Contact tracing and disease control

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Contact tracing, followed by treatment or isolation, is a key control measure in the battle against infectious diseases. It is an extreme form of locally targeted control, and as such has the potential to be highly efficient when dealing with low numbers of cases. For this reason it is frequently used to combat sexually transmitted diseases and new invading pathogens. Accurate modelling of contact tracing requires explicit information about the disease-transmission pathways from each individual, and hence the network of contacts. Here, pairwise-approximation methods and full stochastic simulations are used to investigate the utility of contact tracing. A simple relationship is found between the efficiency of contact tracing necessary for eradication and the basic reproductive ratio of the disease. This holds for a wide variety of realistic situations including heterogeneous networks containing core-groups or super-spreaders, and asymptomatic individuals. Clustering (transitivity) within the transmission network is found to destroy the relationship, requiring lower efficiency than predicted.

Keywords: tracing efficiency; transmission networks; pairwise approximations; stochastic simulation

1. INTRODUCTION

A range of control measures, from prophylactic vaccination (Fine & Clarkson 1983; McLean & Anderson 1988; Anderson & May 1992) to pre-emptive culling (Woolhouse & Donaldson 2001; Keeling et al. 2001), have been successfully used to combat the spread of disease or even eradicate infection entirely (Nokes & Swinton 1997; Bazin 2000). It is now well accepted that targeting such control measures, rather than applying them randomly, leads to an often dramatic increase in efficiency (Hethcote & Yorke 1984; Kault 1995). Contact tracing is an extreme form of targeted control, where the potential next-generation cases are the primary focus. Contact tracing has proved to be a highly successful strategy when the number of infectious cases is low. As such, contact tracing is used in the control of sexually transmitted diseases (STDs; Clarke 1998; FitzGerald et al. 1998; Macke & Maher 1999); it is the recommended policy for tackling outbreaks of new or re-emerging infections such as the recent severe acute respiratory syndrome (SARS) epidemic (Donnelly et al. 2003); and it is the standard tool for eliminating minor outbreaks in the latter stages of disease eradication. Spatially explicit contact tracing, in the form of either ring-vaccination or ring-culling, is also commonly used for a variety of livestock infections (Mueller et al. 2000b; Woolhouse & Donaldson 2001; Haydon et al. 2003). With the recent increase in STD prevalence, the perceived risk from bio-terrorism and the aim of global polio eradication, the public-health importance of an understanding of contact tracing has never been greater.

Contact tracing is fundamentally linked to the individual-level spread of infection and, in particular, the network of potential transmission routes. Networks are a particularly important concept in human disease modelling, where social interactions can take place over a wide range of distances and where geographical distance alone is an inadequate measure of infection risk. For this reason we concentrate on human diseases in this paper. While some models have been developed to consider the role of contact tracing in randomly interacting populations (Mueller et al. 2000a; Hyman et al. 2003), only network-based models that consider transmission pathways (Halloran et al. 2002; Huerta & Tsimring 2002) and the associated pairwise equations can provide an accurate mechanistic understanding of the structured nature of human interactions. Here, we use detailed pairwise equations (Eames & Keeling 2002) to gain an analytical insight into contact tracing both for STDs, such as gonorrhoea and chlamydia (which are described by a susceptible–infected–susceptible (SIS) framework), and for airborne infections, such as SARS, smallpox, pox and measles (which follow the susceptible–infected–recovered/dead paradigm). Finally, stochastic individual-based network simulations are used to validate the pairwise predictions and allow us to ascertain the effect of locally clustered networks on contact tracing.

Pairwise correlation models offer a robust mechanism to capture spatial effects by formulating equations for the number of connected pairs rather than just the number of individuals, and hence link individual-level behaviour to population-level dynamics. They have been shown to be in close agreement with full stochastic network simulations (Eames & Keeling 2002), and have been used to study a range of ecological and epidemiological problems (Keeling et al. 1997; Van Baalen & Rand 1998; Keeling 1999; Ferguson et al. 2001), in particular STDs, where pairs of connected individuals (sexual partnerships) are naturally specified (Bauers & Rand 2000; Ferguson & Garnett 2000; Eames & Keeling 2002). The pairwise approach has several key advantages over more complex simulations: most importantly, it provides an analytical framework, can be readily parameterized with available data (Eames & Keeling 2002) and generates results based on simple
properties of the transmission network. These advantages are used here to develop robust and generic results.

2. CONTACT-TRACING MODEL

Many STDs broadly conform to the SIS paradigm (Anderson & May 1992) whereby individuals enter sexually active life as susceptible (S), and infected individuals (I) return to the susceptible class following treatment. Treatment comes from two distinct sources, either the patient seeking medical assistance once symptoms develop, or the tracking of infection within the population by following potential transmission routes (sexual partnerships). Such contact tracing is an important public-health tool (Clarke 1998; FitzGerald et al. 1998; Macke & Maher 1999), especially when the disease may be asymptomatic (but still infectious), as it provides the only means by which such individuals can be easily identified. Contact tracing is incorporated into the model design using an SITS framework, where the treatment class (T) is short-lived and rapidly ‘transmitted’, mimicking the relatively quick speed at which tracing spreads across the network. From an ecological perspective we can view contact tracing as hyperparasitism (Mills & Gutierrez 1996; Sullivan & Voilkl 1999), spreading through the same network as the disease, but affecting only those individuals that are already infected. Tracing can also be considered when individuals acquire immunity from further infection; this adds a recovered (R) state, giving rise to an SITR model. Throughout, it is assumed that while in the treatment class an individual can no longer transmit infection; this may be the result either of treatment or of isolation, whether behavioural (such as avoiding risky sexual practices) or physical (e.g. isolation wards).

We denote by \([S], [I]\) and \([T]\) the numbers of susceptible and infected individuals, and the number in the treatment class, respectively. The population dynamics of each of these classes is determined by the partners to whom they are connected. Hence, to formulate the equations we let \([AB]\) refer to the number of partnerships within the network involving an individual of status A and another of status B (Keeling et al. 1997). The differential equations that govern the numbers in the individual classes are:

\[
\begin{align*}
[S] &= -\tau[S]I + a[T] \\
[I] &= \tau[S]I - g[I] - c(IT) \\
[T] &= g[I] + c(IT) - a[T],
\end{align*}
\]

where recovery occurs only after seeking medical advice, which initiates the contact-tracing process. Here, \(\tau\) is the rate of spread of infection across a partnership, \(g\) is the recovery rate in the absence of tracing, \(c\) is the rate of contact tracing and \(a\) is the rate at which individuals move out of the treatment class. As indicated above, further interpretations are possible: we might consider \(g\) to be an isolation rate, for instance.

To solve these equations the behaviour of pairs of connected individuals (partnerships) also needs to be modelled. These pair equations in principle need information about triples (and triple equations would need information about higher-order terms); however, we can close the system by approximating these triples in terms of pairs and singles (see electronic Appendix A available on The Royal Society’s Publications Web site). We define the infectivity of the disease, \(r\), as the rate of transmission across a contact multiplied by the infectious period, \(\tau = \tau g\), and similarly the tracing and treatment level, \(t\), is the rate that treatment ‘spreads’ multiplied by the treatment time, \(t = c/a\). The probability that infection passes across a contact is \(\tau/(g + \tau) = r/(1 + \tau)\), and the probability that treatment is passed on is \(t/(1 + \tau)\). We refer to this latter quantity, which represents the proportion of contacts that are traced, as the ‘tracing efficiency’.

3. CONTACT TRACING IN HOMOGENEOUS NETWORKS

When all infectious individuals display symptoms and all individuals have exactly \(n\) contacts to whom they can pass infection, analytical results allow an insight into eradication. By looking at the parameter values at which equilibrium prevalence and the initial growth of infection are zero, we can derive the level of tracing required to eliminate infection. In the SIS scenario, the critical level of tracing and treatment needed to eradicate infection is given by

\[
t_c = \frac{rn^2 + 1 - 2rn - 2n + \sqrt{n^4 + 4rn^2 + 1 - 4n^2(n - 1)}}{2(n - 1) - 1}
\]

and hence \(t_c/(1 + t_c)\) is the proportion of contacts that needs to be successfully traced and treated, i.e. the critical tracing efficiency. Figure 1a shows how the critical tracing efficiency dramatically increases with both the transmission probability of the disease and the number of contacts.

Applying the pairwise approximation to SIR-type infections results in a critical level of tracing with a much simpler form

\[
t_c = \frac{rn(n - 2) - n}{n - 1}
\]

Figure 1b shows the associated critical tracing efficiency for diseases of this sort. SIR infections always require less effort to control, although the difference between SIR and SIS models becomes negligible as the number of contacts, \(n\), becomes large—only where \(n\) is small are figures 1a and 1b appreciably different. In both models (most clearly from equation (3.2)), the critical level of contact tracing increases almost linearly with both the infectivity of the disease, \(r\), and the number of contacts, \(n\). We can relate this result to more standard epidemiological measures. The basic reproductive ratio, \(R_0\), is defined as the expected number of secondary cases produced by an infectious individual in a totally susceptible population, and is calculated from the early asymptotic growth rate of the epidemic (Dickmann et al. 1990). For an SIR disease on a network, pairwise models show that the basic reproductive ratio is given by \(R_0 = (n - 2) r\) (Keeling 1999). Hence, when the number of contacts, \(n\), is large, equation (3.2) implies that critical tracing efficiency,

\[
t_c = \frac{1}{R_0}
\]

although this is a slight underestimate (figure 1c). For example, if \(R_0 = 4\), around 75% of contacts must be traced to control infection. The same relationship also holds for
SIS-type infections (equation (3.1)), despite the more complicated forms of $t_c$ and $R_0$ (Eames & Keeling 2002). Surprisingly, this approximation is in direct agreement with the threshold level of random vaccination needed to eradicate infection (Anderson & May 1992) and agrees with that predicted using unstructured partnership models (Müller et al. 2000a). Thus, while the presence of strong local correlations, captured by the pairwise models, has a significant impact on the two emergent properties of $R_0$ and the critical tracing level, the relationship between these two properties remains largely unchanged.

It should be noted that we have implicitly assumed that all contacts are known and therefore theoretically traceable. However, if a proportion, $p$, of contacts are unknown (and therefore untraceable), then, to maintain the same average level of tracing, the tracing efficiency for the remaining known contacts must be increased to

$$\text{critical tracing efficiency } = \frac{1}{1 - p} \left(1 - \frac{1}{R_0}\right).$$

Thus, control of diseases with large reproductive ratios requires not only a high level of contact tracing but also comprehensive knowledge of the potential transmission routes.

**4. CONTROL IN SEXUAL MIXINGNETWORKS**

We demonstrate how the pairwise model may be extended by introducing two more realistic features of STD transmission. First, a heterogeneous network is used so that individuals have differing numbers of contacts, reflecting the important role of super-spreaders or core groups in the transmission and persistence of infection. This adaptation is appropriate for a wide range of infections. We use a network based on observed social networks (Klovdahl et al. 1977), but parameterizations from other networks produce comparable results. Second, a proportion of infected individuals are assumed to be asymptomatic: these individuals seldom seek treatment and consequently have a much longer infectious period (see electronic Appendix A). Figure 2a,b shows the equilibrium prevalence of infection as control measures are varied. The axes reflect the effort expended on each of the two major control mechanisms: identification of those individuals with symptoms ($x$-axis) and the contact-tracing efficiency ($y$-axis). When the proportion of new cases that are asymptomatic is larger (figure 2b) the importance of contact tracing increases. As asymptomatics are rarely detected or treated, it may be impossible to eradicate the infection without contact tracing even if symptomatic individuals are identified and treated immediately. Contact tracing is essential for eradication when each asymptomatic case is expected to give rise to at least one more asymptomatic case, i.e. when

$$(\text{basic reproductive ratio of asymptomatics}) \times (\text{probability of asymptomatic infection}) > 1.$$
Because asymptomatics may subsequently cause symptomatic cases, contact tracing can potentially identify and treat such individuals, dramatically reducing the force of the infection. The prevalence of infection in the population largely determines the optimal balance between the identification of symptomatic individuals and contact tracing. Contact tracing is favoured when infection is globally rare but locally common within connected areas of the network.

The extinction thresholds for the situations shown in figure 2a, b can be redrawn in terms of $R_0$ (figure 2c). Despite the additional levels of complexity that have been included in the model, the simple relationship critical tracing efficiency $\approx 1 - 1/R_0$ still holds. This degree of robustness is remarkable and reflects a fundamental link between the spread of infection and the spread of control.

5. THE EFFECTS OF CLUSTERING IN TRANSMISSION NETWORKS

While pairwise models provide an excellent approximation to the network of sexual contacts involved in the spread of STDs (Eames & Keeling 2002), they are less accurate for airborne diseases where there is a high level of clustering (transitivity) within the network, so that connected individuals are likely to share common contacts. We therefore investigate the role of contact tracing and treatment in full stochastic simulations of disease transmission on a variety of computer-generated networks (Read & Keeling 2003; figure 3). Not only does this provide much more flexibility in the network structure, but it allows us to address many more applied issues. For the simplest situation, where every individual has exactly the same number of neighbours, figure 3a (solid line) shows the tracing efficiency needed to eradicate infection. When the network is highly clustered the necessary level of tracing decreases; this is to be expected as $R_0$ is generally reduced in clustered networks owing to the self-limitation of the epidemic as the local density of susceptibles is depleted. When the number of contacts is heterogeneous and the clustering low (dashed line), the associated value of $R_0$ is increased, as predicted from both standard theory (Anderson & May 1992) and pairwise approximations (Eames & Keeling 2002), and this necessitates an increase in the efficiency of contact tracing.

Once again, these results can be redrawn in terms of $R_0$ (figure 3b), although here there is a clear deviation from the standard pattern. Where there is no clustering within the network, stochastic simulations confirm the analytically derived threshold of $1 - 1/R_0$. However, clustering...
acts very differently on the values of $R_0$ and the critical tracing efficiency. In general, $R_0$ decreases with clustering owing to increased self-limitation, although in the heterogeneous network (dashed line) small amounts of clustering create interconnected core-groups, which promote spread, and therefore $R_0$ initially increases. While the critical tracing efficiency responds to changes in $R_0$, clustering causes an additional reduction as individuals can be traced via a variety of routes owing to multiple partners being infected. Thus, contact tracing performs far better than predicted in highly clustered environments, and may be more effective against the clustered spread of airborne diseases than expected from work with STDs.

6. DISCUSSION

The analytical and simulation results presented have obvious public-health implications. For STDs, because most of the population have few sexual partners (Johnson et al. 1994), tracing is a very efficient means of identifying infected individuals, and hence reducing the number of cases or even eradicating the disease. Contact tracing provides a method of targeted control whereby intervention is automatically focused on the subset of the population most likely to be at risk of infection, i.e. partners of infected individuals. The benefits of contact tracing are even more pronounced when asymptomatic infections can linger unchecked in the population. For airborne infections, where the average number of neighbours and the basic reproductive ratio are larger, contact tracing has to be far more efficient and generally far more rapid.

Robust approximation models and stochastic simulations have shown that in the absence of clustering (when contacts occur independently) the necessary tracing efficiency can be estimated from the basic reproductive ratio (see equation (3.3)). This simple relationship is robust to a number of modifications that increase the realism of the epidemic models. Thus, while spatial correlations, network heterogeneity and asymptomatic infections can dramatically modify the epidemic dynamics (Keeling et al. 1997; Keeling 1999; Eames & Keeling 2002), the relationship between the emergent properties of $R_0$ and the critical tracing efficiency is conserved. This basic relationship can therefore be expected to hold for a wide range of STDs and real-world transmission scenarios. However, the more clustered (transitive) nature of airborne or social transmission networks breaks this relationship—although it still provides a useful upper bound. For the airborne spread of re-emerging diseases, where contact tracing is a recommended control strategy, an informed tracing policy requires an estimation of the degree of clustering within the transmission network. So far this element of transmission networks has not been comprehensively investigated, although diary-based studies of individual contacts may provide results (Edmunds et al. 1997).

The critical tracing efficiency derived for homogeneous networks shows that the level of contact tracing required depends on the number of contacts. This suggests that, in a realistic heterogeneous network, prioritizing tracing from those individuals with many contacts might further reduce the healthcare burden. Such targeted tracing may be a

![Diagram](image-url)

Figure 3. Average results from 500 stochastic simulations on computer-generated networks, focusing on the effects of clustering. Ten thousand individuals are positioned at random on a two-dimensional landscape, and connections are formed with a probability given by a Gaussian kernel. The clustering (the ratio of triangles to all triples in the network; Keeling 1999) of the network is determined by the kernel, with highly clustered configurations corresponding to a narrow kernel (Read & Keeling 2003). The infectivity is set at $r = 1$, so the probability of infection being transmitted across a contact is 0.5. (a) The critical tracing efficiency necessary for control is seen to decrease as the clustering increases. Results for two distinct types of network are shown: a homogeneous network in which everyone has exactly five contacts (solid line), and a heterogeneous network in which the number of contacts per individual varies but the average number of contacts remains five (dashed line). Also shown are examples of a random and a clustered network: for the random network the connections from two individuals are shown, whereas for the clustered network it is feasible to display all the connections. (b) The results from graph (a) shown with respect to the basic reproductive ratio, $R_0$. Clustering causes a substantial deviation away from the simple theoretical ideal $(1 - 1/R_0^2, \text{grey line})$, such that lower tracing efficiency is sufficient. When clustering is equal to zero, the theoretical threshold applies, but as clustering increases, the threshold moves away from the expected curve. For the heterogeneous network, small amounts of clustering actually increase $R_0$ owing to aggregation of high-risk individuals with many partners.
valuable tool in future control strategies, and merits deeper investigation with the techniques discussed here. There are several distinct scenarios under which contact tracing may not be an effective control strategy. In high-risk groups, with many possible transmission routes and a high incidence of infection, random screening may be a more efficient tool. Also, for airborne infections there may be a significant fraction of contacts that are untraceable; in such cases it may be impossible to achieve the necessary threshold, and additional control measures would need to be applied. If the tracing process is significantly slower than the infection process then, no matter how large a proportion of contacts is eventually traced, it will be impossible to keep pace with the epidemic; there is a trade-off between tracing speed and tracing efficiency. Finally, we note that, once above the critical level of contact tracing, the number of cases decreases through time, as the effective reproductive ratio, $R$, is less than 1. However, when this rate of decline is not sufficiently rapid ($R$ is close to 1) the epidemic may be long-lived and many subsequent cases can arise. Hence, the theoretical threshold, which reduces the reproductive ratio to below 1, acts as a lower bound to the tracing efficiency that should be sought. In practice, much higher tracing efficiency may be needed to reduce the epidemic size quickly to a minimal level. Highly efficient contact tracing may place fewer demands on medical resources as a rapidly curtailed outbreak will involve fewer individuals needing to be traced and treated.

The models presented here represent highly simplified versions of real infection processes, but serve to indicate how contact tracing may be effective. More complex disease dynamics may alter the balance between different intervention methods. For example, diseases that are infectious before they become symptomatic will be harder to control through contact tracing: by the time index cases are identified, the epidemic will have spread some distance through the network. However, the ability of contact tracing to identify asymptomatic individuals would prove beneficial in such situations. The efficacy of contact tracing, therefore, is dependent not only on constraints or resources, but also on the disease itself.

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REFERENCES


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