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Adaptive network modeling of social distancing interventions

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ABSTRACT

The COVID-19 pandemic has proved to be one of the most disruptive public health emergencies in recent memory. Among non-pharmaceutical interventions, social distancing and lockdown measures are some of the most common tools employed by governments around the world to combat the disease. While mathematical models of COVID-19 are ubiquitous, few have leveraged network theory in a general way to explain the mechanics of social distancing. In this paper, we build on existing network models for heterogeneous, clustered networks with random link activation/deletion dynamics to put forth realistic mechanisms of social distancing using piecewise constant activation/deletion rates. We find our models are capable of rich qualitative behavior, and offer meaningful insight with relatively few intervention parameters. In particular, we find that the severity of social distancing interventions and when they begin have more impact than how long it takes for the interventions to take full effect.

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1. Introduction

The global COVID-19 pandemic has upended modern life an placed an enormous epidemiological, economic, and social burden on the world’s resources. The gravity of events has brought the need for epidemiological modeling into sharp focus. As the pandemic spread around the world in the absence of a vaccine, non-pharmaceutical interventions including social distancing, quarantine, and lockdown measures proliferated, and bringing these interventions into modeling efforts has remained paramount.

In recent years, network-based models of epidemic spread have become an increasingly popular paradigm (Pastor-Satorras et al., 2015; Kiss et al., 2017), and network science generally has been recognized for its potential to contribute solutions to the current crisis (Eubank et al., 2020). Most network models represent individuals as nodes in a network, and their contacts as edges connecting the nodes. Moreover, many models assume that the network is static—that the edges between nodes don’t change over time—and thus the epidemic spreads from node to node across these edges. Among static network models, pairwise models (Keeling, 1999; Eames and Keeling, 2002) are both frequently used and well-studied. Pairwise models track not only the number of nodes in a given state, but pairs, triples, and higher order motifs as well (Fig. 1). An advantage of pairwise models is that in their full form, they exactly model (in expectation) the continuous time Markov chain formulation of epidemic spread on a network (Taylor et al., 2012).

Pairwise models have been successfully applied to a number of disease natural histories and different network types. Two important network features that play a role in the theory of pairwise models are degree heterogeneity and clustering. The degree of a node in a network is the number of edges to which it is connected, and the degree distribution is the probability distribution of selecting a random node with a given degree. The degree distribution plays a fundamental role in many network models, and is particularly powerful when described as a probability generating function. The clustering coefficient is the ratio of triangles to connected triples in the network. While clustering is an important component of network structure, it has not widely been incorporated to pairwise models. We acknowledge two major benefits of degree heterogeneous, clustered models. First, including both or either as modeling consideration affects epidemic dynamics in a nontrivial way (House and Keeling, 2011; Keeling, 1999) and second, both have been shown to be features of realistic contact networks (Read et al., 2008).

Though static networks model some forms of complexity well, an important aspect of real-world contact networks is that some connections change in response to disease dynamics or public health measures. By relaxing the static network assumption, dynamic or “adaptive” network models (Gross and Sayama, 2009) can capture both the dynamics of the network and the epidemic dynamics on the network. A number of models have been recently proposed that describe a variety of network dynamic pro-
cesses. Gross et al. (2006) introduced a model of edge rewiring, where susceptible individuals break connections with infectious individuals and reconnect to susceptible individuals at random. A related model of adaptive dynamics is “relational exchange” (Scarpino et al., 2016), where an node in contact with an infectious node are rewired to a susceptible node. Another model for network dynamics is random link addition/deletion (Kiss et al., 2012) where individuals break and form new contacts and constant rates. Their approach is notable for its intuitiveness as a simple dynamic model, and also its use of probability generating functions as a tool to describe network dynamics. A related model is link addition/deletion on a fixed network (Tunc et al., 2013; Shkarayev et al., 2014), where individuals can temporarily deactivate contacts with infectious individuals, and reactivate them when their contact is not infectious. While much of the focus of the adaptive network literature has been involved in analyzing the resulting dynamical systems, particularly for SIS-type diseases, some works have focused on the role of network dynamics in controlling or mitigating epidemic spread (Youssef and Scoglio, 2013; Sélley et al., 2015).

Network models in general offer a compromise between two other common modeling techniques: compartment models and agent-based simulations. They are able to capture more complex contact structure than simple compartment models, while offering analytical tractability that many agent-based simulations lack. Despite this, models of non-pharmaceutical interventions have tended to favor simulation or compartment models (Ahmed et al., 2018; Davey et al., 2008). In the early stages of the COVID-19 pandemic, complex individual-based simulations offered major insights about the effectiveness of non-pharmaceutical interventions (Ferguson et al., 2020). However, the high computational cost can make investigating the impacts of intervention policies with a large number of parameters a challenging endeavor. Network models, especially those with a relatively small number of equations, can offer broad insights at reduced cost. While some models of social distancing have incorporated contact network structure as a major consideration (Valdez et al., 2012; Glass et al., 2006), differential equation network models of such interventions are uncommon. And while other authors have investigated similar pressing questions posed by the COVID-19 pandemic (Lauro et al., 2021, for one), many model social distancing through a time-varying transmission rate. Adaptive network models have the potential to model social distancing directly, and can offer a new perspective on questions surrounding social distancing and other non-pharmaceutical interventions made pressing by the COVID-19 pandemic.

In this paper, we develop simple, novel mechanisms to incorporate social distancing into a network model of epidemic spread, using COVID-19 as the central case study to investigate the impact of a range of interventions. The goal of this paper is to model the effects of social distancing that can be understood on a social level, as well as an epidemiological level. Using network dynamics, we develop social distancing schemes by adding and deleting edges from the network at specified rates. The resulting social distancing schemes can be concretely understood as reducing (and eventually increasing) the average number of contacts of nodes in the network, rather than the more abstract alternative of reducing (and eventually increasing) the transmission rate. Our approach offers potential fruitful avenues of model calibration—with increasing availability of mobile phone mobility data, modeling social distancing policies directly through network dynamics may yield fruitful and flexible real-world models. This paper offers an early theoretical step in such a formulation.

First, we develop a pairwise SEIR model with random link activation/deletion dynamics—that is edges are added and deleted at constant rates independent of the epidemic dynamics on the network. Furthermore, the model incorporates degree heterogeneity and clustering, which offers increased realism over simpler network or compartment models. To apply the model, we use bipartite mixing networks to generate large heterogeneous, clustered contact networks coupled with disease dynamics given by epidemiological parameters estimated for COVID-19. Next, we develop two mechanisms of social distancing using piecewise constant link activation and deletion rates. The first is a single intervention event, where the average number of contacts decreases, is held constant, and then recovers; the second allows for multiple interventions which restart depending on the prevalence of the disease. While we investigate the implications of these policies for COVID-19 on a specific type of heterogeneous, clustered network, both the adaptive network model and the social distancing schemes are more generally applicable to a variety of networks and epidemiological parameters. Finally, we consider the public health implications of the latter model, finding that certain intervention parameters are more important than others in achieving an effective reduction in overall infections.

2. Model

To begin construction of the full model, we consider SEIR dynamics on a static network. Pairwise equations for an SEIR epidemic can be found in Keeling et al. (1997) and Rand (1999). Model variables include the expected number of susceptible, exposed, infectious, and recovered nodes (S, E, I, and R respectively) as well as the expected number of pairs in each state. For example, $\langle SS \rangle$ is the expected number of connected pairs of susceptible nodes, while $\langle SI \rangle$ is the expected number of connected pairs of susceptible and infectious nodes. The expected number of connected triples is also considered $\langle SSI \rangle$, although differential equa-

![Diagram](image.png)

**Fig. 1.** Diagrams of network structures whose evolution is modeled by the pairwise model: (a) node in state $A$, (b) pair in state $A-B$, (c) triple in state $A-B-C$. 
tions for these variables are not written. The full SEIR pairwise model is
\[ S = -\beta[S][E], \]  
\[ E = \beta[S][E] - \gamma[E], \]  
\[ [I] = \eta[E] - \gamma[I], \]  
\[ [SS] = -2\rho[SS], \]  
\[ [SE] = \beta[SE] - \beta[ES] - \eta[SE], \]  
\[ [SI] = \eta[SE] - \beta[SI] - \beta[IS] - \gamma[SI], \]  
\[ EE = 2\beta[ES] - 2\eta[EE], \]  
\[ [EI] = \beta[IS] + \beta[SE] + \eta[EE] - (\gamma + \eta)[EI], \]  
\[ [II] = 2\eta[E] - 2\gamma[I], \]  
where \( \beta \) is the transmission rate, \( \gamma \) is the recovery rate, and \( \eta \) is the rate at which exposed individuals become infectious. The nodes and edges also obey conservation equations
\[ N = [S] + [E] + [I] + [R] \]  
and
\[ \langle k \rangle N = [SS] + [EE] + [II] + [RR] + 2\langle SE \rangle + [SS] + [SR] + [EI] + [ER] + [RR] \]  
where \( N \) is the number of nodes and \( \langle k \rangle \) is the average degree of the network. We note that with the conservation equations, we do not need terms of the type \( [AR] \) to determine the evolution of \([S],[E],[I], \) and \([R] \).

The full model requires dynamical equations for triples of the form \([AS] \) and higher order motifs as well, leading to a system that is prohibitively large for computations. To make the model tractable, we approximate the expected number of triples \([AS] \) in terms of pairs and individual nodes, thus closing the system (1)-(9). An approximation of this kind is referred to as a triple closure. For triples of the type \( A-S-I \), using the techniques of House and Keeling (2011), we begin with a general triple closure approximation
\[ [AS] \approx \frac{\sum (k^2 - k)[S]}{\sum k[S]} \left( 1 - \phi + \phi \frac{N[k][A]}{\sum k[A]} \right), \]  
where \( [A] \) is the expected number of nodes in state \( A \) with degree \( k \) and \( \phi \) is the clustering coefficient of the network. Tracking \([A] \) for any of the disease states \( A \) requires a significantly larger system of equations, so finding a workaround is a worthwhile goal. To simplify (12), we approximate \( \sum_a (k^2 - k)[A] = \langle k^2 \rangle - \langle k \rangle [A] \) and \( \sum_k k[A] = \langle k \rangle [A] \). For any disease state \( A \), where \( p_k \) is the proportion of nodes with degree \( k \), \( \langle k \rangle = \sum_k k p_k \) is the average degree, and \( \langle k^2 \rangle - \langle k \rangle = \sum_k (k^2 - k) p_k \). The resulting triple closure is
\[ [AS] \approx \frac{\langle k^2 \rangle - \langle k \rangle}{\langle k \rangle} \frac{\sum (k^2 - k)[S]}{\sum k[S]} \left( 1 - \phi + \phi \frac{N[k][A]}{\langle k \rangle [A][A]} \right). \]  

To the authors knowledge, this closure is novel; however it bears great similarity to a triple closure given in Keeling (1999), which replaces \( \langle k^2 \rangle - \langle k \rangle \) and \( \langle k \rangle - 1 \) by \( \langle k \rangle \) and \( \langle k^2 \rangle \). The two are equivalent if \( \langle k^2 \rangle = \langle k \rangle^2 \), as is the case with homogeneous networks. For heterogeneous networks, we propose (13) is an improvement, and we will use (13) to close the system (1)-(9).

While we will use a simplified approximation to (12) we note for interested readers that House and Keeling (2011) offer an approach to use (12) exactly. They use the probability generating function of the degree distribution \( g(x) = \sum x^k p_k \) and new dynamical variables such as \( \theta \), the probability that an edge has not transmitted the infection and \( Y \), the sum \( \sum_t k_i \) to develop an SIR model for heterogeneous, clustered networks. In the Supplementary Material, we develop an analogous heterogeneous, clustered SEIR model complete with link activation and deletion. However, the added complexity of the model is not necessary for the primary investigations of this paper, and we continue developing the SEIR model using the triple closure (13).

With the static model closed, we now incorporate the effects of network dynamics. Kiss et al. (2012) introduced a simple model of network dynamics, termed random link activation/deletion (RLAD). In this model, independent of epidemic dynamics nonexistent edges are added to the network (or activated) at a constant rate \( \alpha \) and existing edges are removed from the network (or deleted) at a constant rate \( \omega \). Ignoring epidemic spread and looking at the effects of activation/deletion only, the equation for edges of type \([AB] \) is
\[ [AB] = \alpha([A][B] - [AB]) - \omega[AB], \]  
and for type \([AB] \) we have
\[ [AB] = \alpha([A][B] - [AB]) - \omega[AB]. \]  

Next, we have to consider the effect of activation/deletion on the now time-dependent network quantities: degree distribution moment terms \( \langle k \rangle(t), \langle k^2 \rangle(t) \) and the clustering coefficient \( \phi(t) \). Following the example of Kiss et al. (2012), dynamical equations for the first two can be easily derived by finding the partial differential equation for the degree distribution generating function
\[ g(x,t) = \sum_{k=0}^{N-1} p_k(t)x^k. \]  

The Kolmogorov equations describe the evolution of \( p_k(t) \), the proportion of degree \( k \) nodes at time \( t \):
\[ p_k(t) = \alpha(N - k)p_{k-1} - (\alpha(N - 1 - k) + \omega k)p_k + \omega(k + 1)p_{k+1}. \]  

With some straightforward algebra, we derive a partial differential equation for the degree distribution generating function:
\[ \frac{\partial g}{\partial t} = (x - 1) \left( \alpha(N - 1)g - (\alpha x + \omega) \frac{\partial g}{\partial x} \right). \]  

The network quantities \( \langle k \rangle \) and \( \langle k^2 \rangle \) can be computed from the generating function as \( \langle k \rangle = g_x(1,t) \) and \( \langle k^2 \rangle = g_{xx}(1,t) \). Then, from (18) we derive the dynamical equations:
\[ \langle k \rangle = \alpha(N - 1) - (\alpha + \omega)\langle k \rangle, \]  
\[ \langle k^2 \rangle = 2\alpha(N - 2)\langle k \rangle - 2(\alpha + \omega)(\langle k^2 \rangle - \langle k \rangle). \]  

The clustering coefficient is defined as the ratio of triangles to connected triples in the network. To compute \( \phi \), we start with the Kolmogorov equations for \( q_k(t) \), the probability that there are \( k \) triangles in the network at time \( t \):
\[ q_k = \alpha(L - 3(k - 1))q_{k-1} - (\alpha(L - 3k) + 3\omega k)q_k + 3\omega(k + 1)q_{k+1}, \]  
where \( L = N\langle k^2 \rangle / 2 \) is the number of connected triples. From this we derive the differential equation for the expected number of triangles \( \langle T \rangle \) as
\[ \langle T \rangle = 2\alpha L - 3(\alpha + \omega)\langle T \rangle. \]  

We note that writing (22) as \( \langle T \rangle = \alpha(L - 3T) - 3\omega T \) offers a heuristic interpretation of the evolution of triangles: there are
The model equations are less accurate (see Supplementary Material for examples).

Finally, we have a full set of equations for a pairwise SEIR for a heterogeneous, clustered network with random link activation and deletion:

\[
\begin{align*}
\dot{S} &= -\beta SI, \\
\dot{E} &= \beta SI - \eta E, \\
\dot{I} &= \eta E - \gamma I, \\
\dot{R} &= \gamma I - \lambda R - \omega R, \\
\dot{\phi} &= 3\lambda - \left(\alpha + \omega + 2\lambda(N-2)\frac{(k)}{(k^2 - k)}\right)\phi.
\end{align*}
\]

where

\[
\begin{align*}
\beta &= \frac{1}{C_0}L, \\
\lambda &= \frac{1}{C_0}aH, \\
\gamma &= \frac{1}{C_0}T, \\
\phi &= \frac{1}{C_0}\text{deg}, \\
\eta &= \frac{1}{C_0}\text{deg}, \\
\omega &= \frac{1}{C_0}\text{deg}, \\
\lambda &= \frac{1}{C_0}\text{deg}.
\end{align*}
\]

In Figs. 2, we show the results of these equations compared to simulations, and find that they are in excellent agreement. However, we note that there are circumstances where this agreement is not so good. For sparse networks (low average degree), the agreement is not as strong as compared to networks with higher average degree. Moreover, for intervention schemes that involve changing values of \(\alpha\) and \(\omega\), such as those described in Section 3, the model equations are less accurate (see Supplementary Material for examples).

Finally, we have a full set of equations for a pairwise SEIR for a heterogeneous, clustered network with random link activation and deletion:
To demonstrate the validity of this model, we test it against numerical simulations (Fig. 2) on a heterogeneous, clustered network—the construction of which is described in Section 2.1. Clearly, the model (24)-(35) is in excellent agreement with the simulations.

2.1. Network and Epidemiological Parameters

The goal of this paper is to investigate social distancing policies through random link activation/deletion dynamics, which are controlled by the activation and deletion rates $\alpha$ and $\omega$. Moreover, in building intervention schemes in Section 3 new parameters are introduced. In order to consistently compare the efficacy of intervention schemes, network and epidemiological parameters are held the same across schemes. As such, we restrict our attention to a particular heterogeneous, clustered network and epidemiological parameters that are plausible for COVID-19. For completeness, other network types and epidemiological parameters are considered in the electronic supplementary material.

A consistent challenge of network models is constructing realistic contact networks. In particular, degree heterogeneity and significant clustering are observed in real world social networks (Read et al., 2008). To construct such a contact network, we consider a bipartite mixing network (Eubank et al., 2004) with $N$ individuals and $M$ mixing locations (Fig. 3a). Two individuals are in contact if they both connect to the same mixing location, so we form a contact network as the unipartite projection of the bipartite mixing network (Fig. 3b). To introduce degree heterogeneity, we construct a bipartite mixing network where both individuals and mixing locations have Poisson degree distributions Newman.
et al. (2001). The average individual degree $\bar{z}$ and average mixing location degree $\mu$ are related by

$$N\bar{z} = NM\mu,$$  

so only $N$, $M$, and $\bar{z}$ are needed to characterize this network. Using generating function techniques (Newman et al., 2001), we compute

$$\langle k \rangle = \frac{N}{M} \bar{z},$$

$$\langle k^2 - k \rangle = \left( \frac{N}{M} \right)^2 \bar{z}^2 (\lambda + 1),$$

$$\phi = \frac{1}{\lambda + 1},$$

for the unipartite contact network, which exhibits both degree heterogeneity (Fig. 3b) and clustering. Unless otherwise specified, the networks in this article are generated using $N = 10,000$, $M = 2,500$, and $\bar{z} = 4$. We acknowledge that though we use a bipartite mixing network to generate a heterogeneous, clustered unipartite contact network, our network dynamics are limited to the contact network. Mobility networks (Chang et al., 2021) have been used to great effect for COVID-19, and suggest a fruitful path forward for bipartite network dynamics.

Numerous recent studies have estimated important epidemiological quantities for the spread of Sars-CoV-2, including the length of the incubation period, the length of the infectious period, and the basic reproduction number $R_0$. We choose the plausible estimates in line with recent studies: average incubation period of 5 days (Linton et al., 2020; Zhang et al., 2020), average infectious period of 10 days (You et al., 2020), and $R_0 = 2.4$ (Li et al., 2020; Anastassopoulou et al., 2020). To incorporate these into the model, we note that $1/\eta$ and $1/\gamma$ are the average lengths of the incubation and infectious periods respectively, and thus $\eta = 0.2$, $\gamma = 0.1$. We do not derive $R_0$ for the model (24)-35, but instead consider the basic reproduction number for a heterogeneous, clustered population from Miller (2009), which is given as the series

$$R_0 = \frac{\langle k^2 - k \rangle}{\langle k \rangle} \frac{\beta}{\beta + \gamma} - \phi \frac{\langle k^2 - k \rangle}{\langle k \rangle} \left( \frac{\beta}{\beta + \gamma} \right)^2 + \cdots$$

Ignoring higher order terms, we can compute $\beta$ from (43) when $R_0 = 2.4$. With these parameters, we plausibly model the spread of COVID-19 through a moderately sized heterogeneous, clustered population in the following sections, while introducing various social distancing interventions to mitigate or control the epidemic.

3. Analysis

Social distancing and lockdown measures have been used to curb the spread of infectious diseases throughout history, and are some of the most ubiquitous non-pharmaceutical interventions in the current COVID-19 pandemic. Many compartmental models that incorporate social distancing do so in a phenomenological manner through the transmission rate, but adaptive network models present an opportunity to describe a social distancing mechanism in a fundamental way. A simple model of such interventions can be naturally characterized by the link activation/deletion process. During periods of social distancing and lockdown, individual contacts break; during periods of relaxation of the measures, individual contacts form. In this section, we develop two social distancing schemes (Fig. 4) based on the prevalence $I(t)$. Other models of social distancing may be based on other epidemiological indicators and have different goals—for instance, keeping the effective reproduction number $R_e$ below one, or mitigating transmissions by highly-connected individuals. However, the schemes presented here are in response to prevalence alone, seeking to model social distancing policies that respond to solely to case numbers. Both social distancing schemes begin when the prevalence $|I(t)|$ reaches some specified threshold level. For the simple intervention scheme, contacts break as the intervention is implemented, then contacts stay fixed as the intervention is in place, and finally contacts form until they reach their pre-intervention levels. The prevalence-dependent scheme unfolds similarly, but with two notable differences. First, after the intervention, contacts do not start forming again until the prevalence has dropped below the threshold. Second, any time the prevalence reaches the threshold again, the intervention restarts. This allows for multiple implementations of a social distancing intervention throughout the course of the epidemic.

Critically, we do not treat these schemes as a mere modeling exercise, but are interested in the impact of each intervention scheme at the end of the epidemic. We develop two simple metrics to evaluate the effectiveness of the simple and prevalence-dependent interventions. First, we consider each intervention’s ability to reduce the cumulative number of infections, known as the final size of the epidemic. Second, we also consider how many infections occur above the threshold value for prevalence. These two measures reflect two different yet crucial public health goals, and do not necessarily agree on which interventions are the most effective. Both must be considered to get a complete picture of an intervention’s impact. In this section, we derive these two metrics mathematically, and describe the simple and prevalence-dependent interventions while assessing their overall effects.

3.1. Evaluation Metrics

The first measure of intervention effectiveness we introduce is the Relative Change in the Final Size (RCFS). The final size of an epidemic is the cumulative number of infections that occur over the course of the epidemic. In terms of the model, the final size can be found as the limiting value of the recovered individuals $|R|$: 

$$\lim_{t \to \infty} |R(t)| = R_{\infty}.$$ 

We compare the final size of the epidemic with no intervention $R_{\infty}$ to the final size where an intervention has been implemented $R_{\infty}^{\text{int}}$. We then define the RCFS as

$$\text{RCFS} = \frac{R_{\infty}^{\text{int}} - R_{\infty}}{R_{\infty}}.$$  

An effective intervention will lead to a decrease in final size, so an RCFS near 0 is unsuccessful, while an RCFS near −1 is extraordinarily successful. However, it is important to note that for brief, intense intervention schemes, it is possible that the final size actually increases. In this case, the network parameters change quickly, before significant disease spread, so the epidemic unfolds on a fundamentally different static network.

While the relative change in the final size provides an overall measure of the effectiveness of interventions, reducing cumulative infections alone is not the only public health goal that an intervention scheme might seek to accomplish. In some schemes, a large number of infections occur above the threshold despite a large reduction in the final size of the epidemic. This can be particularly pernicious if the threshold represents some fixed resource such as healthcare capacity, where a large number of infections above the threshold could lead to higher mortality and other negative outcomes. To account for this, we compute the Cumulative Infections Above Threshold (CIAT). Infections are above threshold whenever $|I(t)| - qN \geq 0$, and thus the CIAT can be defined by

$$\text{CIAT} = \int_0^\infty \max (|I(t)| - qN, 0) \, dt.$$
**Simple Intervention**

For computation purposes, we can characterize (45) summing the integrals over time periods when \(|I| - qN ≥ 0\). Let \(t_1, t_2, \ldots\) be the sequence of times when \(|I| = qN\). Assuming \(|I| ≠ 0\) at any time in the sequence, the continuity of \(|I|(t)\) implies that the prevalence is above the threshold on the intervals \([t_{2i-1}, t_{2i}]\) for \(i = 1, 2, 3, \ldots\). Thus, the CIAT is

\[
\text{CIAT} = \sum_{i=1}^{\infty} \int_{t_{2i-1}}^{t_{2i}} |I|(t) - qN dt. \tag{46}
\]

We note that the units of CIAT are person-time—for a metric with units of population, we compute the Average Infections Above Threshold (AIAT):

\[
\text{AIAT} = \sum_{i=1}^{\infty} \int_{t_{2i-1}}^{t_{2i}} |I|(t) \tag{47}
\]

Using the relation \(\sum |R|_i = \gamma I_0\), Eq. (47) becomes

\[
\text{AIAT} = \frac{\sum_{i=1}^{\infty} |R|(t_{2i}) - |R|(t_{2i-1})}{\gamma \sum_{i=1}^{\infty} t_{2i} - t_{2i-1}} - qN, \tag{48}
\]

which is convenient for computations.

**Prevalence-Dependent Intervention**

To achieve this effect in the evolution of the average number of contacts, we consider link activation rate \(\alpha(t)\) and deletion rate \(\omega(t)\) that are piecewise constant. These rate functions can be seen in the bottom two panels of Fig. 5. Since contacts are only broken in the first phase, \(\omega(t) = \omega'\) in the first phase and \(0\) otherwise. Since contacts are only formed in the third phase, \(\alpha(t) = \alpha'\) in the third phase and \(0\) otherwise. As the dynamical function for \(\langle k \rangle(33)\) is a first-order linear ODE, the resulting curve for \(\langle k \rangle(t)\) will be piecewise exponential, and the values of \(\alpha'\) and \(\omega'\) are easily computed for a given \(p\). Other than \(p\), four other parameters characterize the simple intervention: the lengths of the three phases \(L_i\), \(L_h\), and \(L_r\), and the threshold proportion of the population \(q \in [0, 1]\) to initiate the intervention. The full simple intervention scheme can be described as follows:

- **No intervention:** The epidemic spreads unabated until \(|I|\) increases through \(qN (\alpha = \omega = 0)\).
- **Intervention Phase (length \(L_i\)):** Intervention occurs, edges are removed at a constant rate \((\alpha = 0, \omega = \omega')\).
- **Holding Phase (length \(L_h\)):** Intervention holds, edges are neither removed nor added \((\alpha = \omega = 0)\).
- **Relaxation Phase (length \(L_r\)):** Interventions are relaxed, edges are added at a constant rate \((\alpha = \alpha', \omega = 0)\).

To emphasize the value of using activation and deletion rates to model social distancing, we compare the social distancing model under consideration to two alternate approaches. First, we consider the static network equivalent of the model (24)-(35), which requires only \(24)-(32)\) and \(\alpha = \omega = 0\). Second, we consider the standard SEIR model with homogeneous mixing. To model social distancing in each case, we allow for the transmission rate \(\beta\) to be time-varying, and in particular we vary it proportionally to \(\langle k \rangle(t)\) for the adaptive model. That is to say, if we let \(\beta_{ad}\) denote the constant transmission rate in the adaptive network model, the time-varying transmission rate for the alternative models is \(\beta(t) = \beta_{ad} \langle k \rangle(t)\). Fig. 6 compares and contrasts the social distancing mechanisms as \(\langle k^2 - k \rangle(0)\) varies. In Fig. 6a, where
The adaptive network model. This suggests that not only is incorporating network structure into the model essential, but also that controlling social distancing through an adaptive process instead of merely a time-varying transmission rate is important for increasingly heterogeneous networks.

As the simple intervention scheme requires five "intervention" parameters, p, q, LI, LR, and LH, exploring the full impact of the interventions is difficult. To better see the effects, we consider an example scheme where we fix two parameter values in each and allow the other three to vary. To focus on the impact of the severity parameter p and the lengths of the intervention and relaxation phases LI and LR, we set LI = 15 and q = 0.01 for the remainder of this section. Thus, the intervention begins when infections reach one percent of the population, and the holding phase is fixed at 15 days for all interventions. The other three parameters are allowed to vary. This allows for both abrupt and gradual implementations of interventions and relaxation of measures, and different levels of intervention intensity. Fig. 7 shows the prevalence of some example intervention schemes, showing rich qualitative behavior. To assess the effectiveness of the simple intervention we plot the RCFs and the AIAT for a large number of parameter combinations. We allow the lengths of both the intervention and relaxation periods LI and LR to vary from 2 to 180 days, and consider three different intensities p = 0.125, 0.25, 0.5. The results are shown in Fig. 8.

A significant common feature of the plots in Fig. 8a is a qualitative boundary (solid white) that divides (LI, LR) space into two distinct classes of the resulting infection curve (for p = 0.125, this occurs outside the boundaries of the plot). To the right of the boundary, infection curves are characterized by a single "uniform spike," defined by an prevalence curve \( I(t) \) with two inflection points and a single local maximum (Fig. 9a). To the left of the boundary, infection curves take the form of either a single "nonuniform spike" (Fig. 9b), with more than two inflection points but only one local maximum, or multiple spikes (Fig. 9c), with more than two inflection points and multiple local maxima. For p = 0.125 and p = 0.25, only multiple spikes occur to the left of the boundary. For small LI, the first spike is small and the second spike is large, and occasionally the final size of the epidemic surpasses the static case due to network alterations. As LI approaches the qualitative boundary, the second spike becomes shorter and occurs later until negligible. This phenomenon can also be seen in Fig. 8b: as LI increases until the second spike drops below the threshold qN, at which point the AIAT increases as the first spike grows taller. For p = 0.5 on the other hand, both nonuniform spikes and multiple spikes are possible to the left of the boundary. Multiple spikes occur in the region of (LI, LR) space enclosed by the dashed white curve, while a single nonuniform spike occurs elsewhere left of the qualitative boundary.

A few other observations warrant comment. First, the length of the intervention LI appears to be more important in determining epidemic's final size compared to LR. This is intuitive, as the most significant changes to network structure occur during the intervention phase. Second, as p increases, the qualitative boundary shifts generally left. This means that for less severe interventions, single uniform spikes will occur for smaller LI values. This observation carries weight for repeated interventions, explored in Section 3.3, as single uniform spikes are heavily penalized by the AIAT. Third, nonuniform spikes occur for p = 0.5, but not for p = 0.125 or p = 0.25. We hypothesize that there may exist some threshold p' where nonuniform spikes don't occur below p', but do above p'.

### 3.3. Prevalence-Dependent Intervention

While the simple intervention scheme provides a simple yet general model of social distancing, its implementation lacks a degree of realism. Interventions are put into place only once, and the epidemic continues, often with infections spiking after measures begin to relax. In reality, we would expect public health measures to be responsive to rising prevalence. Moreover, continued interventions might be triggered by some indicator, such as case
numbers, deaths, hospital capacity, etc... In this section, we adapt the intervention scheme from Section 3.2 so that it may be reimplementation when a prevalence-based condition is satisfied, forming the prevalence-dependent intervention. We begin with two more realistic assumptions about how a public health response might unfold. First, interventions are reimplemented any time the prevalence increases through some threshold. Second, the relaxation phase of an intervention doesn’t begin until the prevalence has dropped below the threshold. We incorporate these assumptions into a new prevalence-dependent intervention scheme. The scheme is determined by four parameters: \( q, p, L_t, \) and \( L_R \). As before, interventions begin when \( |I| \) reaches \( qN \), \( p \) is the severity of the intervention, and \( L_t \) and \( L_R \) are now the maximum lengths of the intervention and relaxation periods, which determine \( \omega^* \) and \( \chi^* \), as in Section 3.2. We can define the new scheme as follows:

- **As \( |I| \) increases through \( qN \), a new intervention is implemented.**
- **Intervention Phase:** Once an intervention is implemented, edges are deleted at rate \( \omega = \omega^* \) until \( \langle k \rangle = p \langle k \rangle_0 \).
- **Holding Phase:** At the end of the intervention period, a holding period begins \( (\alpha = \omega = 0) \) until the prevalence has dropped below the threshold \( qN \). If the prevalence drops below the threshold during the intervention period, the holding period has length 0.
- **Relaxation Phase:** Edges are added at rate \( \chi = \chi^* \) until \( \langle k \rangle = \langle k \rangle_0 \), or a new intervention is implemented.

It worth noting that compared to the simple intervention in Section 3.2, the intervention, holding, and relaxation phases can all be of variable length. For instance, if the average number of contacts \( \langle k \rangle \) has not rebounded to \( \langle k \rangle_0 \) by the time a new implementation begins, the resulting relaxation period is shorter than \( L_R \). Moreover, in the subsequent intervention phase, edges delete until \( \langle k \rangle = p \langle k \rangle_0 \) and the phase is shorter than \( L_t \). In sum, while \( \omega^* \) and \( \chi^* \) are fixed, the average number of contacts is never less than \( p \langle k \rangle_0 \) and the effective lengths of different intervention and relaxation phases may vary. An example implementation of the prevalence-dependent scheme is shown in Fig. 10, which shows both holding
periods of nonzero length as well as intervention and relaxation periods that are shorter than $L_I$ and $L_R$ respectively. A notable feature of the prevalence-dependent intervention is its ability to generate infection curves with multiple spikes as the epidemic progresses. Examples of this behavior are shown in Fig. 11. To fully explore the intervention, we again consider the RCFS for a variety of parameter combinations. Fig. 12 shows the RCFS for different thresholds ($q = 0.005, 0.01, 0.02$) and intensities ($p = 0.125, 0.25, 0.5$) as $L_I$ and $L_R$ both vary from 2 to 180 days. Though not shown, as with the simple intervention each case has a qualitative boundary, to the right of which infection curves are single, uniform spikes. The most significant departure from the simple intervention though is to the left of the qualitative boundary. In the simple case, infection curves from this region took the form of either two spikes or a single nonuniform spike. With the prevalence-dependent intervention, the infection curve behavior is richer. The region is characterized by “waves” in the RCFS, particularly for lower values of $p$. The boundaries of these waves can be described by the number of spikes that occur over the course of

Fig. 8. Plots of the RCFS (a) and AIAT (b) for the $L_I = 15$ and $q = 0.01$. For intensities $p = 0.125, 0.25,$ and 0.5, the intervention period and relaxation period lengths $L_I$ and $L_R$ vary from 2 to 180 days. In (a), the solid white curve denotes the qualitative boundary, to the right of which uniform spikes occur. The dashed white line in the third panel denotes the boundary of the region where two spikes occur. Initial conditions for the state variables are $S(0) = 9998, E(0) = I(0) = 1, SS = 639784$. $[SE] = 60, [SI] = 48, [EE] = [EI] = [II] = 0$. Epidemiological parameters are $\beta = 0.003, \eta = 0.2, \gamma = 0.1$, and starting network parameters are $N = 10,000, k(0) = 64, (k^2 - k) = 5120, \phi(0) = 0.2$.

Fig. 9. Types of infection curves with the simple intervention: (a) uniform spike, (b) non-uniform spike, (c) multiple spikes. Black dots denote inflection points.
the epidemic. Holding $L_R$ fixed and increasing $L_I$ through one of these contours helps explain the behavior of the infection curve in this region (Fig. 13). At the crest, the final spike peaks just below the threshold $q_N$ (Fig. 13a). As $L_I$ increases, the final spike occurs later and peaks lower (Fig. 13a) and the RCFS decreases until the spike vanishes. Then, the penultimate spike becomes the new final spike, peaking just below the threshold (Fig. 13a) and the RCFS jumps up as a new wave crests. This underscores a potential limitation of a threshold-based intervention: if a spike does not reach the threshold and no intervention occurs, the spike occurs over a longer period of time and more infections accumulate than if the spike had triggered an intervention. A practical implication of this observation is that any spike in infections should not go undressed by interventions if the goal is to reduce the number of cumulative infections. We also consider the AIAT for the same parameter combinations (Fig. 14), though the conclusions by this metric are less complex. For any combination of $p$ and $q$, increasing $L_I$ leads to a larger AIAT. This suggests that when considering interventions with the same RCFS, more abrupt interventions (smaller $L_I$) are preferable. However, an interesting observation is that the

Fig. 10. Prevalence-Dependent Intervention. The intervention begins when $|I| = qN$, and edges are deleted at a constant rate $x$ until $|I|$ decreases to $p|I|$, at which point there is no change to the network until $|I|$ drops below the threshold $qN$. Then, edges are added at a constant rate $\alpha$ until $|I|$ returns to $|I|_0$, or $|I|$ increases through the threshold $qN$, at which point the intervention begins again.

Fig. 11. Example infection curves $|I|(t)$ for the prevalence-dependent intervention. Parameters shown are (a) $q = 0.005, p = 0.125, L_I = 60, L_R = 60$, (b) $q = 0.01, p = 0.5, L_I = 15, L_R = 60$, (c) $q = 0.02, p = 0.25, L_I = 30, L_R = 120$. Solid orange curves are $|I|(t)$ under the intervention, while dashed orange curves are $|I|(t)$ without any intervention. Dashed gray lines denote times when $|I| = qN$. Initial conditions for the state variables are $S(0) = 9998, |E|/C = |I|/C = 1, |SS| = 639784, |SE| = 60, |SI| = 48, |EE| = |EI| = |II| = 0. Epidemiological parameters are $\beta = 0.0031, \gamma = 0.2, \tau = 0.1$, and starting network parameters are $N = 10,000, \langle k \rangle(0) = 64, \langle k^2 - k \rangle = 5120, \phi(0) = 0.2$. 

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Fig. 12. Relative change in final size (RCFS) for the prevalence-dependent intervention. Each plot represents a choice of $p$ and $q$, with $LI$ and $LR$ on the axes, ranging from 2 for 180. Initial conditions for the state variables are $S(0) = 9998$, $|E(0)| = |I(0)| = 1$, $|SS| = 639784$, $|SE| = 60$, $|SF| = 48$, $|EE| = |EI| = |II| = 0$. Epidemiological parameters are $\beta = 0.0031$, $\eta = 0.2$, $\gamma = 0.1$, and starting network parameters are $N = 10000$, $|k(0)| = 64$, $|k^2 - k| = 5120$, $\phi(0) = 0.2$.

Fig. 13. Progression of the infection curve $|I(t)|$ as $L_I$ increases, showing the shrinking of the final spike and the penultimate spike dropping below the threshold $qN$. Parameters are $q = 0.01$, $p = 0.25$, $L_I = 90$ and $L_I = 70$ (a), 78 (b), 92 (c). Solid orange curves are $|I(t)|$ under the intervention, while dashed orange curves are $|I(t)|$ without any intervention. Dashed gray lines denote times when $|I| = qN$. Initial conditions for the state variables are $S(0) = 9998$, $|E(0)| = |I(0)| = 1$, $|SS| = 639784$, $|SE| = 60$, $|SF| = 48$, $|EE| = |EI| = |II| = 0$. Epidemiological parameters are $\beta = 0.0031$, $\eta = 0.2$, $\gamma = 0.1$, and starting network parameters are $N = 10000$, $|k(0)| = 64$, $|k^2 - k| = 5120$, $\phi(0) = 0.2$.  

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AIAT increases rapidly as the epidemic changes from three to two spikes.

While Figs. 12 and 14 show the overall behavior of the prevalence-dependent intervention, by considering fixed values of $L_I$ and $L_R$ and allowing $p$ and $q$ to vary, we get a more pointed perspective on the effectiveness of this type of intervention. Fig. 15 shows increasingly gradual interventions from left to right with plots of the RCFS and AIAT as $p$ and $q$ vary on the axes. Notably, regardless of $L_I$, low values of $p$ and $q$ are able to produce interventions that both greatly decrease the final size of the epidemic, and the average infections above threshold. An important factor in this decrease in the final size is the time at which the measurements of the RCFS and AIAT are taken. In Fig. 15, the epidemic is allowed to run for three years before both are measured. Measuring earlier results in a larger portion of $p,q$ space that greatly decreases the RCFS, while measuring later results in a smaller portion. This may have important control impacts. For instance, if a vaccine is expected to be designed, produced, and distributed, the estimated time frame for that to occur can influence which levels of $p$ and $q$ can produce highly effective interventions. Regardless, the region of highly effective interventions appears mostly the same for the different values of $L_I$. This suggests that for sufficiently low thresholds ($q$) and sufficiently severe decreases in contacts ($p$), the length over which the decrease in contacts occurs ($L_I$) does not play an important role in the effectiveness of interventions. However, as $q$ or $p$ increases, $L_I$ has a more pronounced impact. In particular, for low values of $p$ and large values of $q$, a longer, more gradual intervention can lead to more average infections above threshold. Moreover, a stark change in both effectiveness metrics occurs for large values of $p$, (around $p = 0.5$ for $L_I = 15$ and $L_I = 30$). This suggests that if an intervention doesn’t reduce average contacts sufficiently, a highly effective intervention isn’t possible, regardless of the other parameter values.

4. Discussion

In this paper, we have developed a new SEIR model on a network with random link activation/deletion dynamics. Using piecewise constant activation and deletion rate functions, we propose two simple mechanisms for social distancing interventions. The simple intervention models a single intervention event, where contacts are decreased over a period of time, stay constant, and then return to pre-intervention levels. The prevalence-dependent intervention expands the simple case to more complex scenarios, where interventions can be reintroduced in the face of rising prevalence. Using the unipartite projection of a bipartite network, and epidemiological parameters representative of COVID-19, we examine

![Fig. 14](image.png)

Fig. 14. Average infections above threshold (AIAT) for the prevalence-dependent intervention. Each plot represents a choice of $p$ and $q$, with $L_I$ and $L_R$ on the axes, ranging from 2 for 180. Initial conditions for the state variables are $S(0) = 9998$, $E(0) = 1$, $I(0) = 1$, $SS = 639784$, $SE = 60$, $SI = 48$, $EE = 0$, $EI = 0$. Epidemiological parameters are $\beta = 0.0031$, $\eta = 0.2$, $\gamma = 0.1$, and starting network parameters are $N = 10,000$, $\langle k \rangle = 64$, $\langle k^2 \rangle - k = 5120$, $\psi(0) = 0.2$. 

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the effectiveness of a wide range of potential social distancing poli-

cies on relatively large heterogeneous, clustered networks.

Both intervention schemes are shown to capture a wide variety of behaviors in the prevalence "curve," which has received considerable attention in both academic studies and public health mes-
saging. The simple intervention manifests curves with one or two spikes, while the curves for prevalence-dependent intervention can have many more. Moreover, the behavior of the prevalence curve is consistent across a number of parameters and can be described qualitatively with success. This is despite the simplicity of social distancing mechanism introduced by the piecewise constant activation and deletion rates $\alpha(t)$ and $\omega(t)$, which take on values $\alpha'$ or $\omega'$ respectively, or zero. We have not considered the cases where the values of $\alpha'$ and $\omega'$ may change over time, or where $\alpha(t)$ and $\omega(t)$ are not piecewise constant. As such, our model has natu-
al extensions that may capture an even richer variety of qualitative behaviors.

Furthermore, the mechanisms proposed in this paper offer insights into what makes for a successful intervention. We have used two metrics as simplified public health goals to evaluate the effectiveness of interventions: the relative change in final size (RCFS) and the average infections above threshold (AIAT). For the more realistic prevalence-dependent intervention scheme, we find that the most effective interventions come when the threshold number of infections is low and the intervention severely decreases average contacts. When these conditions are met, the relative change in the final size is greatly decreased and the length over which the intervention is implemented has little impact on the effectiveness. However, even small increases in the threshold value can greatly impact the effectiveness of interventions over a fixed period of time. As well, if interventions do not sufficiently reduce contacts (around fifty percent), they are rendered signifi-
cantly less effective by both measures.

While this is a first foray into the use of adaptive networks to model social distancing for an SEIR disease, we acknowledge some limitations of our model. First, there is a trade-off between com-
plexity of the disease natural history model and the number of equations of the pairwise model; age-structured models or other more complex compartmental models are popular for COVID-19, but added compartments require tracking an increasing number of edge types. However, even simple extensions (such as the inclu-
sion of an asymptomatic infectious state) present interesting opportunities. Second, while the random link activation/deletion process is simple to implement, it has some unrealistic features. In particular, in the $t \to \infty$ limit, one can show from the degree dis-
tribution generating function that the resulting network approaches an Erdős-Rényi random graph, with vanishing cluster-
ing and an approximately Poisson degree distribution. One mani-
ifestation of this property is a rapidly declining clustering coefficient over time. While the piecewise constant activation and deletion rates mitigate this to an extent, the network resulting from these social distancing policies is fundamentally different than the initial network state. To overcome this limitation, future investigations might involve new processes for network dynamics, such as activation/deletion on a fixed network or network dynam-
ics on an underlying bipartite mixing network. Finally, the choice of triple closure in this paper incorporates network structure only through the parameters $\langle k \rangle$, $\langle k^2 - k \rangle$, and $\phi$ for the entire network. The real-world effect of social distancing measures may affect high-degree nodes differently than low-degree nodes, and thus modeling a social distancing scheme for the entire degree distribu-
tion may be more realistic and flexible.
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Appendix A. Supplementary data
Supplementary data associated with this article can be found, in the online version, at https://doi.org/10.1016/j.jtbi.2022.111151.

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