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A class of pairwise models for epidemic dynamics on weighted networks

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Abstract In this paper, we study the SIS (susceptible-infected-susceptible) and SIR (susceptible-infected-removed) epidemic models on undirected, weighted networks by deriving pairwise-type approximate models coupled with individual-based network simulation. Two different types of theoretical/synthetic weighted network models are considered. Both models start from non-weighted networks with fixed topology followed by the allocation of link weights in either (i) random or (ii) fixed/deterministic way. The pairwise models are formulated for a general discrete distribution of weights, and these models are then used in conjunction with network simulation to evaluate the impact of different weight distributions on epidemic threshold and dynamics in general. For the SIR dynamics, the basic reproductive ratio $R_0$ is computed, and we show that (i) for both network models $R_0$ is maximised if all weights are equal, and (ii) when the two models are “equally-matched”, the networks with a random weight distribution give rise to a higher $R_0$ value. The models are also used to explore the agreement between the pairwise and simulation models for different parameter combinations.

1 Introduction

Conventional models of epidemic spread consider a host population of identical individuals, each interacting in the same way with each of the others (see [1,17,33] and references therein). At the same time, in order to develop more realistic mathematical models for the spread of infectious diseases, it is important to obtain the best possible representation of the corresponding transmission mechanism. To achieve this, more recent models have included some of the many complexities that
have been observed in mixing patterns. One such approach consists in splitting the population into a set of different subgroups, each with different social behaviours. Even more detail is included within network approaches which allow to include differences between individuals, not just between sub-populations. In such models, each individual is represented as a node, and interactions that could permit the transmission of infection appear as edges linking nodes. The last decade has seen a substantial increase in the research of how infectious diseases spread over large networks of connected nodes [34,39], where networks themselves can represent either small social contact networks [38] or larger scale travel networks [15,19], including global aviation networks [41,42]. Importantly, the characteristics of the network, such as the average degree and the node degree distribution have a profound effect on the dynamics of the infectious disease spread, and hence significant efforts are made to capture properties of realistic contact networks.

One of the simplifying assumptions often put into network models is that all links are equally likely to transmit infection [10,24,34,45]. However, a more detailed consideration leads to an observation that this is often not the case, as some links are likely to be far more capable of transmitting infection than others due to closer contacts (e.g. within households [7]) or long-duration interactions [23,44,45,46]. To account for this heterogeneity in properties of social interactions, network models can be adapted, thus resulting in weighted contact networks, where connections between different nodes have different weights. These weights may be associated with the duration, proximity, or social setting of the interaction, and the key point is that they are expected to be correlated with the risk of disease transmission. The precise relationship between the properties of an interaction and its riskiness is hugely complex; here, we will consider a “weight” that is exactly proportional to the transmission rate along a link. Although consideration of weighted networks may seem as an additional complication for the analysis of epidemic dynamics, in fact it provides a much more realistic representation of actual contact networks.

Substantial amount of work has been done on the analysis of weighted networks [3,4,5,37] and scale-free networks with different types of weight distribution [48]. In epidemiological context, Britton et al. [11] have derived an expression for the basic reproductive ratio in weighted networks with generic distributions of node degree and link weight, and Deijfen [16] has performed a similar analysis to study vaccination in such networks. In terms of practical epidemiological applications, weighted networks have already been effectively used to study control of global pandemics [13,14,22] and the spread of animal disease due to cattle movement between farms [26]. Eames et al. [22] have considered an SIR model on an undirected weighted network, where rather than using some theoretical formalism to generate an idealized network, the authors have relied on social mixing data obtained from questionnaires completed by members of a peer group [44] to construct a realistic weighted network. Having analysed the dynamics of epidemic spread in such a network, they showed how information about node-specific
infection risk can be used to develop targeted preventative vaccina-
79 tion strategies. Yan et al. [49] analysed a model on weighted scale-free
networks and found that heterogeneity in weight distribution leads
79 to a slowdown in the spread of epidemics. Furthermore, they have
shown that for a given network topology and mean infectivity, epidemics
spread fastest in unweighted networks. Yang et al. [51] have shown that
disease prevalence can be maximized when the edge weights are chosen
to be inversely proportional to the degrees of receiving nodes but, in this
case, the transmissibility was not directly proportional to the weights
and weights were also asymmetric. Yang & Zhou [50] have considered
SIS epidemics on homogeneous networks with uniform or power-law
edge weight distribution and shown how to derive a certain type of
mean-field description for such models.

In this paper, we consider the dynamics of an infectious disease
spreading on weighted networks with different weight distributions.
Since we are primarily concerned with the effects of weight distribution
on the disease dynamics, the connection matrix will be assumed to be
symmetric, representing the situation when the weights can only be
different for different network edges, but for a given edge the weight is
the same irrespective of the direction of infection. From epidemiological
perspective, we consider both the case when the disease confers per-
manent immunity (represented by an SIR model), and the case when
the immunity is short-lived, and upon recovery the individuals return
to the class of susceptibles (SIS model). For both of these cases we
derive the corresponding ODE-based pairwise models and their closure
approximations. Numerical simulation of both the epidemic spread on
the network and the pairwise approximations are performed.

The outline of this paper is as follows. In the next section, the con-
struction of specific weighted networks to be used for the analysis of epi-
demic dynamics is discussed. This is complemented by the derivation of
corresponding pairwise models and their closure approximations. Sec-
tion 3 contains the derivation of the basic reproductive ratio $R_0$ for the
SIR model and for different weight distributions as well as numerical
simulation of both network models and their pairwise ODE counter-
parts. The paper concludes in Section 4 with discussion of results and
possible further extensions of this work.

2 Model derivation

2.1 Network construction and simulation

There are two conceptually different approaches to constructing weighted networks
for modelling infectious disease spread. In the first approach, there is a seed or a
primitive motif, and the network is then grown or evolved from this initial seed
according to some specific rules. In this method, the topology of the network is
coevolving with the distribution of weights on the edges [4,5,6,37,51]. Another
approach is to consider a weighted network as a superposition of an unweighted
network with a distribution of weights across edges which could be independent
of the original network or it may be correlated with node metrics, such as their degree, [11,16,25,?]. In this paper we use the second approach in order to investigate the particular role played by the distribution of weights across edges, rather than network topology, in the dynamics of epidemic spread. Besides computational efficiency, this will allow us to make some analytical headway in deriving and analysing low-dimensional pairwise models which are likely to perform better when weights are attached according to the scenarios described above.

Here we consider two different methods of assigning weights to network links: a network in which weights are assigned to links at random, and a network in which each node has the same distribution of weighted links connected to it. In reality, there is likely to be a great deal more structure to interaction weights, but in the absence of precise data and also for the purposes of developing models that allow one to explore a number of different assumptions, we make these simplifying approximations.

2.1.1 Random weight distribution

First we consider a simple model of an undirected weighted network with \( N \) nodes where the weights of the links can take values \( w_i \) with probability \( p_i \), where \( i = 1, 2, \ldots, M \). The underlying degree distribution of the corresponding unweighted network can be chosen to be of the more basic forms, e.g. homogeneous random or Erdös-Rényi-type random networks.

The generation of such networks is straightforward, and weights can be assigned during link creation in the unweighted network. For example, upon using the configuration model for generating unweighted networks, each new link will have a weight assigned to it based on the chosen weight distribution. This means that in a homogeneous random network with each node having \( k \) links, the distribution of link weights of different type will be multinomial, and it is given by

\[
P(n_{w_1}, n_{w_2}, \ldots, n_{w_M}) = \frac{k!}{n_{w_1}! n_{w_2}! \cdots n_{w_M}!} p_1^{n_{w_1}} p_2^{n_{w_2}} \cdots p_M^{n_{w_M}},
\]

where, \( n_{w_1} + n_{w_2} + \cdots + n_{w_M} = k \) and \( P(n_{w_1}, n_{w_2}, \ldots, n_{w_M}) \) stands for the probability of a node having \( n_{w_1}, n_{w_2}, \ldots, n_{w_M} \) links with weights \( w_1, w_2, \ldots, w_M \), respectively. While the above expression is applicable in the most general set-up, it is worth considering the case of weights of only two types, where the distribution of link weights for a homogeneous random network becomes binomial

\[
P(n_{w_1}, n_{w_2} = k - n_{w_1}) = \binom{k}{n_{w_1}} p_1^{n_{w_1}} (1 - p_1)^{k-n_{w_1}},
\]

where, \( p_1 + p_2 = 1 \) and \( n_{w_1} + n_{w_2} = k \). The average link weight in the model above can be easily found as

\[
w_{av}^{random} = \sum_{i=1}^{M} p_i w_i,
\]

which for the case of weights of two types \( w_1 \) and \( w_2 \) reduces to

\[
w_{av}^{(2r)} = p_1 w_1 + p_2 w_2 = p_1 w_1 + (1 - p_1) w_2.
\]
2.1.2 Fixed deterministic weight distribution

As a second example we consider a network, in which each node has \( k_i \) links with weight \( w_i \) (\( i = 1, 2, \ldots, M \)), where \( k_1 + k_2 + \cdots + k_M = k \). The different weights here could be interpreted as being associated with different types of social interaction: e.g. home, workplace, and leisure contacts, or physical and non-physical interactions. In this model all individuals are identical in terms of their connections, not only having the same number of links (as in the model above) but also having the same set of weights. The average weight in such a model is given by

\[
 w_{av}^{fixed} = \frac{1}{M} \sum_{i=1}^{M} p_i w_i, \quad p_i = \frac{k_i}{k},
\]

where \( p_i \) is the fraction of links of type \( i \) for each node. In the case of links of two types with weights \( w_1 \) and \( w_2 \), the average weight becomes

\[
 w_{av}^{(2f)} = p_1 w_1 + p_2 w_2 = \frac{k_1}{k} w_1 + \frac{k_2}{k} w_2 = \frac{k_1}{k} w_1 + \frac{k - k_1}{k} w_2.
\]

2.1.3 Simulation of epidemic dynamics

In this study, the simple SIS and SIR epidemic models are considered. The epidemic dynamics is specified in terms of infection and recovery events. The rate of transmission across an unweighted edge between an infected and susceptible individual is denoted by \( \tau \). This will then be adjusted by the weight of the link which is assumed to be directly proportional to the strength of the transmission along that link. Infected individuals recover independently of each other at rate \( \gamma \).

The simulation is implemented using the Gillespie algorithm [27] with inter-event times distributed exponentially with a rate given by the total rate of change in the network, with the single event to be implemented at each step being chosen at random and proportionally to its rate. All simulations start with most nodes being susceptible and with a few infected nodes chosen at random.

2.2 Pairwise equations and closure relations

In this section we extend the classic pairwise model for unweighted networks [32, 43] to the case of weighted graphs with \( M \) different link-weight types. Pairwise models successfully interpolate between classic compartmental ODE models and full individual-based network simulation with the added advantage of high transparency and a good degree of analytical tractability. These qualities make them an ideal tool for studying dynamical processes on networks [20, 28, 30, 32], and they can be used on their own and/or in parallel with simulation. The original versions of the pairwise models have been successfully extended to networks with heterogeneous degree distribution [21], asymmetric networks [47] and situations where transmission happens across different/combined routes [20, 28] as well as when taking into consideration network motifs of higher order than pairs and triangles [29]. The extension that we propose is based on the previously established precise counting procedure at the level of individuals, pairs and triples, as well as on a careful and systematic account of all possible transitions needed to derive the full
set of evolution equations for singles and pairs. These obviously involve the precise
dependency of lower order moments on higher order ones, e.g. the rate of change
of the expected number of susceptible nodes is proportional to the expected num-
ber of links between a susceptible and infected node. We extend the previously
well-established notation [32] to account for the added level of complexity due to
different link weights. In line with this, the number of singles remains unchanged,
with \( [A] \) denoting the number of nodes across the whole network in state \( A \). Pairs
of type \( A - B, [AB] \), are now broken down depending on link weights, i.e. \( [AB]_i \),
represents the number of links of type \( A - B \) with the link having weight \( w_i \),
where as before \( i = 1, 2, \ldots, M \) and \( A, B \in \{S, I, R\} \) if an SIR dynamics is used.
As before, links are doubly counted (e.g. in both directions) and thus the follow-
ing relations hold: \( [AB]_m = [BA]_m \) and \( [AA]_m \) is equal to twice the number of
uniquely counted links of weight \( w_m \) with nodes at both ends in state \( A \). From this
extension it follows that \( \sum_{i=1}^{M} [AB]_i = [AB] \). The same convention holds at the
level of triples where \( [ABC]_{mn} \) stands for the expected number of triples where a
node in state \( B \) connects a node in state \( A \) and \( C \) via links of weight \( w_m \) and \( w_n \),
respectively. The weight of the link impacts on the rate of transmission across that
link, and this is achieved by using a link-specific transmission rate equal to \( \tau w_i \),
where \( i = 1, 2, \ldots, M \). In line with the above, we construct two pairwise models,
one for SIS and one for SIR dynamics.

The pairwise model for the SIS dynamics can be written in the form:

\[
\begin{align*}
\dot{S} &= \gamma[I] - \tau \sum_{n=1}^{M} w_n [SI]_n, \\
\dot{I} &= \tau \sum_{n=1}^{M} w_n [SI]_n - \gamma[I], \\
\dot{SI}_m &= \gamma([I]_m - [SI]_m) + \tau \sum_{n=1}^{M} w_n ([SSI]_{mn} - [ISI]_{mn}) - \tau w_m [SI]_m, \quad (3) \\
\dot{IS}_m &= -2\gamma[I]_m + 2\tau \sum_{n=1}^{M} w_n [ISI]_{nm} + 2\tau w_m [SI]_m, \\
\dot{SS}_m &= 2\gamma[SI]_m - 2\tau \sum_{n=1}^{M} w_n [SSI]_{mn},
\end{align*}
\]

where \( m = 1, 2, 3, \ldots, M \) and \( [AB]_m \) denotes the expected number of links with
weight \( w_m \) connecting two nodes of type \( A \) and \( B \), respectively \( (A, B \in \{S, I\}) \).

In the case when upon infection individuals recover at rate \( \gamma \) and once recovered
they maintain a life-long immunity, we have the following system of equations
describing the dynamics of a pairwise SIR model:
\[
\dot{S} = -\tau \sum_{n=1}^{M} w_n [SI]_n,
\]
\[
\dot{I} = \tau \sum_{n=1}^{M} w_n [SI]_n - \gamma [I],
\]
\[
\dot{R} = \gamma [I],
\]
\[
\dot{SS}_m = -2\tau \sum_{n=1}^{M} w_n [SSI]_{mn},
\]
\[
\dot{SI}_m = \tau \sum_{n=1}^{M} w_n ([SSI]_{mn} - [ISI]_{nm}) - \tau w_m [SI]_m - \gamma [SI]_m,
\]
\[
\dot{SR}_m = -\tau \sum_{n=1}^{M} w_n [ISR]_{nm} + \gamma [SI]_m,
\]
\[
\dot{II}_m = 2\tau \sum_{n=1}^{M} w_n [ISI]_{mn} + 2\tau w_m [SI]_m - 2\gamma [II]_m,
\]
\[
\dot{IR}_m = \tau \sum_{n=1}^{M} w_n [ISR]_{nm} + \gamma ([II]_m - [IR]_m),
\]
\[
\dot{RR}_m = \gamma [IR]_m,
\]

where again \(m = 1, 2, 3, \ldots, M\) with the same notation as above. As a check and a reference back to previous pairwise models, in Appendix A we show how systems (3) and (4) reduce to the standard unweighted pairwise SIS and SIR model \([32]\) when all weights are equal to each other, \(w_1 = w_2 = \cdots = w_M = W\).

The above systems of equations (3) and (4) are not closed, as equations for the pairs require knowledge of triples, and thus, equations for triples are needed. This dependency on higher-order moments can be curtailed by closing the equations via approximating triples in terms of singles and pairs \([32]\). For both systems, the agreement with simulation will heavily depend on the precise distribution of weights across the links, the network topology, and the type of closures that will be used to capture essential features of network structure and the weight distribution.

A natural extension of the classic closure is given by

\[
[ABC]_{mn} = k \frac{1}{k} \frac{[AB]_m [BC]_n}{[B]},
\]

where \(k\) is the number of links per node for a homogeneous network or the average nodal degree for networks with other than homogenous degree distributions.

However, even for the simplest case of homogenous random networks with two weights (i.e. \(w_1\) and \(w_2\)), the average degree is split according to weight. Namely, the average number of links of weight \(w_1\) across the whole network is \(k_1 = p_1 k \leq k\), and similarly, the average number of links of weight \(w_2\) is \(k_2 = (1 - p_1) k \leq k\), where \(k = k_1 + k_2\). Attempting to better capture the additional network structure generated by the weights, the closure relation above can be recast to give the following,
potentially more accurate, closures

\[
\begin{align*}
[ABC]_{11} &= [AB]_1 (k_1 - 1) \frac{[BC]_1}{k_1 [B]} = \frac{k_1 - 1}{k_1} \frac{[AB]_1 [BC]_1}{[B]}, \\
[ABC]_{12} &= [AB]_1 k_2 \frac{[BC]_2}{k_2 [B]} = \frac{[AB]_1 [BC]_2}{[B]}, \\
[ABC]_{21} &= [AB]_2 k_1 \frac{[BC]_1}{k_1 [B]} = \frac{[AB]_2 [BC]_1}{[B]}, \\
[ABC]_{22} &= [AB]_2 (k_2 - 1) \frac{[BC]_2}{k_2 [B]} = \frac{k_2 - 1}{k_2} \frac{[AB]_2 [BC]_2}{[B]},
\end{align*}
\]

(6)

where, as in Eq. (5), the form of the closure can be derived by considering the central individual in the triple, \( B \). The first pair of the triple \([AB]_1\) effectively “uses up” one of \( B \)’s links of weight \( w_1 \). For triples of the form \([ABC]_{11}\), the presence of the pair \([AB]_1\) means that \( B \) has \((k_1 - 1)\) remaining links of weight \( w_1 \) that could potentially connect to \( C \). For triples of the form \([ABC]_{12}\), however, \( B \) has \( k_2 \) weight \( w_2 \) links that could potentially connect to \( C \). Furthermore, expressions such \( \frac{[BC]_i}{k_i [B]} \) simply denote the fraction of edges of weight \( w_i \) that start at a node \( B \) and connect this to \( C \). The specific choice of closure will depend on the structure of the network and, especially, how the weights are distributed. For example, for the case of the homogeneous random networks with links allocate randomly, both closures offer a viable alternative. For the case of a network where each node has a fixed pre-allocated number of links with different weights, e.g. \( k_1 \) and \( k_2 \) links with weights \( w_1 \) and \( w_2 \), respectively, the second closure \((6)\) offers the more natural/intuitive avenue towards closing the system and obtaining good agreement with network simulation.

3 Results

In this section we present analytical and numerical results for weighted networks and pairwise representations of SIS and SIR models in the case of two different link-weight types (i.e. \( w_1 \) and \( w_2 \)).

3.1 Threshold dynamics for the SIR model - the network perspective

The basic reproductive ratio, \( R_0 \) (the average number of secondary cases produced by a typical index case in an otherwise susceptible population), is one of the most fundamental quantities in epidemiology ([1,18]). Besides informing us on whether a particular disease will spread in a population, as well as quantifying the severity of an epidemic outbreak, it can be also used to calculate a number of other important quantities that have good intuitive interpretation. In what follows, we will compute \( R_0 \) and \( R_0 \)-like quantities and will discuss their relation to each other, and also issues around these being model-dependent. First, we compute \( R_0 \) from
an individual-based or network perspective by employing the next generation matrix approach as used in the context of models with multiple transmission routes such as household models [2].

Random weight distribution: First we derive an expression for $R_0$ when the underlying network is homogeneous, and the weights of the links are assigned at random according to a prescribed weight distribution. In the spirit of the proposed approach, the next generation matrix can be easily computed to yield

$$NGM = (a_{ij})_{i,j=1,2} = \begin{vmatrix} (k-1)p_1r_1 & (k-1)p_1r_1 \\ (k-1)p_2r_2 & (k-1)p_2r_2 \end{vmatrix},$$

where

$$r_1 = \frac{\tau w_1}{\tau w_1 + \gamma}, \quad r_2 = \frac{\tau w_2}{\tau w_2 + \gamma}$$

represent the probability of transmission from an infected to a susceptible across a link of weight $w_1$ and $w_2$, respectively. Here, the entry $a_{ij}$ stands for the average number of infections produced via links of type $i$ (i.e. with weight $w_i$) by a typical infectious node who itself has been infected across a link of type $j$ (i.e. with weight $w_j$). Using the fact that $p_2 = 1 - p_1$, the basic reproductive ratio can be found from the leading eigenvalue of the $NGM$ matrix as follows

$$R_0 = (k-1)(p_1r_1 + (1-p_1)r_2). \quad (7)$$

In fact, the expression for $R_0$ can be simply generalised to more than two weights to give $R_0 = (k-1)\sum_{i=1}^{M} p_ir_i$, where $w_{m}$ has frequency given by $p_{m}$ with the constraint that $\sum_{i=1}^{M} p_i = 1$. It is straightforward to show that upon assuming uniform weight distribution $w_i = W$ for $i = 1, 2, \ldots, M$, the basic reproduction number on a homogeneous graph reduces to $R_0 = (k-1)r$ as expected, and where, $r = \frac{\tau W}{\tau W + \gamma}$. 

Deterministic weight distribution: The case when the number of links with given weights for each node is fixed can be captured with the same approach, and the next generation matrix can be constructed as follows

$$NGM = \begin{vmatrix} (k_1-1)r_1 & k_1r_1 \\ k_2r_2 & (k_2-1)r_2 \end{vmatrix}.$$ 

As before, the leading eigenvalue of the $NGM$ matrix yields the basic reproductive ratio,

$$R_0^2 = \frac{(k_1-1)r_1 + (k_2-1)r_2 + \sqrt{[(k_1-1)r_1 - (k_2-1)r_2]^2 + 4k_1k_2r_1r_2}}{2} \quad (8)$$

It is worth noting that the calculations above are a direct result of a branching process approximation of the pure transmission process which differentiates between individuals depending on whether he/she was infected via a link with of weight $w_1$ or $w_2$, with obvious generalisation to more than two weights. This separation used in the branching process leads to the offspring or next generation matrix of the branching process [2]. Using the two expressions for the basic reproductive ratio, it is
possible to prove the following result.

**Theorem 1.** Given the setup for the fixed weight distribution and using $p_1 = k_1/k$, $p_2 = k_2/k$ and $k_1 + k_2 = k$, if $1 \leq k_1 \leq k - 1$ (which implies that $1 \leq k_2 \leq k - 1$), then $R_0^2 \leq R_0^1$.

The proof of this result is sketched out in Appendix B. This Theorem effectively states that provided each node has at least one link of type 1 and one link of type 2, then independently of disease parameters, it follows that the basic reproductive ratio as computed from (7) always exceeds or is equal to an equivalent $R_0$ computed from (8).

It is worth noting that both $R_0^1$ and $R_0^2$ values reduce to

$$R_0^1 = R_0^2 = R_0 = (k - 1)r = \frac{(k - 1)\tau W}{\tau W + \gamma},$$

(9)

if one assumes that weights are equal, i.e. $w_1 = w_2 = W$. As one would expect, the first good indicator of the impact of weights on the epidemic dynamics will be the average weight. Hence, it is worth considering the problem of maximising the values $R_0$ under assumption of a fixed average weight:

$$p_1w_1 + p_2w_2 = W.$$

(10)

Under this constraint the following statement holds.

**Theorem 2.** For weights constrained by $p_1w_1 + p_2w_2 = W$ (or $(k_1/k)w_1 + (k_2/k)w_2 = W$ for a fixed weights distribution), $R_0^1$ and $R_0^2$ attain their maxima when $w_1 = w_2 = W$, and the maximum values for both is $R_0 = (k - 1)r = \frac{(k - 1)\tau W}{\tau W + \gamma}$.

The above results suggest that for the same average link weight and when the one-to-one correspondence between $p_1$ and $k_1/k$, and $p_2$ and $k_2/k$ holds, the basic reproductive ratio is higher on networks with random weight distribution than on networks with a fixed weight distribution. This, however, does not preclude the possibility of having a network with random weight distribution with smaller average weight exhibiting an $R_0$ value that is bigger than the $R_0$ value corresponding to a network where weights are fixed and the average weight is higher. The direct implication is that it is not sufficient to know just the average link weight in order to draw conclusions about possible epidemic outbreaks on weighted networks; rather one has to know the precise weight distribution that provides a given average weight.

Figure 1 shows how the basic reproductive ratio changes with the transmission rate $\tau$ for different weight distributions. When links on a homogeneous network are distributed at random (upper panel), the increase in the magnitude of one specific link weight (e.g. $w_1$) accompanied by a decrease in its frequency leads to smaller $R_0$ values. This is to be expected since the contribution of the different link types in this case is kept constant ($p_1w_1 = p_2w_2 = 0.5$) and this implies that the overall weight of the network links accumulates in a small number of highly weighted links with most links displaying small weights and thus making transmission less likely. The statement above is more rigorously underpinned by
the results of Theorem 1 & 2 which clearly show that equal or more homogeneous weights lead to higher values of the basic reproductive ratio. For the case of fixed weight distribution (lower panel), the changes in the value of $R_0$ are investigated in terms of varying the weights, so that the overall weight in the network remains constant. This is constrained by fixing values of $p_1$ and $p_2$ and, in this case, the highest values are obtained for higher values of $w_1$. The flexibility here is reduced due to $p_1$ and $p_2$ being fixed, and a different link breakdown may lead to different observations. The top continuous line in Fig. 1 (upper panel) corresponds to the maximum $R_0$ value achievable for both models if the $p_1w_1 + p_2w_2 = 1$ constraint is fulfilled.

3.2 $R_0$-like threshold for the SIR model - a pairwise model perspective

To compute the value of $R_0$-like quantity from the pairwise model, we use the approach suggested by Keeling [32], which utilises the local spatial/network structure and correctly accounts for correlations between susceptible and infectious nodes early on in the epidemics. This can be achieved by looking at the early behaviour of $[SI]_1/[I] = \lambda_1$ and $[SI]_2/[I] = \lambda_2$ when considering links of only two different weights. In line with Eames [20], we start from the evolution equation of $[I]$

$$[I] = (\tau w_1 [SI]_1/[I] + \tau w_2 [SI]_2/[I] - \gamma)[I],$$

where from the growth rate $\tau w_1 \lambda_1 + \tau w_2 \lambda_2 - \gamma$ it is easy to define the threshold quantity $R$ as follows,

$$R = \frac{\tau w_1 \lambda_1 + \tau w_2 \lambda_2}{\gamma}. \quad (11)$$

For the classic closure (5), one can compute the early quasi-equilibria for $\lambda_1$ and $\lambda_2$ directly from the pairwise equations as follows

$$\lambda_1 = \frac{\gamma(k-1)p_1 R}{\tau w_1 + \gamma R} \quad \text{and} \quad \lambda_2 = \frac{\gamma(k-1)(1-p_1) R}{\tau w_2 + \gamma R}.$$

Substituting these into (11) and solving for $R$ yields

$$R = \frac{R_1 + R_2 + \sqrt{(R_1 + R_2)^2 + 4R_1R_2Q}}{2}, \quad (12)$$

where

$$R_1 = \frac{\tau w_1 [(k-1)p_1 - 1]}{\gamma}, \quad R_2 = \frac{\tau w_2 [(k-1)p_2 - 1]}{\gamma},$$

$$Q = \frac{k - 2}{[(k-1)p_1 - 1][(k-1)p_2 - 1]},$$

with details of all calculations presented in Appendix D. We note that $R > 1$ will result in an epidemic, while $R < 1$ will lead to the extinction of the disease. It is straightforward to show that for equal weights, say $W$, the expression above reduces to $R = \frac{\tau W (k-2)}{\gamma}$ which is in line with $R_0$ value in [32] for unclustered, homogeneous networks. Under the assumption of a fixed total weight $W$, one can show that similarly to the network-based basic reproductive ratio, $R$ achieves its maximum when $w_1 = w_2 = W$.  

In a similar way, for the modified closure (6), we can use the same methodology to derive the threshold quantity as

\[ R = \frac{R_1 + R_2 + \sqrt{(R_1 + R_2)^2 + 4R_1R_2(Q - 1)}}{2}, \]  

(13)

where

\[ R_1 = \frac{\tau w_1(k_1 - 2)}{\gamma}, \quad R_2 = \frac{\tau w_2(k_2 - 2)}{\gamma}, \quad Q = \frac{k_1k_2}{(k_1 - 2)(k_2 - 2)}. \]

For this closure once again, \( R > 1 \) results in an epidemic, while for \( R < 1 \), the disease dies out. Details of this calculation are shown in Appendix D. It is noteworthy that one can derive expressions (12) and (13) by considering the leading eigenvalue based on the linear stability analysis of the disease-free steady state of system (4) with the corresponding pairwise closures given in (5) and (6).

Finally, we note that this seemingly \( R_0 \)-lookalike, \( R = \tau W(k - 2)/\gamma \) for the equal weights case \( w_1 = w_2 = W \) is a multiple of \((k - 2)\) as opposed to \((k - 1)\) as is the case for the \( R_0 \) derived based on the individual-based perspective, where, for equal weights, \( R_0 = R_0^* = \tau W(k - 1)/(\tau W + \gamma) \). This highlights the importance, in models that are based on an underlying network of population interactions, of the way in which an \( R_0 \)-like quantity is defined. In simple mass-action-type models the same value is derived whether \( R_0 \) is thought of as the number of new cases from generation-to-generation (the NGM method), or as the growth rate of the epidemic scaled by the infectious period. In a network model the two approaches have the same threshold behaviour, but the clusters of infection that appear within the network mean that they produce different values away from the threshold. It is important therefore to be clear about what we mean by “\( R_0 \)” in a pair-approximation model. It is also important when using empirically-derived \( R_0 \) values to inform pair-approximation models to be clear about how these values were estimated from epidemiological data, and to consider which is the most appropriate way to incorporate the information into the model.

3.3 The performance of pairwise models and the impact of weight distributions on the dynamics of epidemics

To evaluate the efficiency of the pairwise approximation models, we will now compare numerical solutions of models (3) and (4) (with closures given by Eq. (5) and Eq. (6) for random and deterministic weights distributions, respectively) to results obtained from the corresponding network simulation. The discussion around the comparison of the two models is interlinked with the discussion of the impact of different weight distributions/patterns on the overall epidemic dynamics. We begin our numerical investigation by considering weight distributions with moderate heterogeneity. This is illustrated in Fig. 2, where excellent agreement between simulation and pairwise models is obtained. The agreement
remains valid for both SIS and SIR dynamics, and networks with higher average link weight lead to higher prevalence levels at equilibrium for SIS and higher infectiousness peaks for SIR.

Next, we explore the impact of weight distribution under the condition that the average weight remains constant (i.e. \( p_1 w_1 + p_2 w_2 = 1 \), where without loss of generality the average weight has been chosen to be equal to 1). First, we keep the proportion of edges of type one (i.e. with weight \( w_1 \)) fixed and change the weight itself by gradually increasing its magnitude. Due to the constraint on the average weight and the condition \( p_2 = 1 - p_1 \), the other descriptors of the weight distribution follow. Fig. 3 shows that concentrating a large portion of the total weight on a few links leads to smaller epidemics, since the majority of links are low-weight and thus have a small potential to transmit the disease. This effect is exacerbated for the highest value of \( w_1 \); in this case 95% of the links are of weight \( w_2 = (1 - p_1 w_1)/(1 - p_1) = 0.5/0.95 \) leading to epidemics of smallest impact (Fig. 3(a)) and smallest size of outbreak (Fig. 3(b)).

While the previous setup kept the frequency of links constant while changing the weights, one can also investigate the impact of keeping at least one of the weights constant (e.g. the larger one) and changing its frequency. To ensure a fair comparison, here we also require that the average link weight over the whole network is kept constant. When such highly weighted links are rare, the system approaches the non-weighted network limit where the transmission rate is simply scaled by \( w_2 \) (the most abundant link type). As Fig. 4 shows, in this case, the agreement is excellent, and as the frequency of the highly weighted edges/links increases, disease transmission is less severe.

Regarding the comparison of the pairwise and simulation models, we note that while the agreement is generally good for a large part of the disease and weight parameter space, the more extreme scenarios of weight distribution result in poorer agreement. This is illustrated in both Figs. 3 and 4 (see bottom curves), with the worst agreement for the SIS dynamics. The insets in Fig. 3 show that increasing the average connectivity improves the agreement. However, the cause of disagreement is due to a more subtle effect driven also by the weight distribution. For example, in Fig. 4, the average degree in the network is 10, higher then used previously and equal to that in the insets from Fig. 3, but despite this, the agreement is still poor.

The two different weighted network models are compared in Fig. 5. This is done by using the same link weights and setting \( p_1 = k_1/k \) and \( p_2 = k_2/k \). Epidemics on network with random weight distribution grow faster and, given the same time scales of the epidemic, this is in line with results derived in Theorem 1 & 2 and findings concerning the growth rates. The difference is less marked for larger values of \( \tau \) where a significant proportion of the nodes becomes infected.

In Fig. 6 the link weight composition is altered by decreasing the proportion of highly-weighted links. As expected, the reduced average link weight across the network leads to epidemics of smaller size while keeping the excellent agreement between simulation and pairwise model results.
4 Discussion

The present study has explored the impact of weight heterogeneity and highlighted that the added heterogeneity of link weights does not manifest itself in the same way as most other heterogeneities in epidemic models on networks. Usually, heterogeneities lead to an increase in $R_0$ but potentially for final epidemic size to fall [35]. However, for weighted networks the concentration of infectiousness on fewer target link, and thus target individuals, leads to a fall in $R_0$ for both homogeneous random and fixed weight distribution models. Increased heterogeneity in weights accentuates the locality of contact and is taking the model further from the mass-action type models. Infection is concentrated along a smaller number of links, which results in wasted infectivity and lower $R_0$. This is in line with similar results [11, 12, 49] where different modelling approaches have been used to capture epidemics on weighted networks.

The models proposed in this paper are simple mechanistic models with basic weight distributions, but despite this they provide a good basis for analysing disease dynamics on weighted networks in a rigorous and systematic way. The modified pairwise models have performed well, and provide good approximation to direct simulation. As expected, the agreement with simulations typically breaks down at or close to the threshold, but away from it, pairwise models provide a good counterpart or alternative to simulation. Disagreement only appears for extreme weight distributions, and we hypothesise that this is mainly due to the network becoming more modular with islands of nodes connected by links of low weight being bridged together by highly weighted links. A good analogy to this is provided by considering the case of a pairwise model on unweighted networks specified in terms of two network metrics, node number $N$ and average number of links $k$.

The validity of the pairwise model relies on the network being connected up at random, or according to the configuration model. This can be easily broken by creating two sub-networks of equal size both exhibiting the same average connectivity. Simulations on such type of networks will not agree with the pairwise model, and highlights that the network generating algorithm can push the network out of the set of ‘acceptable’ networks. We expect that this or similar argument can more precisely explain why the agreement breaks down for significant link-weight heterogeneity.

The usefulness of pairwise models is illustrated in Fig. 7, where the $I/N$ values are plotted for a range of $\tau$ values and for different weight distributions. Here, the equilibrium value has been computed by finding the steady state directly from the ODEs (3) by finding numerically the steady state solution of a set on nonlinear equations (i.e. $\dot{A} = 0$ and $\dot{AB} = 0$). To test the validity, the long term solution of the ODE is plotted along with results based on simulation. The agreement away from the threshold is excellent and illustrates clearly the impact of different weight distributions on the magnitude of the endemic threshold.

The models proposed here can be extended in a number of different ways. One potential avenue for further research is the analysis of correlations between link weight and node degree. This direction has been explored but in the context of classic compartmental mean-field models based on node degree [31, 40]. Given that pairwise models extend to heterogeneous networks such avenues can be further explored to include different type of correlations or other network dependent weight distributions. While this is a viable direction, it is expected that the extra
complexity will make the pairwise models more difficult to analyse and disagree-
ment between pairwise and simulation model more likely. Another theoretically
interesting and practically important aspect is the consideration of different types
of time delays, representing latency or temporary immunity [9], and the analysis
of their effects on the dynamics of epidemics on weighted networks. The method-
ology presented in this paper can be of wider relevance to studies of other natural
phenomena where overlay networks provide effective description. Examples of such
systems include the simultaneous spread of two different diseases in the same pop-
ulation [8], the spread of the same disease but via different routes [35] or the spread
of epidemics concurrently with information about the disease [28,36]. These areas
offer other important avenues for further extensions.

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5 Appendix

5.1 Appendix A - Reducing the weighted pairwise models to the unweighted equivalents

We start from the system

\[
\dot{S} = \gamma [I] - \tau \sum_{n=1}^{M} w_n [SI]_n,
\]
\[
\dot{I} = \tau \sum_{n=1}^{M} w_n [SI]_n - \gamma [I],
\]

\[
[\dot{SI}]_m = \gamma ([II]_m - [SI]_m) + \tau \sum_{n=1}^{M} w_n ([SSI]_{mn} - [ISI]_{nm}) - \tau w_m [SI]_m,
\]

\[
[\dot{II}]_m = -2\gamma [II]_m + 2\tau \sum_{n=1}^{M} w_n [ISI]_{nm} + 2\tau w_m [SI]_m,
\]

\[
[\dot{SS}]_m = 2\gamma [SI]_m - 2\tau \sum_{n=1}^{M} w_n [SSI]_{mn},
\]

(14)

where \( m = 1, 2, \ldots, M \). To close this system of equations at the level of pairs, we use the approximations

\[
[ABC]_{mn} = \frac{k - 1}{k} \frac{[AB]_m [BC]_n}{[B]}.
\]

To reduce these equations to the standard pairwise model for unweighted networks we use the fact that \( \sum_{m=1}^{M} [AB]_m = [AB] \) for \( A, B \in \{S, I\} \) and aim to derive the evolution equation for \( [AB] \). Assuming that all weights are equal to some \( W \), the following relations hold,

\[
[\dot{SI}] = \sum_{m=1}^{M} [\dot{SI}]_m
\]

\[
= \sum_{m=1}^{M} \left( \gamma ([II]_m - [SI]_m) + \tau \sum_{n=1}^{M} w_n ([SSI]_{mn} - [ISI]_{nm}) - \tau w_m [SI]_m \right)
\]

\[
= \gamma ([II] - [SI]) - \tau W [SI] + \tau W \sum_{m=1}^{M} \sum_{n=1}^{M} ([SSI]_{mn} - [ISI]_{nm}),
\]

where the summations of the triples can be resolved as follows,

\[
\sum_{m=1}^{M} \sum_{n=1}^{M} [SSI]_{mn} = \frac{k - 1}{k} \frac{M - 1}{M} \frac{M - 1}{M} [SI]_m \sum_{n=1}^{M} [SI]_n
\]

\[
= \frac{k - 1}{k} \frac{[SS][SI]}{[S]} = [SSI].
\]

Using the same argument for all other triples, the pairwise model for weighted networks with all weights being equal (i.e. \( W = 1 \)) reduces to the classic pairwise
model, that is
\[
\dot{S} = \gamma[I] - \tau[S], \\
\dot{I} = \tau[S] - \gamma[I], \\
\sum_{m=1}^{M} \dot{SI} = \dot{SI} = \gamma([II] - [SI]) + \tau[SSI] - [SI], \\
\sum_{m=1}^{M} \dot{II} = \dot{II} = -2\gamma[II] + 2\tau([ISI] + [SI]), \\
\sum_{m=1}^{M} \dot{SS} = \dot{SS} = 2\gamma[S] - 2\tau[SSI].
\]

A similar argument holds for the pairwise model on weighted networks with SIR dynamics.

5.2 Appendix B - Proof of Theorem 1

We illustrate the main steps needed to complete the proof of Theorem 1. This revolves around starting from the inequality itself and showing via a series of algebraic manipulations that it is equivalent to a simpler inequality that holds trivially. Upon using that \(p_1 k = k_1, p_2 k = k_2\) and \(p_2 = 1 - p_1\), the original inequality can be rearranged to give
\[
\sqrt{(k_1 - 1)r_1 - (k_2 - 1)r_2}^2 + 4k_1 k_2 r_1 r_2 \leq (k_1 - 1)r_1 + (k_2 - 1)r_2 + 2r_1 p_2 + 2r_2 p_1.
\]

Based on the assumptions of the Theorem, the right-hand side is positive, and thus this inequality is equivalent to the one where both the left- and right-hand sides are squared. Combined with the fact that \(p_2 = 1 - p_1\), after a series of simplifications and factorizations this inequality can be recast as
\[
4p_1 (1 - p_1)(r_1^2 + r_2^2) + 8kp_1 (1 - p_1)r_1 r_2 \leq 4kp_1 (1 - p_1)(r_1^2 + r_2^2) + 8p_1 (1 - p_1)r_1 r_2,
\]
which can be further simplified to
\[
4p_1 (1 - p_1)(r_1 - r_2)^2(k - 1) \geq 0,
\]
which holds trivially and thus completes the proof. We note that in the strictest mathematical sense the condition of the Theorem should be \((k_1 - 1)r_1 + (k_2 - 1)r_2 + 2r_1 p_2 + 2r_2 p_1 \geq 0\). This holds if the current assumptions are observed since these are stronger but follow from a practical reasoning whereby for the network with fixed weight distribution, a node should have at least one link with every possible weight type.

5.3 Appendix C - Proof of Theorem 2

First, we show that \(R_1^0\) is maximised when \(w_1 = w_2 = W\). \(R_1^0\) can be rewritten to give
\[
R_1^0 = (k - 1) \left( p_1 \frac{\tau w_1}{\tau w_1 + r} + (1 - p_1) \frac{\tau w_2}{\tau w_2 + r} \right).
\]
Maximising this given the constraint \( w_1 p_1 + w_2 (1 - p_1) = W \) can be achieved by considering \( R_0^2 \) as a function of the two weights and incorporating the constraint into it via the Lagrange multiplier method. Hence, we define a new function \( f(w_1, w_2, \lambda) \) as follows

\[
f(w_1, w_2, \lambda) = (k - 1) \left( p_1 \frac{\tau w_1}{\tau w_1 + r} + (1 - p_1) \frac{\tau w_2}{\tau w_2 + r} \right) + \lambda (w_1 p_1 + w_2 (1 - p_1) - W).\]

Finding the extrema of this function leads to a system of three equations

\[
\frac{\partial f}{\partial w_1} = \frac{(k - 1) p_1 \tau \gamma}{(\tau w_1 + \gamma)^2} + \lambda p_1 = 0,
\]

\[
\frac{\partial f}{\partial w_2} = \frac{(k - 1) (1 - p_1) \tau \gamma}{(\tau w_2 + \gamma)^2} + \lambda (1 - p_1) = 0,
\]

\[
w_1 p_1 + w_2 (1 - p_1) - W = 0.
\]

Expressing \( \lambda \) from the first two equations and equating these two expressions yields

\[
\frac{(k - 1) \tau \gamma}{(\tau w_1 + \gamma)^2} = \frac{(k - 1) \tau \gamma}{(\tau w_2 + \gamma)^2}.
\]

Therefore,

\[
w_1 = w_2 = W.
\]

and it is straightforward to confirm that this is a maximum.

Performing the same analysis for \( R_0^2 \) is possible but it is more tedious. Instead, we propose a more elegant argument to show that \( R_0^2 \) under the constraint of constant total link weight achieves its maximum when \( w_1 = w_2 = W \). The argument starts by considering \( R_0^2 \) when \( w_1 = w_2 = W \). In this case, and using that

\[
r_2 = r_1 = r = \tau W/(\tau W + \gamma)
\]

we can write,

\[
R_0^{2*} = \frac{(k_1 - 1) r_1 + (k_2 - 1) r_2 + \sqrt{[(k_1 - 1) r_1 - (k_2 - 1) r_2]^2 + 4 k_1 k_2 r_1 r_2}}{2}
\]

\[
= \frac{r (k_1 + k_2 - 2) + \sqrt{r^2 [(k_1 - 1) - (k_2 - 1)]^2 + 4 r^2 k_1 k_2}}{2}
\]

\[
= \frac{r (k_1 + k_2 - 2) + r \sqrt{(k_1 + k_2)^2}}{2}
\]

\[
= \frac{r (2k_1 + 2k_2 - 2)}{2} = r (k_1 + k_2 - 1) = (k - 1) r.
\]

However, it is known from Theorem 1 that \( R_0^2 \leq R_0^1 \), and we have previously shown that \( R_0^2 \) under the present constraint achieves its maximum when \( w_1 = w_2 = W \), and its maximum is equal to \((k - 1) r\). All the above can be written as

\[
R_0^2 \leq R_0^1 \leq (k - 1) r.
\]
Now taking into consideration that $R_0^2 = (k-1)r$, the inequality above can be written as

$$R_0^2 \leq R_0^1 \leq (k-1)r = R_0^2,$$

and this concludes the proof.

5.4 Appendix D - The $R_0$-like threshold $R$

Let us start from the evolution equation for $[I](t)$,

$$[\dot{I}] = \tau(w_1[S]_1 + w_2[S]_2) - \gamma[I]$$

$$= \left[\tau w_1 \left( \frac{[S]_1}{[I]} \right) + \tau w_2 \left( \frac{[S]_2}{[I]} \right) - \gamma \right][I]$$

$$= (\tau w_1 \lambda_1 + \tau w_2 \lambda_2 - \gamma)[I],$$

where $\lambda_1 = \frac{[S]_1}{[I]}$ and $\lambda_2 = \frac{[S]_2}{[I]}$, and let $R$ be defined as

$$R = \frac{\tau w_1 \lambda_1 + \tau w_2 \lambda_2}{\gamma}. \quad (23)$$

Following the method outlined by Keeling [32] and Eames [20], we calculate the early quasi-equilibrium values of $\lambda_1, \lambda_2$ as follows:

$$\dot{\lambda}_1 = 0 \Leftrightarrow \dot{[SI]}_1[I] = [\dot{I}][SI]_1,$$

$$\dot{\lambda}_2 = 0 \Leftrightarrow \dot{[SI]}_2[I] = [\dot{I}][SI]_2.$$

Upon using the pairwise equations and the closure, consider $[\dot{SI}]_1[I] = [\dot{I}][SI]_1$:

$$[\dot{SI}]_1[I] = (\tau w_1 [SS]_1_{11} + \tau w_2 [SS]_1_{12} - \tau w_1 [IS]_1_{11} - \tau w_2 [IS]_1_{21} - \tau w_1 [SI]_1 - \gamma [SI]_1[I])$$

$$= (\tau w_1 [SI]_1 + \tau w_2 [SI]_2 - \gamma[I])[SI]_1.$$

Using the classical closure

$$[ABC]_{12} = \frac{k - 1}{k} \frac{[AB]_1 [BC]_2}{[B]},$$

$$[ABC]_{21} = \frac{k - 1}{k} \frac{[AB]_2 [BC]_1}{[B]},$$

and making the substitution : $[SI]_1 = \lambda_1[I]$, $[SI]_2 = \lambda_2[I]$, $[I] \ll 1$, $[S] \approx N$,

$[SS]_1 \approx kNp_1$, $[SS]_2 \approx kN(1-p_1)$ together with $\gamma R = \tau w_1 \lambda_1 + \tau w_2 \lambda_2$, we have

$$(\tau w_1 \lambda_1 + \tau w_2 \lambda_2)kp_1 - (\tau w_1 \lambda_1 + \tau w_2 \lambda_2)p_1 - (\tau w_1 \lambda_1 + \tau w_2 \lambda_2)\lambda_1 - \tau w_1 \lambda_1 = 0,$$

which can be solved for $\lambda_1$ to give

$$\lambda_1 = \frac{\gamma (k - 1)p_1 R}{\tau w_1 + \gamma R}. $$
Similarly, $\lambda_2$ can be found as
\[
\lambda_2 = \frac{\gamma(k-1)(1-p_1)R}{\tau w_2 + \gamma R}.
\] (25)

Substituting the expressions for $\lambda_1, \lambda_2$ into the original equation for $R$ yields
\[
R = \frac{A + B + \sqrt{(A + B)^2 + 4\tau^2 w_1 w_2 (k-2)}}{2\gamma}
\]
where $A = \tau w_1 [(k-1)p_1 - 1]$ and $B = \tau w_2 [(k-1)p_2 - 1]$. If we define
\[
R_1 = \frac{\tau w_1 [(k-1)p_1 - 1]}{\gamma}, \quad \text{and} \quad R_2 = \frac{\tau w_2 [(k-1)p_2 - 1]}{\gamma},
\]
the expression simplifies to
\[
R = \frac{R_1 + R_2 + \sqrt{(R_1 + R_2)^2 + 4R_1 R_2 Q}}{2}
\]
where $Q = \frac{(k-2)}{[(k-1)p_1 - 1][(k-1)p_2 - 1]}$.

Substituting the modified closure
\[
[ABC]_{11} = \frac{k_1 - 1}{k_1} \frac{[AB]_1[BC]_1}{[B]},
\]
\[
[ABC]_{12} = \frac{[AB]_1[BC]_2}{[B]},
\]
\[
[ABC]_{21} = \frac{[AB]_2[BC]_1}{[B]},
\]
\[
[ABC]_{22} = \frac{k_2 - 1}{k_2} \frac{[AB]_2[BC]_2}{[B]},
\]
into (24) and making further substitution: $[SI]_1 = \lambda_1[I], [SI]_2 = \lambda_2[I], [I] \ll 1, [S] \approx N, [SS]_1 \approx k_1 N, [SS]_2 \approx k_2 N$, we have
\[
(\tau w_1 \lambda_1 + \tau w_2 \lambda_2)k_1 - (\tau w_1 \lambda_1 + \tau w_2 \lambda_2)\lambda_1 - 2\tau w_1 \lambda_1 = 0 \implies \lambda_1 = \frac{\gamma k_1 R}{2\tau w_1 + \gamma R}.
\]
Similarly, the equation $[SI]_2[I] = [I][SI]_2$ yields
\[
\lambda_2 = \frac{\gamma k_2 R}{2\tau w_2 + \gamma R}.
\]

Substituting these expressions for $\lambda_1, \lambda_2$ into (23), we have
\[
R = \frac{\tau (w_1 k_1 + w_2 k_2) - 2\tau (w_1 + w_2)}{2\gamma}
\]
\[
+ \frac{\sqrt{[2\tau (w_1 + w_2) - \tau (w_1 k_1 + w_2 k_2)]^2 + 8\tau^2 w_1 w_2 (k_1 + k_2 - 2)}}{2\gamma}.
\]
If we define
\[ R_1 = \frac{\tau w_1(k_1 - 2)}{\gamma}, \quad R_2 = \frac{\tau w_2(k_2 - 2)}{\gamma}, \]
the above expression for \( R \) simplifies to
\[ R = \frac{R_1 + R_2 + \sqrt{(R_1 + R_2)^2 + 4R_1R_2(Q - 1)}}{2} \quad (26) \]
where
\[ Q = \frac{k_1k_2}{(k_1 - 2)(k_2 - 2)}. \]
References


Fig. 1 Basic reproductive ratio $R_0$ for random (upper) and deterministic (lower) weight distributions with different weight and weight frequency combinations, but with $p_1 w_1 + p_2 w_2 = 1$. Upper panel: the case of homogenous networks with weights assigned at random considers the situation where the contribution of the two different weight types is equal ($p_1 w_1 = p_2 w_2 = 0.5$) but with weight $w_1$ increasing and its frequency decreasing (top to bottom with $(p_1, w_1) = \{(0.5, 1), (0.2, 25), (0.05, 10)\}$). Increasing the magnitude of weights but reducing their frequency leads to smaller $R_0$ values. Lower panel: the case of homogeneous networks with fixed number of links of type $w_1$ and $w_2$ illustrates the situation where $w_1$ increases while $p_1 = k_1/k = 1/3$ and $p_2 = (k - k_1) / k = 2/3$ remain fixed (bottom to top with $w_1 = \{0.1, 0.5, 1.4\}$). Here the opposite tendency is observed with increasing weights leading to higher $R_0$ values. Finally, for the randomly distributed weights case, setting $p_1 = 1/3$, $w_1 = 1.4$ and observing $p_1 w_1 + p_2 w_2 = 1$, we obtain $R_0$ $(\star)$ values which compare almost directly to the fixed-weights case (top continuous line). Other parameters are set to $k = 6$, $k_1 = 2$ and $\gamma = 1$. 
Fig. 2 The infection prevalence ($I/N$) from the pairwise and simulation models for homogeneous random networks with random weight distribution (ODE: solid line, simulation: dashed line and (o)). All nodes have degree $k = 5$ with $N = 1000$, $I_0 = 0.05N$, $\gamma = 1$ and $\tau = 1$. From top to bottom, the parameter values are: $w_1 = 5, p_1 = 0.2, w_2 = 1.25, p_2 = 0.8$ (top), and $w_1 = 0.5, p_1 = 0.5, w_2 = 1.5, p_2 = 0.5$ (bottom). The left and right panels represent the SIS and SIR dynamics, respectively.

Fig. 3 The infection prevalence ($I/N$) from the pairwise and simulation models for homogeneous networks with random weight distribution (ODE: solid line, simulation: dashed line and (o)). All numerical tests use $N = 1000$, $I_0 = 0.05N$, $k = 5$, $\gamma = 1$, $\tau = 1$ and $p_1 = 0.05$ ($p_2 = 1 - p_1 = 0.95$). From top to bottom, $w_1 = 2.5, 5, 10, w_2 = 0.875/0.95, 0.75/0.95, 0.5/0.95$. The weight distributions are chosen such that the average link weight, $p_1w_1 + p_2w_2 = 1$, remains constant. Insets of (a) and (b): the same parameter values as for the lowest prevalence plots but, with $k = 10$ and $\tau = 0.5$. The left and right panel represent the SIS and SIR dynamics, respectively.
Fig. 4 The infection prevalence ($I/N$) from the pairwise and simulation model for homogenous networks with random weight distribution (ODE: solid line, simulation: dashed line and (o)). All numerical tests use $N = 1000$, $I_0 = 0.05N$, $k = 10$, $\gamma = 1$, $\tau = 0.5$ and $w_1 = 10$. From top to bottom, $P(w_1) = 0.01, 0.05, 0.09$, $w_2 = 0.9/0.99, 0.5/0.95, 0.1/0.91$. Here also $p_2 = 1 - p_1$ and $p_1w_1 + p_2w_2 = 1$. The left and right panel represent the SIS and SIR dynamics, respectively.

Fig. 5 The infection prevalence ($I/N$) based on random (model 1) and fixed (model 2) weight distribution (ODE: black (1) and blue (2) solid line, simulation results: same as ODE but dashed lines, and (o) and (+)). All numerical tests use $N = 1000$, $I_0 = 0.05N$, $k = 10$, $k_1 = 2$, $k_2 = 8$, $p_1 = k_1/k$, $p_2 = k_2/k$, $w_1 = 10$, $w_2 = 1.25$ and $\gamma = 1$. The rate of infection $\tau = 0.5$ (top) and $\tau = 0.1$ (bottom). The left and right panel represent the SIS and SIR dynamics, respectively.
Fig. 6 The infection prevalence ($I/N$) for a fixed weight distribution (ODE: solid lines, simulation results: dashed lines and (o)). All numerical tests use $N = 1000$, $I_0 = 0.05N$, $k = 6$, $\gamma = 1$, $\tau = 1$ and $w_1 = 1.4$, $w_2 = 0.8$. From top to bottom: $k_1 = 5, 4, 3, 2, 1$ and $k_2 = k - k_1$. The left and right panel represent the SIS and SIR dynamics, respectively.

Fig. 7 Endemic steady state from the SIS model on networks with random weight distribution. The continuous lines correspond to the steady state computed numerically by setting all evolution equations in the pairwise system to zero. These are complemented by finding the endemic steady state through direct integration of the ODE system for a long-enough time (o), as well as direct simulation (*). The first marker corresponds to $\tau = 0.3$ followed by $\tau = 0.5, 1.0, \ldots, 3.0$. All results are based on: $k = 5$, $\gamma = 1$ and $w_1 = 10$, $w_2 = 1$. From top to bottom: $p_1 = 0.9, 0.5, 0.1, 0.01$ and $p_2 = 1 - p_1$. 