Worst-Case Scenarios and Epidemics

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Chapter 2

Worst-Case Scenarios and Epidemics

Gerardo Chowell* and Carlos Castillo-Chavez†

2.1 Introduction

The potential deliberate release of biological agents such as smallpox, influenza, and foot and mouth disease (FMD) is a source of continuous concern. It is believed that “worst” epidemics (single outbreaks) are most likely to occur in populations where individuals mix randomly (proportionate mixing). Therefore, the definition and identification of landscapes or topologies that support worst-case scenarios is critical. Here, SIR (susceptible-infected-recovered) epidemics that result from the introduction of single or multiple infectious sources are studied on various topologies including small-world and scale-free networks.

Most of the models studied in classical mathematical epidemiology fall in the class of compartmental models because the population under consideration is divided into classes or compartments defined by epidemiological status [1]. The simplest version assumes that individuals mix uniformly (homogeneous mixing) within each compartment. The study of the transmission dynamics of communicable diseases in human populations via mathematical (compartmental) models can be traced back to the work of Kermack and McKendrick (1927). Their simple SIR (susceptible-infected-removed) epidemic model (Figure 2.1) not only was capable of generating realistic single-epidemic outbreaks but also provided important theoretical epidemiological insights.

Figure 2.1. Simple SIR epidemic model.

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The Kermack and McKendrick (K-M) model in essence captures the theoretical underpinnings associated with the framework that it is currently used to define (and model) worst-case epidemic outbreaks. The K-M model is given by the following system of non-linear differential equations:

\[
\begin{align*}
\dot{S} &= -\lambda(N)SI, \\
\dot{I} &= \lambda(N)SI - \gamma I, \\
\dot{R} &= \gamma I,
\end{align*}
\]

where \( S(t) \) denotes susceptible individuals at time \( t \); \( I(t) \) infective (assumed infectious) individuals at time \( t \); \( R(t) \) recovered (assumed permanently immune) individuals at time \( t \); \( \lambda(N) \) the transmission rate when the total population is \( N(N = S + I + R) \); and \( \gamma \) the recovery rate. In the case of a fatal disease, \( N = S + I \) as \( R(t) \) would denote those removed by death and \( \gamma \) the per-capita death rate.

The K-M threshold theorem establishes (quantitatively) the conditions required for successful disease invasion. This threshold theorem says that a disease will invade whenever its basic reproductive number

\[
R_o = \frac{\lambda(N(0))N(0)}{\gamma} > 1.
\]

\( R_o \) is interpreted as the number of secondary infectious individuals generated by a “typical” infectious individual when introduced into a fully susceptible population [2, 3]. An alternative interpretation is that, in a randomly mixing population, a disease will invade if there are enough susceptibles, that is, if

\[
N(0) > \frac{\gamma}{\lambda(N(0))}.
\]

Typically, it has been assumed that either \( \lambda(N) = \frac{\beta_0}{N} \) or \( \lambda(N) = \beta_0 \), and consequently, the interpretation of \( R_o \) depends on the definition of \( \lambda(N) \) [5, 1, 2].

Nold [6] introduced the concept of proportionate mixing as a way of modeling a simple form of heterogeneous mixing. She divided the population into \( K \) groups, each with population size \( N_i(t) = S_i(t) + I_i(t) + R_i(t) \) \((i = 1, \ldots, K)\) and modeled an SIR epidemic that considered interactions of various intensities among individuals. In order to describe her framework we introduce the mixing matrix \( P(t) = (P_{ij}(t)) \), where \( P_{ij}(t) \) denotes the proportion of contacts of individuals in group \( i \) with individuals in group \( j \) given that \( i \)-individual had a contact with a member of the population at time \( t \). Nold’s proportionate mixing corresponds to the case where \( P_{ij} \) is independent of \( i \), that is,

\[
P_{ij} = \tilde{P}_j = \frac{C_jN_j}{\sum_{l=1}^{k} C_lN_l}, \quad (\Pi)
\]

where \( C_l \) denotes the average activity level (contact rate) of individuals in group \( l = 1, \ldots, k \).

Other forms of mixing can be found in [2, 7, 8, 9] and references therein.

A slightly modified version of Nold’s generalization of the K-M model (assumes that
\( \lambda_j = \frac{\beta_j}{N_j}, j = 1, \ldots, K \) is given by

\[
\begin{align*}
\dot{S}_t &= -S_t(t) \sum_{j=1}^{K} \beta_j \tilde{P}_j \frac{l_j}{N_j}, \\
\dot{I}_t &= S_t(t) \sum_{j=1}^{K} \beta_j \tilde{P}_j \frac{l_j}{N_j} - \gamma_t I_t, \\
\dot{R}_t &= \gamma_t I_t, \quad i = 1, \ldots, K.
\end{align*}
\] (2.3)

Examples of mathematical studies for compartmental models of this type can be found in [5, 1, 2, 6]. Extensions of these models to (local) populations interconnected via migrating individuals (metapopulation models) have been carried out in some situations (see [9] and references therein). In the K-M model and Nold’s models, individuals mix at random (see Figure 2.2), an uncommon and extreme situation. Recent analyses of worst-case scenarios for the deliberate release of biological agents (smallpox in particular) have been carried out under the assumption that random mixing supports the worst epidemic outbreaks [10]. The focus of this paper (as suggested to us by Ed Kaplan) consists of the preliminary examination of the validity of this assumption.

![Figure 2.2. Fully mixed transmission network.](image)

The identification of worst-case scenarios would require an approach that considers “all” measures of mixing [6]. The study of the meaning of worst-case scenarios using “mean” field models like the K-M or Nold’s model is problematic (mixing assumes a predetermined number of groups or types). Hence, we look at this question in the context of simple individual-based models where mixing is embedded in a preselected (fixed) topology. This approach also has its limitations as the nodes (individuals) of the network have no dynamics. However, we feel that this approach is useful if the goal is to study the “strength” of a single epidemic outbreak, that is, in the study of situations where transient disease dynamics are critical.

### 2.2 Individual-Based Models

Population structures are often represented by networks (graphs) composed of nodes (individuals) and edges (representing predefined relationships between nodes). Examples include family trees, traffic networks that describe street intersections by nodes and traffic direction...
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(a)

(b)

Figure 2.3. The Erdős and Rényi random graph with $N = 16$, (a) $p_{ER} = 0.25$, (b) $p_{ER} = 0.5$.

by arrows (edges), and airline traffic networks. Graphs (networks) can be represented by the adjacency matrix $T$, where $T(i, j) = 1$ implies that vertex $i$ is connected to $j$. If the network is undirected (edges have no direction), then $T$ is symmetric; that is, $T = T^t$ (transpose of $T$). An adjacency list, a compilation of the vertices of a graph and the vertices adjacent to such a vertex, is also used to represent graphs. The analysis of network models can be traced back to the work of Erdős and Rényi in the 1960s. These researchers introduced the following simple algorithm for the construction of random networks [11]: start with a fixed number of disconnected nodes $N$ and connect each pair of nodes independently by an edge with probability $p_{ER}$. Hence, $p_{ER} = 0$ corresponds to the case where no node is connected to any of the other $N - 1$ nodes, while $p_{ER} = 1$ corresponds to the case where every node is connected to all other nodes in the network (complete graph). Examples are depicted in Figure 2.3. The total number of edges when $p_{ER} = 1$ is $\binom{N}{2}$; the average number of edges is $\frac{N(N-1)p_{ER}}{2}$; and the average degree of a node (number of edges incident from a node) is $z = (N - 1)p_{ER} \approx Np_{ER}$ (for large $N$).

Erdős and Rényi [11] showed that for large systems (large $N$) the probability that a node has $k$ edges follows the Poisson distribution $P(k) = \frac{\exp(-z)c^k}{k!}$ ($k = 0, 1, \ldots, N$). Furthermore, they showed that there is a critical value of $z$ ($z_c$) such that whenever $z > z_c$, a connected component (a subset of vertices in the graph each of which is reachable from the others by some path through the network) forms. Such a component is often referred to as a spanning cluster [14]. The Erdős and Rényi random graph provides a null-model for the “comparative” study of the disease transmission on other network topologies. The case $p_{ER} = 1$ (totally connected network) is naturally believed to be the generator of the landscape most conducive to disease spread and, in some sense, “corresponds” to Nold’s version of the K-M model. Hence, here we address, via simulations, whether or not populations modeled by graphs comparable to those of the Erdős and Rényi model support the worst possible epidemics. It is not clear, however, whether or not diseases spread at a faster rate in highly clustered networks, that is, in networks where there is a higher probability that neighbors of a particular node are also neighbors of each other. The importance of this possibility
Figure 2.4. Small-world networks with $N = 16$, $K = 1$, (a) $p = 0.1$, (b) $p = 0.3$.

comes in part from the fact that Watts and Strogatz [15] showed that networks with high degree of aggregations (clustering), a characteristic absent in random networks (see Erdős and Rényi [11]), are not uncommon. In order to explain our simulations of epidemics in various network setups, we need to describe their construction.

Watts and Strogatz's [15] algorithm for the construction of networks is as follows: a one-dimensional ring lattice of $N$ nodes connected to its $2K$ nearest neighbors ($K$ is known as the coordination number) and periodic boundary conditions are preselected. The algorithm goes through each of the edges in turn and, independently with probability $p_{WS}$, "rewires" it to a randomly selected node. That is, the Watts–Strogatz algorithm shifts one end of the edge to a new randomly chosen node from the whole lattice (except that no two nodes can have more than one edge running between them, and no node can be connected by an edge to itself (see Figure 2.4). Watts and Strogatz [15] classified networks by their level of randomness, as measured by their own disorder parameter $p_{WS}$ (from "regular" $p_{WS} = 0$ to completely random, $p_{WS} = 1$). Whenever each node in a network is just connected to its nearest two neighbors one on its right and on the other on its left, the network is regular [15]. A completely random network has $p_{WS} = 1$; that is, all nodes are randomly connected to each other. Watts and Strogatz showed that the introduction of a small number of random connections ($p_{WS} \approx 0.01$) significantly reduces the average distance between any two nodes (characteristic path length), a property that facilitates disease spread. In fact, Watts and Strogatz showed that such average distance grows like $O(\log(N))$ and not as $O(N)$. Networks constructed via the Watts–Strogatz algorithm also support high levels of clustering. The small-world effect (short average distance between nodes and high levels of clustering) has been detected in networks that include a network of actors in Hollywood, the power generator network in the western US, and the neural network of Caenorhabditis elegans [15]. This “small-world effect” had already been documented by the psychologist Stanley Milgram, using data from the letter-passing experiments that he conducted in the 1960s [16]. Newman and Watts [17] studied a slight variation of the Watts–Strogatz model. They added shortcut edges with probability $\phi$ per edge in the underlying ring lattice instead of "rewiring" the existing edges. The Newman and Watts model turned out to be easier to
analyze since the network cannot become disconnected after rewiring. Figure 2.5 shows a small-world network with \( N = 16, K = 1, \) and \( \phi = 0.1. \) The degree or connectivity distribution \( P(k) \) (\( k \) is the degree or number of connections per node) of the small-world network model depends on the disorder parameter \( p_{\text{WS}}. \) That is, \( p_{\text{WS}} = 0 \) implies that the connectivity distribution is given by the delta function \( \delta(k - 2K), \) where \( K \) is the coordination number in the network. As \( p_{\text{WS}} \) approaches 1, the connectivity distribution converges to that
Figure 2.7. The power-law distributions observed in (a), (b) scientific collaboration networks (The Los Alamos e-Print Archive) [12]; (c) the World Wide Web (nd.edu domain) [13]; and (d) the location-based network of the city of Portland [25].

obtained from the Erdős and Rényi model. Figure 2.6 shows the degree distribution for a small-world model as a function of the disorder parameter $p_{ws}$, when $N = 10^4$ and $K = 3$. “Unfortunately,” small-world networks do not exhaust all possibilities. The bell-shaped node degree distributions observed in the Erdős–Rényi, Watts–Strogatz, and Newman–Watts models are in contrast with the power-law degree distributions observed in a number of biological [18], social [19, 20, 21, 22, 23, 24, 25], and technological [19, 20, 26, 27] networks (see Figure 2.7). (Power-law degree distributions—also known as Pareto distributions—are given by the parametric family

$$P(k) = Ck^{-\gamma},$$

where $\gamma$ is typically between 2 and 3 (infinite variance) and $C$ is a normalization constant (makes $P(k)$ a probability density function). Networks that fit power-law degree distributions (free of characteristic scale) well have a small number of highly connected nodes; that is, most nodes have a small number of connections. Barabási and Albert [19] dubbed this type of structures scale-free networks.

The number of sexual partners in the 1996 Swedish survey of sexual behavior [24] fits a power-law distribution and the number of sexual partners of Cornell University undergraduates from the 1990 Cornell Undergraduate Social and Sexual Patterns (CUSSP) Survey [28] can also be fitted well by such distribution (see Figure 2.8). These observations, for example, support the view that sex-education campaigns must target individuals with the highest number of partners (core group) [29]. The location-based network of the city of Portland, Oregon also exhibits a scale-free structure (Chowell et al. [25]). Here, nodes represent locations while directed connections between locations represent the average movement
of individuals in the city. The scale-free (Figure 2.7(d)) topology implies the existence of a high number of locations with a low number of connections (i.e., households) and a small number of highly connected hubs (i.e., schools, hospitals, etc.). Barabási and Albert [19, 20] introduced a simple theoretical model that generates networks with a power-law degree distribution (see Figure 2.9). The Barabási–Albert (BA) algorithm starts with a small number of nodes \( m_0 \), and at each time step, a new node connects (with \( m \) links), with higher probability, to nodes that have already accumulated a higher number of connections. The resulting network has a power-law exponent of 3 and a mean connectivity of \( 2m \). Thus, the BA model captures features that seem characteristic of real-world networks, namely, growth and preferential attachment. Figure 2.10 shows a network generated using the BA model.

Several modifications of the BA model have been studied, including edge rewiring [30], edge removal [31], growth constraints [32, 33], and edge competition [34]. Klemm and Eguíluz [35] developed an alternative algorithm for the generation of scale-free networks. These researchers incorporated memory as part of a node's ability to acquire additional links. The Klemm–Eguíluz model produces scale-free networks with high clustering coefficients, a property not generated by the BA model.

The capacity of networks to maintain essential properties when nodes are removed is a measure of their robustness. In scale-free networks, most of the nodes have low degree; hence their removal "typically" does not impact the connectivity of the remaining vertices. However, the removal of nodes with the highest degree (pressure points) of connectivity can have dramatic consequences (see Figure 2.11). This effect was first demonstrated independently and numerically by Albert [36] and Broder et al. [37], using subsets of data of the World Wide Web. The practical relevance of network robustness was highlighted by the February 2000 service denial of highly connected web servers, including Yahoo, CNN, Amazon, eBay, Excite, and E*TRADE, following a network attack.

**Figure 2.8.** Number of sexual partners of Cornell University undergraduates from the 1990 Cornell Undergraduate Social and Sexual Patterns (CUSSP) Survey [28]. The power-law exponent for females is 2.86, for males is 2.90, and for the overall distribution is 2.78.
Figure 2.9. Connectivity distribution of the BA model decays as a power law with $\gamma = 3$. Here, $N = 10000$, $m_0 = 3$, and $m = 2$.

Figure 2.10. Network of size $N = 23$ generated using the BA model described in the text with $m_0 = 2$ and $m = 1$.

Important work on the use of worst-case scenarios in the development of response policy has been carried out by Kaplan [38, 39, 10] in the context of HIV and smallpox. However, the nature of his approach does not allow for the incorporation of population structures such as those identified in [19, 20, 21, 22, 23, 25]. The importance and frequency of these networks is the main motivation behind our efforts to look at the rate of growth of epidemic outbreaks on these graphs. The focus of this paper (instigated by Ed Kaplan) is driven by these questions:
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- How is the initial rate of epidemic growth affected by a population’s structure?
- What are the roles of “social” topologies and the number of initial infectious sources on the rate of growth of an epidemic?

The following sections represent an initial attempt to address these questions in the context of small-world [40, 15] and scale-free networks [19]. The organization of the rest of this paper is as follows: Section 2.3 corroborates Kaplan’s view of mixing in worst-case epidemics when the transmission topology is given by small-world networks; section 2.4 focuses on the study of epidemics in scale-free networks where a natural “node” hierarchy often emerges. (This structure, in some sense, “equivalent” to the concept of core group developed by Hethcote and Yorke [29], seems to provide a fluid landscape for disease spread); section 2.5 collects our conclusions, caveats, and views on implications of these results in the study of the impact of deliberate releases of biological agents.

2.3 Epidemics on Small-World Networks

Simple epidemic models such as the susceptible-infected-recovered (SIR) model have been studied on small-world networks. Moore and Newman [41] studied SIR epidemics on small world networks via site and bond percolation. In site percolation, nodes (sites) are occupied (by spins) or not and any two spins occupying nearest neighbor sites are connected by an open bond. In bond percolation, the relevant entities are bonds or edges. Bonds are sequentially visited and set open with probability $p$ or closed with probability $1 - p$ (independently). The percolation threshold is the smallest probability $p$ at which an infinite cluster of sites emerges when sites or bonds (depending on the type of percolation) are occupied with that probability.

Figure 2.11. The size of the largest component in a scale-free networks as a function of the number of nodes removed randomly (dashed line) or in decreasing order of degree; that is, hubs are removed first (solid line). The network was constructed using the BA model with $N = 10000$, $m_o = 3$, and $m = 2$. 

![Figure 2.11](image-url)
SIR epidemic processes are built on the assumption that nodes are occupied by individuals who can be infected by neighbors connected by edges or bonds (Grassberger [42]). When the total probability of transmission from one individual to another is greater than this threshold, the disease explodes; that is, a “giant” component (whose size is the size of the epidemic) appears [14, 43]. In an epidemic that starts with a single infectious source and spreads as a bond percolation process, the subset of nodes (individuals) that can be reached from the initial infective individuals by traversing only open bonds is the size of the epidemic outbreak. Newman, Jensen, and Ziff [44] studied SIR epidemics on a two-dimensional small-world network via bond percolation. His investigation was motivated by the study of disease transmission in plants coming from two sources: from nearest neighbor (plants) and long-distance contacts (vectors). Epidemics on small-world networks can exhibit phase transition behavior; that is, there is a critical value of the disorder parameter ($p_c$) such that for values of $p_{WS} > p_c$ self-sustained oscillations in the number of infected individuals in susceptible-infected-susceptible (SIS) epidemics are possible (Kuperman and Abramson [45]).

In order to study the role of the disorder parameter $p_{WS}$ (small-world networks) on the initial rate of growth of disease spread, the following algorithm is used to compute the initial (empirical) rate of growth. The number of infected individuals is computed as a function of time $I(t)$ (time is discrete) for a small time range ($t < t_c$). The value of $t_c$ is selected so that $I(t)$ is still in its exponentially growing phase. The algorithm follows three steps:

1. **Computation of $t_c$.** The time $t_c$ at which $I(t)$ changes concavity, that is, the value of $t$ at which the second derivative of $I(t)$ changes from positive to negative (see Figure 2.12).

2. **Rescaling of $I(t)$.** $\hat{I}(t) \equiv \log_e[I(t)]$ for $t < t_c$, where $t_c$ is the value computed in Step 1.

3. **Regression on $\hat{I}(t)$.** Compute the average slope $r$ of the best fitting line to $\hat{I}(t)$. $r$ is the average “$r$” that results from 50 realizations. The average initial rate of growth $r$ is computed as a function of the disorder parameter ($p_{WS}$) of small-world networks ($p_{WS} \in [0, 1]$ is changed in increments of 0.01).

![Figure 2.12. Computation of the empirical rate of growth of epidemics on networks.](image-url)
2.3.1 Epidemiological Model

We consider an stochastic SIR epidemiological model. Hence, individuals can be in one of three epidemiological states: susceptible (S), infected (I), or recovered (R). A susceptible individual in contact with \( i \) infectious individuals may become infected in a short period of time \( \delta t \) with a probability given by \( \hat{\beta}i\delta t \), where \( \hat{\beta} \) is the constant risk of infection per unit of time and \( \delta t = 1 \) in this discrete time model. Similarly, infected recover with a probability given by \( \gamma \delta t \), where \( \frac{1}{\gamma} \) is the mean period of infectivity. After recovery, individuals get full immunity to the disease.

![Figure 2.13](image)

**Figure 2.13.** Rate of growth of epidemics in small-world networks of size \( N = 10^4 \) with \( K = 2, K = 3, \) or \( K = 5 \) and as a function of the disorder parameter \( p \in [0, 1] \). Averages are taken from 50 realizations. \( p \) is incremented by 0.01. Disease parameters are \( \hat{\beta} = \frac{4}{7}, \gamma = \frac{2}{7}, \) and \( I(0) = 1 \).

Epidemics were simulated on small-world networks of size \( N = 10^3 \) with \( K = 2, K = 3, \) and \( K = 5 \) (\( K \) is the coordination number of small-world networks). Empirical results on the average rate of growth were obtained from the mean of 50 realizations with disease parameters \( \hat{\beta} = \frac{4}{7} \) and \( \gamma = \frac{2}{7} \). Simulations were started by placing a single infectious source randomly at a node. The average rate of growth increases in a nonlinear fashion as the disorder in the network grows. It saturates when it is close to 1 (totally random networks) with \( r_{\text{random}} \approx 0.7481 \) and \( K = 2 \). Figure 2.13 shows the average (from 50 realizations) rate of growth of epidemics in small-world networks of size \( N = 10^3 \) (with \( K = 2, K = 3, \) and \( K = 5 \)) as a function of the disorder parameter \( p_{\text{WS}} \). The rate of growth in small-world networks also increases as the coordination number (\( K \)) increases.

The initial rate of growth of epidemics depends on the network topology. The simulations start with an initial (small) group of infective individuals chosen from those with highest connectivity. The resulting growth rate is computed and compared to that resulting from epidemics where the initial infectious sources are chosen (uniformly) at random. Naturally, epidemics that started at the most connected nodes exhibited a higher average rate of growth (see Figure 2.14). Higher rates of growth are observed as the number of initial infec-
Figure 2.14. Rate of growth is higher when the epidemics start at the individuals (nodes) with the highest connectivity (*) rather than chosen uniformly at random (.) with $N = 10^4$ and $K = 2$. Averages are taken from 50 realizations. Disease parameters are $\beta = \frac{4}{3}$, $\gamma = \frac{3}{2}$, and $I(0) = 1$.

tious sources (always small compared to the size of the network) in the network increases (see Figure 2.15).

2.4 Epidemics on Scale-Free Networks

Pastor-Satorras and Vespignani [46] studied an SIS epidemic model on scale-free networks (generated using the BA model) and found that the disease may persist independently of its transmissibility. That is, the basic reproductive number $R_0$, routinely computed in classical mathematical epidemiology sometimes loses its meaning in their setting. The small number of nodes with a high connectivity (hubs) observed in scale-free networks are responsible for “zero” threshold behavior. This observation gives rise to the following question: Can a control strategy be implemented that restores a positive epidemic threshold?

This question was studied by Pastor-Satorras and Vespignani [47] and independently by Dezso and Barabási [48]. Both groups concluded that targeted immunization campaigns towards the most connected nodes or hubs increase the probability of recovering finite epidemic threshold behavior. A similar result has been observed on the spread of foot and mouth disease (FMD) in three distinct regions of Uruguay (Rivas et al. [49]). A contrasting result has been established on alternative highly clustered scale-free networks [35]. Here, a finite epidemic threshold has been observed on (SIS) epidemics (Eguiluz and Klemm [50]).

An extensive number of simulations have been carried out of SIR epidemics on scale-free networks. We compute the average rate of growth from the mean of 50 realizations of the epidemic process with two sets of initial conditions: We place the infective source at a randomly selected node or at the most connected node (highest degree). Simulations are carried out on small-world and scale-free networks of the same size ($N = 10^4$) and
average connectivity ($\bar{k} = 4$). Significantly higher rates of growth are observed in scale-free networks (see Figure 2.16). Hence, scale-free topologies seem to provide ideal scenarios for the study of worst-case epidemics. The existence of highly connected nodes or hubs in scale-free networks plays a central role on the rate at which viruses (or information) spreads. Or, in other words, the concept of core group [29] is still critical to disease spread on topologically defined networks.

2.5 Conclusions and Caveats

The development and implementation of policies that deal with the deliberate release of biological agents must consider worst possible situations and such scenarios are highly dependent on the network of individual interactions (social topology). Hence, gaining some understanding of the nature of social structures that facilitate disease spread is critical. Kaplan et al. [10] assume that random mixing corresponds to a worst-case scenario and (using such a setup) conclude that, in the case of a bioterrorist smallpox attack, mass vaccination is a better policy than ring vaccination. Halloran et al. [51], using a stochastic model with a structured community of 2000 people, conclude that targeted vaccination outperforms mass vaccination. The disagreement in results may be directly related to the preassumed population structure and mixing topology (network of interactions).

Here, we have tried to identify under what conditions random mixing can be used to model worst-case scenarios. We have found that on small-world networks, random mixing indeed supports epidemics with the highest average rate of growth. However, this is not necessarily the case on scale-free networks. The nature of the mixing between individuals (the connectivity hierarchy in scale-free networks) plays a key role on the initial average
Figure 2.16. Average number of infected individuals from 50 realizations over time in small-world ($p = 0.1$) and scale-free networks of the same size ($N = 10^4$) and average connectivity ($k = 4$). Two different initial conditions are considered: The initial infectious source is placed in a randomly selected or in the most connected node (highest degree). The rates of growth of the epidemics are higher in scale-free networks. Disease parameters are $\beta = \frac{2}{7}$, $\gamma = \frac{2}{7}$, and $I(0) = 1$.

rate of growth of an initial epidemic outbreak. The inherent connectivity hierarchy of scale-free networks and the “sensitivity” (lack of robustness) to the removal of key nodes (most connected individuals), in some sense, corresponds to the concept of core group [29]. Highly connected nodes are pressure points in the network, and consequently, their identification and management must be considered in the development and implementation of a logistic plan of response to the threat of a bioterrorist attack.

The location-based network of the simulated city of Portland possesses a scale-free nature (Chowell et al. [25]). Hence, the initial rate of growth of epidemics in this city can be expected to be (on average) significantly higher whenever a source is placed at a hub (see Figure 2.16).

While the use of classical epidemiological approaches has been and will continue to be central in the study of disease spread [5, 1, 4], it is clear that the study of the potential initial impact of the deliberate release of pathogens on unsuspecting populations must be addressed on various setups. The reasons for such an approach are multiple, the interest is no longer on long-term disease dynamics but on the elaboration of policies that result on timely responses (see Rivas et al. [49]). Such a degree of urgency requires the identification of the most sensitive points of release (pressure points) and the use of models that account for multiple releases. The approach used here has its own drawbacks. The most important comes from the fact that temporal dynamics are not explicitly considered. The incorporation of temporal dynamics on networks poses challenges and opportunities for serious theoretical work. Although we have fallen short in addressing the challenge posed by Ed Kaplan, we hope that our results have at least helped clarify it because the development of contingency plans in worst-case scenarios is fundamental.
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