Contact tracing and epidemics control in social networks

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A generalization of the standard susceptible-infectious-removed (SIR) stochastic model for epidemics in sparse random networks is introduced which incorporates contact tracing in addition to random screening. We propose a deterministic mean-field description which yields quantitative agreement with stochastic simulations on random graphs. We also analyze the role of contact tracing in epidemics control in small-world networks and show that its effectiveness grows as the rewiring probability is reduced.

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Properties of complex networks recently attracted much attention in physical community\textsuperscript{[1]}. Although perhaps it was prompted by the advent of the Internet and World-Wide Web, the importance of this subject goes far beyond computer networks. Indeed, daily commute, power and goods traffic, wired and wireless communication, disease spreading occur within certain physical or social networks. The theory of disease spreading, which is known as mathematical epidemiology, has a long and rich history (see, e.g.,\textsuperscript{[2]}). However, until recently the epidemiological studies have been mostly concerned with so-called mean-field description of epidemics, in which it is assumed that at any time the probability to get infected is the same for all individuals. In some other works, spreading of a disease on relatively simple lattices of individuals has been studied within so called “forest-fire” models\textsuperscript{[3]}. Only recently, the studies which elucidate the role of underlying network structure in the disease spreading began to appear in the literature\textsuperscript{[4,5]}.

Most of the epidemiological models are based on several simple assumptions regarding disease contracting and cure. In particular, the most common mechanism of infection is through a contact with another infectious individual, and the mechanism of recovery is either deterministic or purely stochastic with a certain typical time of recovery. In the simplest Susceptible-Infectious-Susceptible (SIS) model, a recovered individual immediately becomes susceptible again, while in a more complicated Susceptible-Infectious-Removed (SIR) model, cured individuals become immune and effectively excluded from further dynamics.

While these models give a good description of evolution of many common infectious diseases, they usually neglect the role of intelligent strategies to stop nascent epidemics. Few epidemiological models take into account prevention strategies such as, for example, mass and ring vaccination\textsuperscript{[6]}. In practice, one of the main counter-epidemics measures is the contact tracing, when individuals which have been in contact with an infected (and identified) individual, are found and thoroughly checked. It applies, among others, to the treatment of sexually transmitted diseases, tactics of law-enforcement organizations trying to uncover criminal or terrorists networks, cleaning of computer virus infection, etc. We only are aware of one theoretical paper\textsuperscript{[7]} where a model of this kind has been studied. The model\textsuperscript{[8]} is based on the assumption that infection is a slow branching process, while the contact tracing occurs at a much shorter time scale. This leads to a familiar SIR-type model with rescaled parameters and similar dynamics. In this Letter we consider a more realistic model in which infection and contact tracing occur concurrently, and their interplay determines the dynamics of the system.

\textit{Stochastic model.} We assume that the population consists of $N$ hosts whose connections to each other form a fixed graph. The hosts are enumerated with index $n = 1, ..., N$. A node $n$ is said to have a degree $k(n)$ if it is connected to $k$ other hosts. In case of random graphs the degree distribution is Poissonian with a certain mean degree $K = \langle k(n) \rangle$.

For simplicity we assume that there in no spontaneous recovery, an infectious individual can only be disinfected externally through screening. Immediately upon disinfecting, the individual becomes traced ($T$) for a certain period of time during which its neighbors are checked for possible infection. After that time, the individual spontaneously becomes removed, and its neighbors are no longer traced.

\textbf{Infection} $S \to I$. Initially, the whole population except for one host is assumed to be susceptible to infection. The probability of host infection depends on the state of its nearest neighbors. The infection dynamics is modeled as a simple contact process: if a susceptible node $n$ has $k_i(n)$ infectious neighbors, the probability that it becomes infectious during a small $\Delta t$ time interval is $\alpha k_i(n) \Delta t$.

\textbf{Tracing} $I \to T$. The process of infection elimination consists in finding infectious hosts and then curing them. Hosts are being checked with certain probability $\beta$ which depends on the state of its neighbors. We postu-
late that if an infectious host is checked, it is immediately cured, eliminated or at least isolated so it cannot infect other hosts. We introduce two non-exclusive strategies of checking for infectious hosts: random checking and contact tracing. Random checking means choosing an arbitrary host with probability \( \beta_t \Delta t \), while contact tracing of host \( n \) is done with probability \( \beta_t k_t(n) \Delta t \) where \( k_t(n) \) is the number of neighbors of \( n \) which are in the traced state \( T \). The random checking process is equivalent to the removal process of general epidemics [3].

**Removal** \( T \to R \). With certain probability \( \gamma \Delta t \), traced hosts are transformed into the removed state, in which they also cannot be infected, but they are no longer under observation, so they do not initiate contact tracing.

**Stochastic simulations** of the described process were performed using an event-driven scheme [10]. It is significantly superior over synchronous and asynchronous update schemes both in terms of accuracy and computational speed. In the event-driven scheme we select the time lapsed between two consecutive events from a Poisson distribution with a combined probability of all events (infection, tracing, recovery, etc.), then choose a node (each node has its own probability to be chosen depending on its state and the states of its neighbors), and apply the transition from one state to another according to the ratio of individual transition probabilities.

In our simulations with random graph based networks, we typically built networks with average degree \( K = 10 \) and 1000 nodes. For every random graph we ran 100 simulations starting every time from a single (but different in each run) infected host. Then we averaged the results for 50 random graphs. The time bin was \( \Delta t = 10^{-6} \) for most simulations. In all simulations we varied the tracing parameters \( \beta_t \) and \( \beta_r \), while the infection constant was set at \( \alpha = 0.1 \), and the transition rate from \( T \) to \( R \), \( \gamma = 0.5 \). The latter parameter is important for the effectiveness of the targeting elimination, because the longer a node remains in the traced state, the more probable it is to trace its neighboring infectious nodes. However, since tracing presumably bears a significant cost, an optimal choice of the tracing parameters (\( \beta_r, \beta_t \), and \( \gamma \)) for a given epidemics is an important issue.

In Fig. 1 we present the "prevalence" of epidemics (the fraction of infectious nodes in the whole population \( i = I/N \)) as a function of time for several values of \( \beta_r \) and \( \beta_t \). When \( \beta_r = \beta_t = 0.0 \) we have a simple SI process, and all the nodes eventually get infected (thick solid line in Fig. 1). Other curves show the fraction of infectious nodes as a function of time for \( \beta_r = 0.02 \) and different values of \( \beta_t \). The ratio \( \alpha/\beta_t \) is chosen to be above the epidemics threshold [2]. For \( \beta_t = 0 \) we obtain the classical SIR process with randomly removed infectious (dashed line in Fig. 1). The epidemic eventually saturates, and the fraction of infectious nodes decays exponentially. The lower lines display the evolution of the infection fraction for values of \( \beta_t \) ranging from 0 to 2.5 with a step value of 0.1. The initial (exponential) phase of the epidemics growth is nearly independent of \( \beta_t \), because the contact tracing process is intrinsically nonlinear (it requires the presence of I-T connected pairs and therefore only begins after the first infected node is randomly screened).

As expected, the tracing process significantly reduces the magnitude of the epidemics (maximal value of \( i \)), but at large times the infection decays with the same exponential rate as for \( \beta_t = 0.0 \) (again, because we return to the linear regime at small \( i \)). The most interesting feature of the process at large \( \beta_t > 0.35 \) is the presence of a second maximum of \( i \) which indicates a second epidemic. Due to this second epidemic, the percentage of the infectious population at large times \( t > 40 \) may actually increase with increase of \( \beta_t \). It means that the range of \( \beta_t \) values from 0.4 to 0.9 are not better to control the epidemics than values smaller than 0.4.

**Mean-field equations.** At the first sight, it seems that the mean-field approach cannot be applied to the contact tracing, since it does not take into account the non-uniform distribution of infection in the population. Nevertheless, a more sophisticated mean-field approach which operates not only with the mean densities of states, but also with the densities of links connecting nodes with different states, can be devised. In the derivation we follow the method of Rand [3] (see also [4]). Let us introduce the number of nodes \( A \), the number of connected pairs \( AB \) and triples \( ABC \) of nodes, where \( A, B \) and \( C \) stand for any of the types \( S, I, T, R \). For example, the number of connected pairs of infectious and traced nodes is denoted \( [IT] \). Note that \( [AB] = [BA] \) and each pair in \( [AA] \) is counted twice. For large \( N \), the ratios \( A/N, [AB]/N, [ABC]/N \) approach deterministic limits which we label \( a, [ab], [abc] \), respectively.

The dynamics of the model is described by the following set of rate equations.
\[ \dot{s} = -\alpha s i, \quad \dot{i} = \alpha s i - \beta s i - \beta i \dot{\tau}, \]
\[ [ss] = -2\alpha [ssi], \]
\[ [si] = \alpha ([ssi] - [s]i - [si]) - \beta s [si] - \beta i \dot{\tau}, \]
\[ [ii] = 2\alpha ([isi] + [si]) - 2\beta s [ii] - 2\beta i [i\tau], \]
\[ [i\tau] = \beta i [ii] + \beta s [i\tau] - [\tau \tau] - [i\tau] - \gamma [i\tau] - \beta r [i\tau]. \]  

Here we used the notation \( \tau = [T]/N \) to avoid confusion between the density of traced nodes and time \( t \). Note that we omit here the equations for \( \tau, [\tau \tau], [s\tau] \), as well as any combinations involving the removed state, since they do not affect the dynamics of the infectious population.

The meaning of these equations is rather straightforward. For example, the terms in the r.h.s. of the last equation can be explained as follows. A \((p, q)\) pair becomes \([i\tau]\) through random screening of the infectious node \( q \) in an \([ii]\) pair, or through contact tracing of node \( q \) from node \( r \) in a \((p, q, r)\) triple in the state \([i\tau]\). On the other hand, we can lose an \([i\tau]\) pair by contact tracing of the \( p \) node in a \([\tau \tau]\) triple \((r, p, q)\), by direct tracing \( p \) by \( q \), by removing of \( q \), or by random screening of \( p \). Other equations can be obtained from similar arguments.

This set of equations is not closed, as the equations for the pair densities contain triple densities. We need to introduce a closure rule. Similarly to Refs. [1][2], we can use the approximation \([abc] = [ab][bc]/b\), which follows from the condition that the influence of a node on the state of its second neighbor in a triple is negligible [1].

Using this closure rule, we arrive at the following set of equations
\[ \dot{s} = -\alpha s i, \]
\[ \dot{i} = \alpha s i - \beta s i - \beta i \dot{\tau}, \]
\[ \dot{\tau} = (\beta p - \beta \dot{\tau} - \gamma - \alpha \hat{\tau}) \hat{\tau} + \beta \dot{p}, \]
\[ \dot{\hat{p}} = \alpha s (2i + 2 - \hat{p}) - (\beta \dot{r} + \beta i \dot{\tau}) \hat{p}. \]  

where \( \dot{i} = [is]/s \) is the mean number of infectious neighbors per susceptible node, \( \dot{\tau} = [i\tau]/i \) is the mean number of traced neighbors per infectious node, and \( \hat{p} = [ii]/i \) is the mean number of infectious neighbors of an infectious node. Notice that the equation for \([ss]\) dropped out as \([ss] = K s^2 \) at all times. We used the initial conditions \( s(0) = 1 - i_0, i(0) = i_0, \dot{i}(0) = (K - 1)i_0, \dot{\tau}(0) = 0, \dot{\hat{p}}(0) = 0 \) which correspond to a small set of disconnected infectious nodes.

During the early stage of an epidemic the contact tracing can be neglected (\( \tau = 0 \)), and Eqs. [1]-[1] are reduced to a set of three equations for \( s, i, \dot{i} \) which coincide with the model has been studied in [3][4]. Independently of \( \beta_i \), the initial epidemics growth is characterized by the basic reproduction number \( K \alpha/(\alpha + \beta_i) \). However, as the number of traced individuals grows, the growth rate is reduced and the epidemics is saturated. In Fig.2 the dynamics of the epidemics calculated from Eqs.[1]-[1] are shown for different values of \( \beta_i \). As seen from the figure, the maximum number of infectious nodes is drastically reduced with increase of \( \beta_i \). In the same figure we show the results of direct stochastic simulations for \( \beta_i = 0, 0.5 \) [2]. The most important question is whether the contact tracing is capable of arresting the exponential growth of the epidemics before it engulfs a finite portion of the total population.

![FIG. 2. Evolution of the infection prevalence for different \( \beta_i \). Mean-field model (lines) and stochastic simulations (symbols), \( \alpha = 0.1, K = 10, \beta_s = 0.02, \gamma = 0.5 \).](image)
node in the graph can be exposed to infection in a short time after an epidemics begins. If the clustering coefficient is large, the infection propagates faster within a certain community but then it make it easier to trace the epidemics. Sparse random networks studied above represent a particular class of networks with short average minimal path and small clustering coefficient. Many social networks are characterized by a relatively large clustering coefficient while keeping the average minimal path low. We studied numerically the effect of the network structure on the contact tracing of epidemics within the small-world model [13]. Changing the re-wiring probability \( p \) allows us to scan the range of networks from regular \((p = 0)\) to random \((p \rightarrow 1)\) through the small-world range \((0.001 < p < 0.1)\) which exhibits a short average minimal path and a large clustering coefficient typical for many social networks. We used the same number of nodes and edges as for the random graph simulations, and fixed the parameter values at \( \alpha = 0.1, \beta_r = 0.02, \) and \( \gamma = 0.5 \). Fig. 3 shows the dependence of the maximum prevalence \( i_{\text{max}} \) on \( p \) for several different \( \beta_r \). As we can see, \( i_{\text{max}} \) changes mostly within the SW range \((0.001 < p < 0.1)\) where the clustering coefficient and the average path undergo large variations.

![FIG. 3. Maximum number of infectious nodes vs. \( \beta_r \) for different \( i_0 \). Parameters are the same as in Fig. 4](image)

In this Letter we studied the role of contact tracing as a part of the epidemics control strategy in complex networks. We demonstrated that by applying this strategy, a major outbreak can be significantly reduced or even eliminated at a small additional cost. Based on the pair correlation approach due to Rand [6], we developed the mean-field model of contact tracing for the case of random graphs. We also studied the influence of network topology on the contact tracing using the small-world model with variable re-wiring probability \( p \), and found that its effectiveness grows as the rewiring probability is reduced. The main change occurs within the small-world regime at \( p \sim 10^{-2} \).

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![FIG. 4. The maximum prevalence as a function of the rewiring probability \( p \). Effectiveness of the contact tracing becomes very significant in the small-world regime.](image)

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