters, which provides an estimate of how many nodes will never get infected. The size of the clusters also affects the performance of the network. Larger clusters mean that a larger part of the network can continue exchanging information. Second, we show that the epidemic threshold that divides non-infected from infected networks favorably increases in networks that display clustering features.

If the infection is spreading very fast and all the clusters get infected, the number of infected nodes in the metastable state is reduced. We discuss the improvement with the respect to the number of removed links.

To illustrate the influence of clustering on epidemic spreading, we use several real-world networks. First, the Internet AS level topology obtained by Route View in 2006 and posted by the University of Oregon is used to demonstrate the effect of clustering on the virus spread in large infrastructural networks.

Further, we used an example of a social network between weblogs on US politics recorded in 2005 by Adamic and Glance [1]. The political blog network is shown in Fig. 1, with nodes belonging to different clusters colored in different colors. Finally, we examine an on-line social network of friends from www.digg.com, collected by the NAS at Delft University of Technology.

In disease modeling, transport networks are frequently used. To illustrate the influence of traveling patterns on virus spread, we investigate the direct airport-to-airport American traffic network maintained by the U.S. Bureau of Transportation Statistics and the European direct airport-to-airport traffic network obtained from the European commission for statistics Eurostat. The number of links and nodes of different networks are given in Table II.

In order to extend our understanding of the effects of clustering on the network robustness against virus spread, we include several artificial networks with $N = 1,000$ nodes.

We consider three Erdős-Rényi (ER) random graphs with a different number of links. Each node in the ER random graph is connected to every other node with probability $p$. The probability $p$ determines the number of links in the network [12]. We model power law networks using the Barabási-Albert model (BA) of preferential attachment for different number of links [2]. Finally, we use an artificial model of clustered networks [27]. The network is constructed in a similar manner as the ER random graph with two probabilities of link existence, one for inter-community connections and the other for intra-community connections. We have generated several different networks with $N = 1,000$, two clusters and different modularity. Further, we have considered networks with 4, 6, 8, 10 clusters. We choose to generate a greater number of networks with two clusters because most of the real-world networks consist of mainly two big clusters.

We additionally consider the square lattice, line, ring and tree topologies.

The networks are not weighted; however, the $N$-intertwined model is extendable to a heterogeneous setting [23].
III. N-INTERTWINED MODEL EPIDEMIC THRESHOLD

To model epidemic spread, we use the N-intertwined SIS model, which was introduced and discussed in [24]. A SIS model is one of the standard epidemic models: a node is susceptible to infection (S), then it becomes infected (I) and, after curing, it is susceptible to infection (S) again.

In order to quantify reduction of the number of infected nodes gained by clustering in the case of slow separation of the network, we use results of the N-intertwined model. A network is modeled as a connected, bidirectional graph $G(N, L)$.

By separately observing each node, the infection spread is modeled in a bidirectional network specified by a symmetric adjacency matrix $A$. A node $i$ at time $t$ can be in one of the two states: infected, with probability $v_i(t) = \Pr[X_i(t) = 1]$ or susceptible, with probability $1 - v_i(t)$. The sum of the probabilities of being infected and susceptible are equal to 1 because a node can only be in one of these two states. The state of a node $i$ is specified by a Bernoulli random variable $X_i \in \{0, 1\}$: $X_i = 0$ for a susceptible node and $X_i = 1$ for an infected node. We assume that the protection process per node $i$ is a Poisson process with rate $\delta$, and that the infection per link is a Poisson process with rate $\beta$ which is imminent for all nodes and thus constant in the network. For a node $i$, we can formulate the following differential equation

$$\frac{dv_i(t)}{dt} = \beta(1 - v_i(t)) \sum_{j=1}^{N} a_{ij} v_j(t) - \delta v_i(t)$$

where $a_{ij}$ is the element of the adjacency matrix $A$ and it is equal to 1 if the nodes $i$ and $j$ are connected, otherwise it is 0. A node is not considered connected to itself, i.e. $a_{ii} = 0$. The probability of a node being infected depends on the probability that it is not infected $(1 - v_i(t))$ multiplied with the probability that a neighbor $j$ is infected $a_{ij} v_j(t)$ and that it tries to infect the node $i$ with the rate $\beta$. Detailed derivations are given in [24] and [23].

In the steady-state, where it holds that $\frac{dv_i(t)}{dt} = 0$, and $\lim_{t \to \infty} v_i(t) = v_i(\infty)$ for each node $1 \leq i \leq N$, we have that

$$v_i(\infty) = \frac{\beta \sum_{j=1}^{N} a_{ij} v_j(\infty)}{\beta \sum_{j=1}^{N} a_{ij} v_j(\infty) + \delta} \quad (1)$$

This system of equations has $2N$ solutions with one positive solution and one solution equal to 0 [24]. The positive solution gives the probability that nodes are in the infected state during the steady-state of the model. The model gives a good approximation of the real epidemic process and the metastable state [24] for a wide range of effective spreading rates $\tau$. Thus, we will refer to the metastable state as a steady-state.

The fraction of infected nodes at any given time $t$ can be calculated as a sum of probabilities that the nodes are infected

$$y(t) = \frac{1}{N} \sum_{j=1}^{N} v_j(t)$$

and in the steady-state $y(\infty) = \frac{1}{N} \sum_{j=1}^{N} v_j(\infty)$.

For a fixed curing rate and spreading rate, the fraction of infected nodes as a function of the effective spreading rate $\tau$ is given in Fig. 2. The model as well as the real epidemic process have a threshold value at $\tau_c$. The threshold can be defined as follows: for effective spreading rates (rate of spread divided by rate of protection) below some critical value the virus in the network with $N$ nodes dies out before a large population is infected with a mean epidemic lifetime of the order of $O(\log N)$. For effective spreading rates above the critical value $\tau_c$, the epidemic persists and the number of infected nodes is large, with a mean epidemic lifetime [14] of the order of $O(e^{N/\tau})$ for a SIS model. The state above the epidemic threshold is referred to as the metastable state. In the metastable state, some constant mean portion of nodes is infected [24].

The epidemic threshold is equal to $\tau_c = \frac{1}{\lambda_{\text{max}}(A)}$, where $\lambda_{\text{max}}(A)$ is the largest eigenvalue of the matrix $A$ [14], [29] and similar results exist for Susceptible Infected Removed SIR model [11], [10]. We denote $\lambda_{\text{max}}(A)$ with $\lambda_{\text{max}}$. \label{eq:threshold}

If $\tau < \tau_c$, the infection will eventually be cured, and for $\tau > \tau_c$ the infection persists with the average number of infected nodes equal to $y(\infty)$.

For example, the largest eigenvalue of a line graph is $\lambda_{\text{max}} \simeq 2$, while that of a star topology is $\lambda_{\text{max}} = \sqrt{N-1}$. These two graphs are interesting examples, because both have the same number of links $L = N - 1$. Thus, the spreading in a star topology is significantly higher than in a line topology with the same number of nodes and links.

Figure 2 shows the threshold behavior for the steady-state of an infected network.

IV. RESULTS

In this section, we examine the case of instant clustering where a network is clustered faster than the worm is spreading, resulting in a single infected cluster. Further, we consider the case where all the clusters are infected before the quarantine

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{threshold.png}
\caption{Fraction of infected nodes as a function of the effective infection rate $\tau$. The epidemic threshold is denoted by $\tau_c$.}
\end{figure}
process clustered the network. Finally, we compare the quar-
tained networks with networks where the same number of
links has been randomly removed.

A. Early clustering

Defending the network and performing quarantines provides
important advantages. First of all, if a network is cut on time
and the infection is limited to one cluster, only a percentage
of nodes will eventually be exposed to infection. Second,
from the interlacing theorem of graph theory [9], the largest
eigenvalue of a subgraph is always smaller than that of the
graph. Thus, the thresholds \( \tau_c = 1/\lambda_{\text{max}} \) will always increase
for any subgraph, making the subgraphs more robust against
epidemic spreading. The case that all the clusters are initially
infected is discussed in section IV-B. Finally, the lifetime of
the metastable state depends on the number of nodes [14] as
\( \Omega(e^{N\alpha}) \), for \( \alpha > 0 \). The number of removed links using the
modularity algorithm ranges from 7\% to 58\% of the links. The
values for different networks are given in Table II.

One of the improvements introduced by clustering is a
reduction of the largest eigenvalue \( \lambda_{\text{max}} \) of the smaller clusters
with respect to the original graph. This increases the threshold
\( \tau_c \), the border between infected and non-infected networks.
The ratio between the largest eigenvalue of a cluster and the
largest eigenvalue of the whole network versus the modularity
\( Q \) for several networks is shown in Fig. 3 and 4.

The behavior of \( \lambda_{\text{max Cluster}} \) for the different network is diverse. For networks with high modularity, such as the
topologies, the improvement, a lowering of \( \lambda_{\text{max Cluster}} / \lambda_{\text{max G}} \) is not so significant. For the same type of networks,
e.g. BA or ER with different number of links, a reduced modu-
ality results in a reduced \( \lambda_{\text{max}} \), which is an improvement. For
either cases, the modularity is reduced by generating topologies
with a larger number of links (by respectively increasing the
parameter \( m \) in the BA model and the parameter \( p \) in the ER
model). In addition, the difference between the two largest
eigenvalues of different clusters is greater for BA than for ER.
The effect can be caused by the homogeneity of the degree
distribution of clusters in the ER case, while BA shows a
significantly heterogeneous cluster degree distribution.

The threshold \( \tau_{c, \text{cluster}} = \frac{1}{\lambda_{\text{max Cluster}}} \) increases as a function of the number of links removed between a cluster and the rest of the network, as shown in Fig. 5 and 6. In order to preserve as much network communication as possible upon link removal, a small number of links should be removed during the quarantine. On the other hand, \( \tau_c \) is inversely proportional to \( \lambda_{\text{max Cluster}} \). Hence the networks with best
performance show clusters with both low \( \lambda_{\text{max Cluster}} \) and low \( L_{\text{out}} \), close to the point \((0,0)\) in the figures. Real-world
networks such as the airline networks and AS network perform
well, while artificial networks perform much better the smaller
the number of clusters in the graph is.

For individual graphs, the dependency of threshold im-
provement versus the number of links removed is close to
linear, which is indicated by change in lower bound on largest
eigenvalue \( \lambda_{\text{max}} \geq \frac{2}{N} \). Sparse ER graphs are clustered easily,
with a small number of removed links, but show no significant
improvement in \( \tau_c \). The artificial, clustered graph with low
modularity shows the worst performance in the number of removed links, as in Figure 4.

The size of the clusters after cutting is an important variable
for the performance of the network. Large clusters will allow
for node communication after a quarantine. But on the other
hand smaller clusters will be more robust to virus spread. The
size of the clusters is decided by the modularity algorithm.

Another parameter to consider is the size of the largest
cluster after the quarantine. The distribution of the fraction
of cluster sizes \( \frac{N_c}{N} \) is shown in Fig. 7. In the case of early
clustering, the network is cut into clusters before the virus can
reach any other cluster except for the one it starts to spread
in. The worst case scenario is when the virus starts to spread
in the largest cluster. Most of the networks have one cluster
that contains half of the nodes. In the case of the European
air network, the three clusters pop up, thus leaving more than
two thirds of network protected. A BA graph has many small clusters of the size one fifth of network, which leaves four fifths of network protected, as shown in Fig.7.

The Digg network has one large cluster which covers half of the network and many significantly smaller ones. The USA air network has the largest cluster with the smallest number of deleted links, while the European air network has 3 clusters.

In Fig. 9 and 10, for the same network, larger clusters tend to have a larger $\lambda_{\text{max, Cluster}}$ than the smaller clusters. This is, however, not true for any graph: compare the path graph of any size with the complete graph of any smaller size.

B. Delayed clustering

We examine the number of infected nodes using the $N$-intertwined model. In order to clean the infected network, it is necessary to apply a protection/cleaning rate $\delta$ such that the effective spreading rate $\tau = \frac{1}{\tau_{\text{max}}}$ is below the threshold $\frac{1}{\lambda_{\text{max}}}$.

If the network is completely infected and then clustered, the amount of cleaning is reduced because $\lambda_{\text{max, Cluster}} < \lambda_{\text{max, G}}$, therefore $\tau_{\text{c}}(G) \leq \tau_{\text{c}}(\text{Cluster})$. Thus, if the network is clustered, it will be easier to clean the network from infection. Fig. 11 presents the percentage of infected nodes as a function of the effective spreading rate $\tau$ for different clusters in the artificial, cluster network with low modularity $Q$.

We calculate the fraction of infected nodes in the clustered network $y_{\text{Cluster}}$ for the effective spreading rate $\tau$ for which the number of infected nodes in the original network $y_{\text{tot, 50%}}$.
there are modularity of the graph. The improvement is different when of infected nodes exhibits the tendency to decrease with the number of removed links in the hole network, shown in Fig. 12. This is not surprising because the power of the largest eigenvalue of the cluster network models.

We calculate the fraction of infected nodes for several networks. Larger networks as the Internet AS and the Digg network are computationally more demanding and are left out of the analysis. In Fig. 12, the upper bound on the reduction of infected nodes reaches 50% and 80%. Then, we calculate the difference between the original value and the improved one:

$$y_{50\%} = y_{\text{tot},50\%} - y_{\text{clust}}, \ y_{80\%} = y_{\text{tot},80\%} - y_{\text{clust}}$$

We calculate the fraction of infected nodes for several networks. Larger networks as the Internet AS and the Digg network are computationally more demanding and are left out of the analysis. In Fig. 12, the upper bound on the reduction of infected nodes exhibits the tendency to decrease with the number of removed links in the hole network, shown in Fig. 13. This is not surprising because the power of spreading in a network decreases with links removal. Real-world networks do not show a significant reduction in number of infected nodes.

C. Random removal of nodes

In this section, we compare the threshold $\tau_c$ between quarantined networks with networks where the same number of links has been randomly removed. We give the largest eigenvalue of the original graph $\lambda_{\text{max},G}$, the size of the giant connected component $N_{\text{big,clust}}$, its largest eigenvalue $\lambda_{\text{max},\text{rand}}$, the size $N_{\text{big,clust}}/N_G$ and the largest eigenvalue $\lambda_{\text{max},\text{clust}}$ of the largest cluster in the clustered network in Table I. Links are removed at random and the average over many simulations of the largest eigenvalue of the largest connected component is calculated together with the variance of the largest eigenvalue. The results are presented in Table I. A large part of the network remains connected and can transmit infection, which is an expected result of random link removal. Between 80% and 90% of the network can be affected compared with at most 50% in case of clustering. Further, the largest eigenvalue of the
calculate the price of quarantine as the number of links that are the communication and reachability of nodes in the network. Shutting down links from the network reduces stopped and annihilated faster. However, protection comes removed from the graph as a result of a modularity clustering.

Random removal. have the same largest eigenvalue, which is the same as for the case of cluster 28s and 49s, two disconnected components. In networks where a small number of links is removed, no significant improvement in the largest eigenvalue is significant. In networks where a small number of links is removed, no significant improvement in the largest eigenvalue is significant. In networks where a small number of links is removed, no significant improvement in the largest eigenvalue is significant. In networks where a small number of links is removed, no significant improvement in the largest eigenvalue is significant. In networks where a small number of links is removed, no significant improvement in the largest eigenvalue is significant. In networks where a small number of links is removed, no significant improvement in the largest eigenvalue is significant. In networks where a small number of links is removed, no significant improvement in the largest eigenvalue is significant. In networks where a small number of links is removed, no significant improvement in the largest eigenvalue is significant.

The variance of the largest eigenvalue for different simula-
tion is also how good its resulting clustering is. We have not examined other algorithms that may perform differently, because we have concentrated on keeping the communities intact.

The largest eigenvalue improvement using the modularity algorithm is comparable with random links removal for several networks; however, in this case the worm can spread to 90% of the network.

V. CONCLUSION

This paper combines the diverse concepts of network clustering, graph spectra and epidemic spread in order to improve the protection against the spread of malware. We have found that real-world networks tend to show a better epidemic threshold \( \tau_c \) after clustering than artificially generated graphs.

For all the networks under study, the curing rate can improve between 29% and 83% for the largest connected component with respect to the original graph. This wide range of values demonstrates the effect of the network topology on the virus spread. Regarding the network clustering features, an easily clustered graph does not guarantee a slower epidemic threshold, but the way the links intertwine between inter- and intra-communities are key.

Overall, network protection against cascading failures can be improved for any kind of graph. However, the number of removed links is, in practice, unacceptably high. The advantages of early quarantine are shadowed by the fact that up to half of the links must be shut down for the quarantine to take effect.

The real-world networks have typically two or three big clusters and several smaller ones, while BA and ER graphs have several smaller ones comparable in size. BA and ER graphs are assumed to model the real-world complex networks. However, in respect of the size of the clusters, BA and ER fail to match real-world networks.

Additional to the epidemic spread analysis, this diversity in results appears valuable to create a general classification of types of networks. The degree distribution of the graph has been so far widely used for this purpose. For instance, a network classification could be generated by taking the largest eigenvalue of the adjacency matrix of clusters \( \lambda_{\text{max}} \text{Cluster} \) vs. links that are removed \( L_{\text{out}} \) as an input.

### Table II

<table>
<thead>
<tr>
<th>Network</th>
<th>( L_{\text{stat}} )</th>
<th>( L_{\text{removed}} % )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Euro</td>
<td>1,237, 14,952</td>
<td>47.27%</td>
</tr>
<tr>
<td>USA</td>
<td>2,179, 31,326</td>
<td>18.11%</td>
</tr>
<tr>
<td>BA 2m</td>
<td>1000, 1,971</td>
<td>42.36%</td>
</tr>
<tr>
<td>BA 3m</td>
<td>1000, 2,073</td>
<td>58.88%</td>
</tr>
<tr>
<td>ER 0.002</td>
<td>868, 982</td>
<td>17.34%</td>
</tr>
<tr>
<td>ER 0.006</td>
<td>1000, 3,054</td>
<td>51.27%</td>
</tr>
<tr>
<td>ER 0.02</td>
<td>1000, 9,938</td>
<td>55.02%</td>
</tr>
<tr>
<td>AS ’06</td>
<td>22,963, 48,436</td>
<td>20.62%</td>
</tr>
<tr>
<td>Pol. Blog</td>
<td>1,222, 19,021</td>
<td>7.16%</td>
</tr>
<tr>
<td>Digg</td>
<td>254, 371, 4,354, 174</td>
<td>25.02%</td>
</tr>
</tbody>
</table>

Fig. 13. The difference between the number of infected nodes in the original network and the clustered network as a function of the relative number of removed links in the case when 50% and 80% of nodes are infected in the original network.
The clustering with random removal of links has led us to conclude that the largest eigenvalue of the largest cluster can be less or comparable to the largest eigenvalue of the biggest component generated by random links removal. However, other clusters have a significantly smaller largest eigenvalue, which leads to a smaller amount of cleaning necessary to completely remove the worm from the network. Furthermore, if only the largest cluster is infected, only up to 50% of the network will need cleaning.

This paper considers modularity to be the partitioning algorithm, but there exists a large number of partitioning algorithms that try to optimize different variables. The investigation of how different clustering algorithms affect the epidemic dynamics stands on the agenda for future work.

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