

Epidemic spread in networks: Existing methods and current challenges

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Abstract

We consider the spread of infectious disease through contact networks of Configuration Model type. We assume that the disease spreads through contacts and infected individuals recover into an immune state. We discuss a number of existing mathematical models used to investigate this system, and show relations between the underlying assumptions of the models. In the process we offer simplifications of some of the existing models. The distinctions between the underlying assumptions are subtle, and in many if not most cases this subtlety is irrelevant. Indeed, under appropriate conditions the models are equivalent. We compare the benefits and disadvantages of the different models, and discuss their application to other populations (*e.g.*, clustered networks). Finally we discuss ongoing challenges for network-based epidemic modeling.

1 Introduction

Mathematical models of infectious disease spread have played a significant role in improving our understanding of epidemics and developing better intervention strategies [1]. Invariably, these models make simplifying assumptions in order to arrive at tractable equations.

Recently, considerable research has focused on eliminating some of the standard assumptions of mathematical models. There has been particular emphasis on understanding how contact networks influence disease spread [40, 36, 25, 26, 39, 15, 16, 19, 5]. In this paper we focus on research into two features of contact networks: Partnerships have nonzero duration and different individuals have different numbers of partners. We will focus our attention on “SIR” diseases, diseases in which individuals begin susceptible, become infected through partnerships with infected individuals, and may eventually recover into an immune class.

The models we investigate in this paper consider the spread of an infectious disease through a population whose partnerships are static. The populations have “Configuration Model” structure: each individual is assigned a number of stubs (its degree, k), and then finds partners for each stub randomly from the available stubs [37, 34]. So if $P(k)$ is the probability a random individual u has k partners, the probability that a partner v has k' partners is $k'P(k')/\langle K \rangle$ where $\langle K \rangle$ is the average degree — there is a size bias because an individual’s partners are selected proportional to their degrees. We assume that infected individuals transmit to each partner at a fixed rate β and recover at fixed rate γ .

Several competing systems of equations have been developed to capture the population-scale dynamics of disease spread in these populations. These models have varying levels of complexity and detail. Each model makes assumptions about the independence of individuals that appear in the form of a closure. The assumptions are subtly different. The resulting models are known to be closely related [45, 11]. We introduce these models and provide their derivations. We clarify these relations and show that in appropriate cases they are identical. We discuss their advantages and disadvantages for different epidemiological questions

*JCM dedicates this work to the memory of Bob Borrelli, who taught him how to use integrating factors and much more.

and present more recent directions of network-based infectious disease modeling and end with a discussion of ongoing challenges. We include an appendix which shows the mathematical relationships between the models by demonstrating how one can derive some models from others.

2 The existing models

We study five different models having three different basic approaches. The three approaches are characterized as “pairwise” models, “effective degree” models, and “edge-based compartmental” models. In all cases, the disease is assumed to transmit at rate β from an infected individual to its partner (which may or may not be susceptible). If the partner is susceptible, the transmission results in immediate infection. Infected individuals recover at rate γ and become immune. One notable model to incorporate partnership duration that we do not investigate is the “renewal equation” approach of [5].

We give a brief heuristic explanation of the approaches here. The “pairwise models” observe that the rate susceptible individuals become infected is proportional to the number of partnerships between susceptible and infected individuals. The main effort of these models is in tracking how the number of such partnerships change in time. The “effective degree” models focus on the number of individuals with a given number of “active” partnerships. These models “discard” partnerships once it is clear that they will no longer play a role in transmission (for example, if a partner v of u recovers, we no longer have to track the v - u partnership). Consequently these models stratify individuals by their “effective” degree. Finally, the “edge-based compartmental” models focus on the probability that a partner v has transmitted to u , creating compartments showing the probability v has transmitted, or — if it has not yet transmitted — whether it is susceptible, infected, or recovered.

2.1 Closures

The concept of a “closure” comes up across many branches of applied mathematics [41, 18, 17, 46, 43, 42]. Often we are interested in how some physical quantity is distributed at one scale, but calculating that requires knowledge about its distribution at a larger (or smaller) scale. However, to calculate that larger (or smaller) scale, requires its distribution at another yet larger (or smaller) scale. This leads to an infinite cascade of scales, and an infinite sequence of equations. To truncate this system, an assumption is made that at some scale the distribution can be calculated in terms of the distribution at some previous scale. This results in a finite system of equations which can be solved. The error, if any, introduced by the closure determines the accuracy of the solution. The closures we will use can be thought of as similar to the concept of maximal entropy [13]. We will assume that at some scale in the network there is no useful information contained in larger scales, so we will express our equations in terms of the smaller scale.

For the problem of epidemic spread on networks, our ultimate goal is equations giving the proportion of the population which is susceptible, infected, or recovered. An epidemic is an inherently stochastic process, so when we write down deterministic equations, we are implicitly assuming that the actual proportion in each state is closely approximated by the expected number in each state. Thus our equations are appropriate only in large populations with a sufficiently large number of infections at time $t = 0$. If the number of infections is too small, we can typically wait until the number infected has grown sufficiently, and then the equations will be accurate.

The rate at which new infections occur clearly depends on the number of partnerships between susceptible and infected individuals. In turn, the distribution of such pairs depends on the distribution of triples (*e.g.*, an S - S pair becomes an S - I pair through infection introduced from a third individual). The frequency of various triples depends on still larger structures. This cascade of scales suggests that a closure may be needed. The various models that have been proposed differ in their choice of closure. There are three distinct closures which we will consider. These closures are faithful to the dynamics of the epidemic in the sense that if the initial distribution of infection satisfies the closure, then at later time the distribution of infection will still satisfy the closure. This faithfulness is a consequence of the assumption that the contact structure is a Configuration Model network. So if the contact structure is of a different type, then these closures may fail.

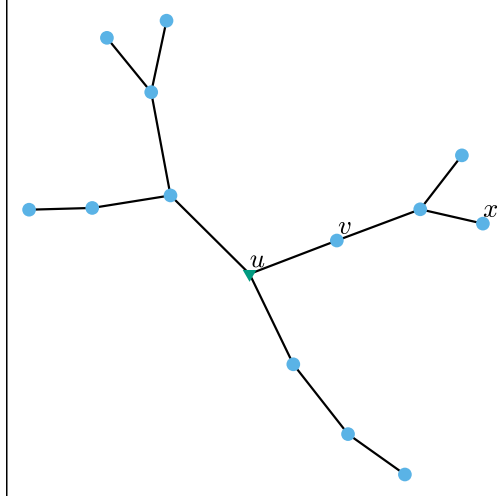


Figure 1: **A demonstration of “reachable” individuals.** Taken from a Configuration Model network with $P(2) = 0.8$ and $P(3) = 0.2$. In this case, we choose u and look at individuals within distance 3 from u . The individual x is reachable from u through v , but not through other partners of u . On a sufficiently large scale there will be cycles that connect the different branches. As the population grows, the size of these cycles grows as well.

Although Configuration Model networks have cycles in them, we are interested in the large network limit, for which the networks are “locally treelike”. The probability an individual is in a short cycle scales like $1/N$ where N is the population size. Thus when we look at u , we may neglect the distant features and think of the network as a tree with root u . If v is a partner of u and the path from u to x passes through v , we refer to x as “reachable” from u through v . Note that v is considered to be reachable from u through v . This is shown in figure 1.

We can understand the closures in terms of what information we gain about one partner of a randomly chosen susceptible individual u by considering other partners of u . We now assume that at time t , u is a randomly chosen susceptible individual and v is a partner of u . Each of our closures makes a different claim about what information about v (and individuals reachable from u through v) is provided by other partners of u (and individuals reachable through those partners). So long as u is susceptible there is no opportunity for dynamics on one branch from u to affect those on another branch. So we might anticipate that there is no information to be had about v from the other partners of a susceptible individual u . However depending on how the initial infections are chosen, there may be correlations built into the initial condition. Depending on what correlations are built into the initial condition, some, or all of our closures may be invalid. If so, then the equations derived assuming the closure is at best an approximation to the true dynamics. If there is no correlation built in to the initial condition then all of our closures hold. Even if there is correlation built in, we anticipate that information about the initial conditions will decay in time, so a closure that does not initially hold is likely to be a good approximation at later time.

Figure 2 demonstrates how a specially selected initial condition can give information about different partners of a random individual. On the left, 3 degree three individuals (denoted by triangles in the figure) are chosen from the population and their partners infected. If we now choose a random individual u from the population and observe whether one partner is infected or not, this gives us some information about whether u is one of the “triangle” individuals, and so we can update our expectation for the probability another partner is infected. On the right, the same number of initial infections are observed, but they are chosen uniformly from the population. If we choose an individual u and look at any partner of u , it gives no information about the other partners.

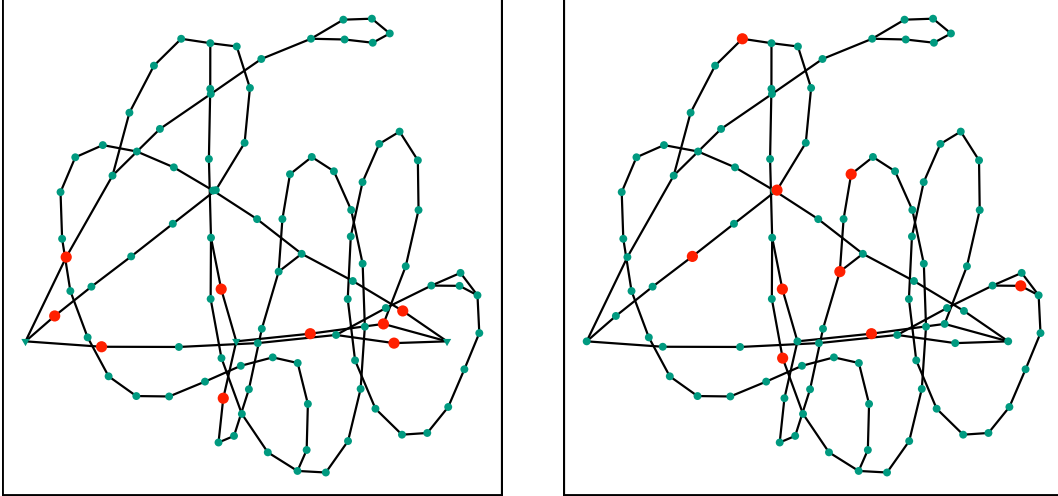


Figure 2: **Two different ways that the initial conditions could be chosen.** A Configuration Model network is used with $P(2) = 0.8$ and $P(3) = 0.2$. On the left, 3 individuals (denoted by triangles) are chosen uniformly from those with degree three and their partners (larger disks) are infected. On the right, the same number of initial infections are chosen, but randomly from the population. On the left, we would anticipate slower growth for the disease due to “wasted” transmissions to the triangular individuals. The star closure applies for the left, but not the pairs or triples closure. All closures apply on the right.

Triples closure We define a triple to be a set of three individuals for which there is a central individual who is a partner of both of the other two. Let $[A_p S_k B_q]$ to be the number of triples in the population for which the central individual has status S and degree k and the other two individuals have status A with degree p and status B with degree q (A and B are any of $S, I,$ or R). Further let $[A_p S_k]$ and $[S_k B_q]$ be the number of pairs of the given types.

The triples closure can be stated as follows: At time t we are given a randomly chosen susceptible individual u and its partner v . The probability that v has a given status may depend on the degrees of u and v . However, information we get about individuals reachable from u through other partners of u does not tell us anything about v or other individuals reachable from u through v .

A consequence of this assumption is that if the triples closure holds at time t , then at all later times the frequency of various triples (with susceptible central individual) can be expressed in terms of the number of pairs as

$$[A_p S_k B_q] = \frac{(k-1)[A_p S_k][S_k B_q]}{k[S_k]}$$

where $[S_k]$ is the number of susceptible individuals of degree k . So if the triples closure holds, everywhere a triple appears in our infinite cascade of equations, we can reduce it to pairs.

Star closure We define a star to be a central individual and all of its partners. Let $x_{s,i,r}$ denote the number of stars having a central susceptible individual and $s, i,$ and r susceptible, infected, and recovered peripheral individuals respectively.

The star closure can be stated as follows: If at time t we are given a randomly chosen susceptible individual u and its partner v , we may gain information about the status of v by knowing the statuses of other partners of u . However, we do not gain any information about v (or those reachable from u through v) by knowing anything about individuals reachable along longer paths from u through these other partners. If we additionally know that v was susceptible at time 0, then we cannot gain information about v from u or the other partners of u .

In other words, it may be possible to get information about what proportion of the peripheral individuals surrounding a susceptible individual u are in each state. However, if we know a partner v is susceptible, then it is not possible to learn anything more about v .

The implication of this assumption is that if the star closure holds at time t , then knowing the distribution of $x_{s,i,r}$ at time t allows us to predict its distribution at all later times.

Pairs closure The pairs closure states that conditional on u being susceptible at time t , there is nothing we can learn about u or individuals reachable from u through partners other than v which will give any information about v .

This assumption means that everywhere a pair involving a susceptible individual appears in our infinite cascade of equations, we can express it in terms of a random partner of a susceptible individual. Note that if the pairs closure holds, so do the other two closures.

One of the models we will investigate uses a slightly stronger assumption, which we refer to as the “strong” pairs closure. In this case, we make the assumption that even if u is infected, if there has been no transmission in either direction between u and v we can learn nothing about v . This assumption does not play a major role and appears only in the context of this particular model.

2.1.1 Validity of the closures

All of our closures are consistent with the dynamics: Assume at time t we cannot learn any information about a certain property of v from other partners of u (or individuals reachable through them). If u is still susceptible at a later time \hat{t} , there is no way for information to have travelled between v and the other partners of u . There is no way for any correlations to develop. This statement can be made more mathematically rigorous by looking at an infinite system of equations which describes the epidemic and introducing the closure to those equations. So long as the equations remain valid, the closure is compatible with the epidemic dynamics.

2.1.2 Examples of initial conditions for various closures

We now give some specific examples of initial conditions which may or may not violate some of the closures.

Assume that we color some subset of the individuals blue. If we infect those individuals at distance 2 from the blue individuals, none of our closures will hold. To see this, consider a randomly chosen susceptible individual u and a partner v . By observing whether those individuals a distance 2 from u which are reachable through partners other than v are infected (or not), we gain information about whether u was colored blue, which gives information about partners of v . This violates all of the closure assumptions.

In contrast, assume that as our initial condition at $t = 0$, we infect all degree 3 individuals which are partners of degree 2 individuals. There is no information that goes into this initial condition which depends on structures larger than pairs. So triples and larger structures can be reduced to pairs. The triples closure holds. The other closures do not hold. The star closure fails because if u is susceptible with degree 3, we immediately know that its partner v does not have degree 2, while if u is susceptible with degree 4 then v may have degree 2. This would give us information about v 's degree which in turn tells us something about the possible stars centered at v , which in turn means that v 's risk of infection depends on the degree of u . So we have learned something about the susceptible partner v . Similarly the pairs closure fails. If u has degree 3 and is susceptible, then we learn things about the degrees of its partners. Note that this example implies it is possible to satisfy the triples closure without satisfying the star closure.

We can also find conditions which violate the triples closure but satisfy the star closure. We color some subset of the population blue and at $t = 0$ infect all (or some uniformly chosen proportion p) of those individuals with at least one blue partner. Knowing the status of one partner of the susceptible individual u gives information about whether u was colored blue, which in turn gives information about the status of other partners of u . This violates the triples closure. However, while knowing the status of one partner w of u does give information about another partner v , we do not gain any information about v if v is initially susceptible. So the star closure holds. The pairs closure again fails for this case.

In all of the above examples, the pairs closure is violated. However, the violation comes because the initial conditions choose to infect individuals based on the location of some other individual who is likely to initially be susceptible. This seems rather artificial — it is difficult to conceive of a relevant scenario where this would happen. The pairs closure is satisfied in many cases, in particular if we initially infect some individuals uniformly chosen at random.

In general, even if the initial condition does not satisfy a given closure, we anticipate that as the disease spreads we will lose information about the initial condition. If so, then the closures will become valid. So in particular, if the initial condition involves infection of a small proportion of the population, we anticipate that the closures will all be valid by the time the epidemic has grown to appreciable size.

2.2 Pairwise models

Pairwise models are based on “pair approximation” methods. The rate at which susceptible individuals become infected clearly depends on the number of susceptible-infected partnerships in the population. If we set $[SS]$ to be the expected number of susceptible-susceptible partnerships in the population and $[SI]$ to be the expected number of susceptible-infected partnerships, we see that there is a flux from $[SS]$ to $[SI]$ resulting from susceptible individuals becoming infected from another partnership. Determining the rate of this flux is a central focus of pair approximation models.

We begin by considering the basic pairwise model and then investigate a more recent version which reduces the number of equations required. Before continuing, we make a note about our accounting. Each partnership may be thought of as ordered. Thus it makes sense to talk about the “first” or “focal” individual of a partnership. In our accounting, each partnership will appear twice. So if both partners are susceptible, this partnership will contribute 2 to $[SS]$, while if one is susceptible and the other infected it will contribute 1 to each of $[SI]$ and $[IS]$. Of course, we know $[SI] = [IS]$, so we do not need to calculate both, but we should be aware that the contribution to $[SS]$ is doubled.

2.2.1 The basic pairwise model

Our initial goal is to track individuals of degree k as they pass through different infection classes. We set $[S_k]$, $[I_k]$, and $[R_k]$ to be the expected number of susceptible, infected, and recovered individuals of degree k in the population. Because recovery occurs at rate γ , it is straightforward to see that the flux from $[I_k]$ to $[R_k]$ is $\gamma[I_k]$. The rate of infection follows easily: Each partnership between a susceptible and an infected individual transmits independently at rate β , so the flux from $[S_k]$ to $[I_k]$ is $\beta[S_k I]$ where $[S_k I]$ is the total number of partnerships between susceptible individuals of degree k and infected individuals. As t increases, $[S_k I]$ changes, so we need to be able to track its evolution.

This is the reason that the basic pairwise model [7] focuses on pairs or partnerships among individuals. We set $[S_k S_{k'}]$ to be the total number of partnerships between susceptible individuals of degrees k and k' . This is given by the sum over all susceptible individuals of degree k of the number of susceptible partners of degree k' . This definition gives an implicit direction, so that we can think of this as the total number of partnerships from susceptible individuals of degree k to susceptible individuals of degree k' . If $k = k'$, each partnership is counted twice.

We define $[S_k I_{k'}]$ and $[I_k S_{k'}]$ to correspond to the number of partnerships from susceptible degree k individuals to infected degree k' individuals and from infected degree k individuals to susceptible degree k' individuals respectively. Note that $[S_k I] = \sum_{k'} [S_k I_{k'}]$. We similarly define $[S_k R_{k'}]$, $[I_k I_{k'}]$, $[R_k S_{k'}]$, $[I_k R_{k'}]$, $[R_k I_{k'}]$, and $[R_k R_{k'}]$.

Putting this together, we get the flow diagrams in figure 3. We note that when a susceptible individual in a partnership of interest transitions to being infected, it may be because of a transmission from within that partnership or because of a transmission from outside the partnership. In figure 3 we denote transmissions from outside the partnership by dashed lines. Solid lines are used for transmissions within a partnership or a recovery of an infected individual within the partnership, events that do not depend on triples.

For each dashed line, the central individual of the relevant triple is susceptible. This means that if the triples closure holds, then we can reduce our triples into pairs. The resulting equations are then closed at

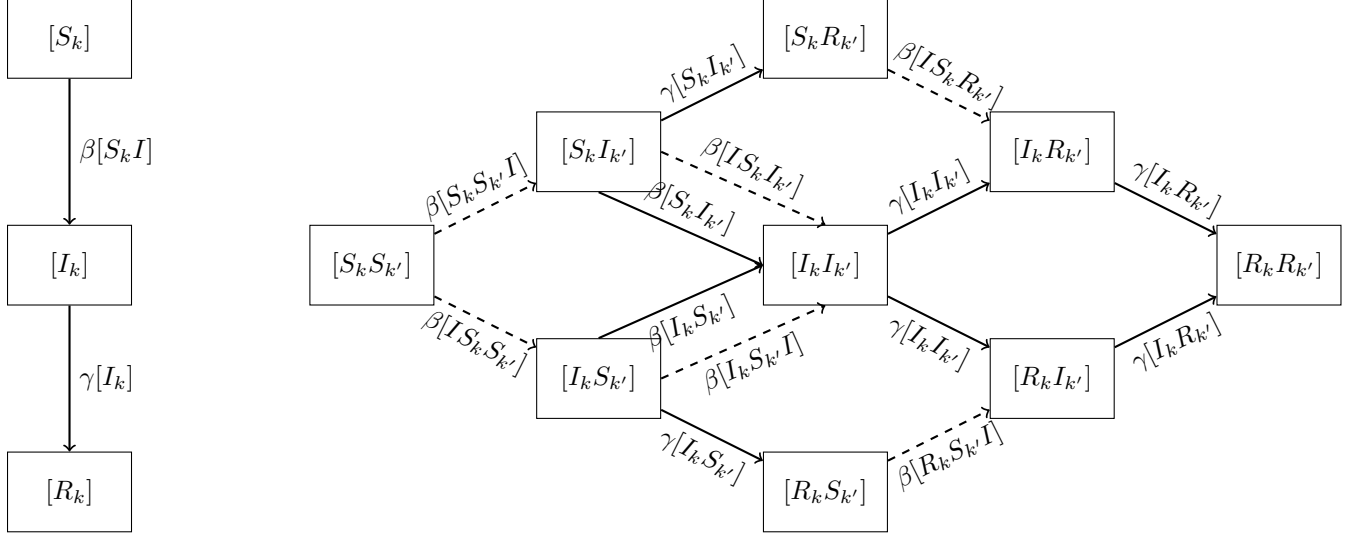


Figure 3: **The flow diagram underlying the basic pairwise model.** Dashed lines denote transitions that rely on infection coming from a source outside the partnership of interest. A similar cascade exists at the level of triples, with some transitions depending on paths of length 4 or on central individuals with three partners.

the pairs level:

$$[S] = \sum_k [S_k] \quad (1)$$

$$[I] = \sum_k [I_k] \quad (2)$$

$$[R] = \sum_k [R_k] = N - [I] - [S] \quad (3)$$

$$[IS_k] = [S_k I] = \sum_{k'} [S_k I_{k'}] \quad (4)$$

$$[\dot{S}_k] = -\beta[S_k I] \quad (5)$$

$$[\dot{I}_k] = \beta[S_k I] - \gamma[I_k] \quad (6)$$

$$[\dot{R}_k] = \gamma[I_k] \quad (7)$$

$$[S_k \dot{S}_{k'}] = -\beta \frac{k' - 1}{k'} \frac{[S_k S_{k'}][S_{k'} I]}{[S_{k'}]} - \beta \frac{k - 1}{k} \frac{[IS_k][S_k S_{k'}]}{[S_k]} \quad (8)$$

$$[S_k \dot{I}_{k'}] = \beta \frac{k' - 1}{k'} \frac{[S_k S_{k'}][S_{k'} I]}{[S_{k'}]} - \gamma[S_k I_{k'}] - \beta \frac{k - 1}{k} \frac{[IS_k][S_k I_{k'}]}{[S_k]} - \beta[S_k I_{k'}] \quad (9)$$

$$[S_k \dot{R}_{k'}] = \gamma[S_k I_{k'}] - \beta \frac{k - 1}{k} \frac{[IS_k][S_k R_{k'}]}{[S_k]} \quad (10)$$

If our goal is to just find $[S]$, $[I]$, and $[R]$, we do not require $[S_k R_{k'}]$, $[I_k]$, or $[R_k]$ directly, so we can avoid calculating them. They are included here for completeness. We can instead write $[I] = N - [S] - [R]$ and $[\dot{R}] = \gamma[I]$. So we can reduce the system to $2K^2 + K + 1$ differential equations where K is the total number of distinct degrees in the population. A particular weakness of these equations is that in many important limiting cases, K tends to ∞ .

It should be noted that if the population structure is not Configuration Model (for example, there may

be correlation in the degrees of partners), the triples closure may still hold. In such cases it may be possible to use a similar process to reduce the model. Depending on how the population structure deviates from Configuration Model, we may not be able to use the star closure or the pairs closure. So this approach may be more flexible than the others introduced below, but it requires more equations than some of the later models.

We note that all of these variables are numbers of individuals or numbers of pairs. Thus they are expected to be proportional to N , the population size. We may divide through by N , in which case S , I , and R become the proportion of individuals in each state. This may be a more natural system to use as we are interested in the limit of large populations, so by dividing by N we arrive at equations that do not change with population size. This is effectively the distinction between intensive and extensive quantities in physical systems. However, in the existing literature these equations are typically introduced in terms of absolute numbers rather than proportions.

2.2.2 A compact pairwise model

A recent paper [10] developed some pairwise equations with a significantly reduced dimension. It was derived for a multiple disease model, but we focus on the restriction to just a single disease.

We begin with the pairwise model using the triples closure, but we additionally make the pairs closure: We assume that if v is a partner of a susceptible individual u , then the probability v is susceptible or infected is independent of the degree of u (or other partners of u).

We define $\langle I \rangle$ to be the probability a partner of a susceptible individual u is infected. By the pairs closure assumption, this is independent of k . It follows that

$$\langle I \rangle = \frac{[SI]}{\sum_k k[S_k]}.$$

Since we know that $\sum_k k[S_k] = [SS] + [SI] + [SR]$, this becomes

$$\langle I \rangle = \frac{[SI]}{[SS] + [SI] + [SR]}.$$

The number of susceptible-susceptible edges between degree k and degree k' individuals is equal to the number of edges from susceptible individuals of degree k to other susceptible individuals times the probability that other susceptible individual has degree k' . That is

$$[S_k S_{k'}] = [SS_k] \frac{k'[S_{k'}]}{\sum_{k''} k''[S_{k''}]}.$$

This further leads us to

$$\frac{\sum_{k'} (k' - 1)[S_k S_{k'}]}{[S_k S]} = \frac{\sum_{k'} k'(k' - 1)[S_{k'}]}{\sum_{k''} k''[S_{k''}]}$$

which we define to be $\langle k \rangle_{\text{ex}}$. This is the average “excess degree” of a susceptible individual. That is, if we reach a susceptible individual by following a partnership this is the expected number of other partners that susceptible individual has. It appropriately accounts for the fact that the probability of reaching an individual along a partnership is proportional to its degree. We can simplify the denominator to $[SS] + [SI] + [SR]$.

Using these assumptions the flux $\beta[S_k I]$ from $[S_k]$ to $[I_k]$ can be represented as $\beta k \langle I \rangle [S_k]$.

$$\begin{aligned} \dot{[S_k]} &= -\beta k \langle I \rangle [S_k] \\ \dot{[I_k]} &= \beta k \langle I \rangle [I_k] - \gamma [I_k]. \end{aligned}$$

We no longer need $[S_k I]$ explicitly for our equations at the singles level. So at the pairs level we can sum equations (8)–(10) over k and k' . We find $\sum_{k,k'} (k - 1)[S_k S_{k'}][IS_k]/k[S_k] = \langle k \rangle_{\text{ex}} \langle I \rangle [SS]$. Similarly if we

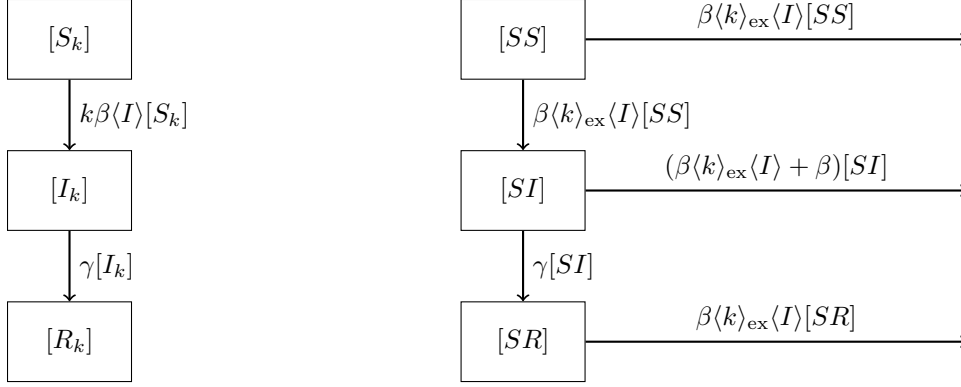


Figure 4: **The flow diagram for the reduced system of [10].** The $[SS]$, $[SI]$, and $[SR]$ compartments correspond to the sum of the $[S_k S_{k'}]$, $[S_k I_{k'}]$ and $[S_k R_{k'}]$ compartments of the basic pairwise model. The fluxes to the right go to compartments $[IS]$, $[II]$, and $[IR]$ which are not needed for our calculations.

switch k and k' in this equation, we get the same result. So we arrive at

$$\begin{aligned} \dot{[S]} &= -2\beta\langle k \rangle_{\text{ex}}\langle I \rangle [SS] \\ \dot{[SI]} &= \beta\langle k \rangle_{\text{ex}}\langle I \rangle [SS] - (\beta + \gamma)[SI] - \beta\langle k \rangle_{\text{ex}}\langle I \rangle [SI] \\ \dot{[SR]} &= \gamma[SI] - \beta\langle k \rangle_{\text{ex}}\langle I \rangle [SR]. \end{aligned}$$

The $[SR]$ equation can be kept if it simplifies calculations, or we can use $[SR] = (\sum_k k S_k) - [SI] - [SS]$.

The variables $[I_k]$, $[R_k]$ and $[SR]$ are not needed to find $[S]$, $[I]$, and $[R]$. If we do not include these, our final system is

$$[S] = \sum_k [S_k] \quad (11)$$

$$[I] = N - [S] - [R] \quad (12)$$

$$[\dot{R}] = \gamma[I] \quad (13)$$

$$[\dot{S}_k] = -\beta k \langle I \rangle [S_k] \quad (14)$$

$$[\dot{S}S] = -2\beta\langle k \rangle_{\text{ex}}\langle I \rangle [SS] \quad (15)$$

$$[\dot{SI}] = \beta\langle k \rangle_{\text{ex}}\langle I \rangle [SS] - (\beta + \gamma)[SI] - \beta\langle k \rangle_{\text{ex}}\langle I \rangle [SI] \quad (16)$$

$$\langle I \rangle = \frac{[SI]}{\sum_k k [S_k]} = \frac{[SI]}{[SS] + [SI] + [SR]} \quad (17)$$

$$\langle k \rangle_{\text{ex}} = \frac{\sum_k k(k-1)[S_k]}{\sum_k k [S_k]} = \frac{\sum_k k(k-1)[S_k]}{[SS] + [SI] + [SR]}. \quad (18)$$

The number of differential equations needed is reduced to $K + 3$. This model is represented by a reduced flow diagram, shown in figure 4.

Our derivation assumed that the pairs closure holds at all times. To be fully rigorous, we would additionally need to show that if the pairs closure holds at $t = 0$, then the basic pairwise model predicts that it continues to hold at all later time.

Similarly to the basic pairwise model, all of the bracketed quantities may be divided by the population size N so that the equations are in terms of “intensive” quantities.

2.3 Effective degree models

We now turn to “effective degree” models. We define an “inactive” partnership to be a partnership which we know will never transmit infection. We then stratify individuals by the number of active partners they have. How “omniscient” we assume we are affects how well we can identify which partnerships will never transmit infection. Depending on the information we choose to track in our model, we may know that an infected individual has an infected partner in which case we know that partnership can be treated as inactive, or we may simply know that a partner of an infected individual may be infected in which case we cannot discard the partnership and instead must track the probability the partner is infected. So we can arrive at different models depending on how we choose our variables.

We focus our attention on two models which have been studied already. By changing the definition of active partnerships other valid models could be derived. Using the pairs closure and a careful definition of active partnerships, we are able to further simplify these models and introduce a new simpler model.

2.3.1 The basic effective degree model

When we introduced the star closure, we used the variable $x_{s,i,r}$ to be the number of susceptible individuals with s susceptible, i infected, and r recovered partners. The star closure implies that the susceptible partners of one individual are indistinguishable *a priori* from the susceptible partners of another. Thus the degree of an individual does not matter, and so the future risk of a person with s susceptible partners and i infected partners is unaffected by the number of recovered partners.

So we can instead use $x_{s,i} = \sum_r x_{s,i,r}$ as our focal variable. This is the number of stars with a susceptible central individual and s susceptible and i infected peripheral individuals. We similarly introduce $y_{s,i}$ to correspond to stars with infected central individuals. These are the variables of [20]: We assume only a very basic amount of information about which partnerships we can discard, we can only identify those partnerships which contain (at least) one recovered individual. So active partnerships are defined to be those partnerships which do not have a recovered member. The star closure was not defined in terms of infected central individuals, and we shall see that in fact the $y_{s,i}$ variables can be eliminated from these equations.

Here we modify the approach of [20] by introducing the variables ξ and ζ : Given a susceptible individual u which is chosen by following a random edge, these are the expected number of infected partners of u that the edge followed to u came from a susceptible or infected individual respectively. When we eliminate $y_{s,i}$ from the system, ζ will no longer be needed. To calculate ξ , we notice that v is chosen by finding a susceptible individual u and then selecting a susceptible partner. So v is chosen with probability proportional to its number of susceptible partners. Accounting for this bias, we have

$$\xi = \frac{\sum_{k_e=1}^M \sum_{s+i=k_e} s i x_{s,i}}{\sum_{k_e=1}^M \sum_{s+i=k_e} s x_{s,i}}$$

where the summation over $s + i = k_e$ means the sum over all pairs s and i that sum to k_e . We similarly find

$$\zeta = \frac{\sum_{k_e=1}^M \sum_{s+i=k_e} i^2 x_{s,i}}{\sum_{k_e=1}^M \sum_{s+i=k_e} i x_{s,i}}.$$

These differ from G and H in [20] by a factor of β . The denominator of ζ appears different from the denominator of H in [20] but it is in fact the same¹. Our interpretation of ζ differs slightly from their interpretation of H because their interpretation requires a slightly stronger assumption than the star closure — they assume that the star closure applies when the central individual is infected. This assumption is consistent with the dynamics, but if the initial condition violates this assumption, this assumption will fail at later times. However, the failure is irrelevant for the purposes of determining the evolution of S , I , and R .

¹Here $\sum_{k_e} \sum_{s+i=k_e} i x_{s,i}$ counts the number of susceptible-infected partnerships from the perspective of susceptible individuals, while $\sum_{k_e} \sum_{s+i=k_e} s y_{s,i}$ in [20] counts the same thing from the perspective of infected individuals

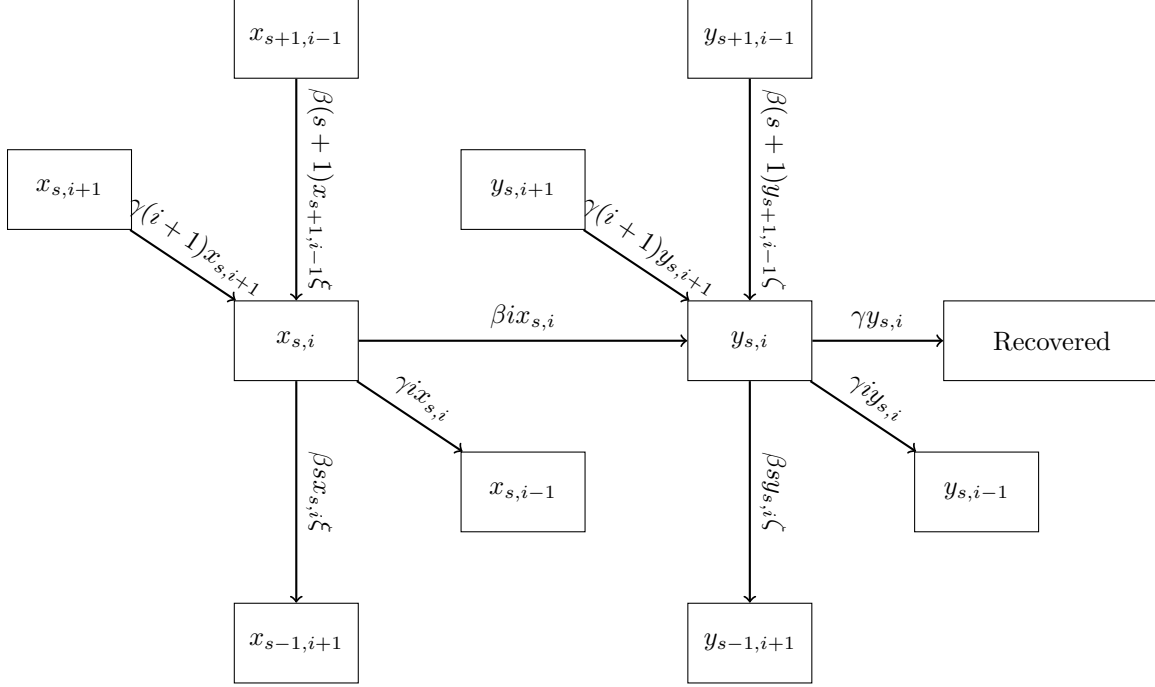


Figure 5: **The flow diagram underlying the effective degree model of [20].** We include just the fluxes involving the $x_{s,i}$ or $y_{s,i}$ compartments. Fluxes between other compartments exist but are not included (they can be deduced by changing s and i and using this figure). Given a susceptible individual u , if v is a susceptible partner of u , the variable ξ is the expected number of infected partners of v . Given an infected individual u , if v is a susceptible partner of u , then ζ is the expected number of infected partners of v (including u). We have made some small changes to the variables as used by [20]: Their $G = \beta\xi$ and their $H = \beta\zeta$.

Following figure 5, the final equations are

$$\dot{x}_{s,i} = -\beta i x_{s,i} + \gamma [(i+1)x_{s,i+1} - i x_{s,i}] + \beta \xi [(s+1)x_{s+1,i-1} - s x_{s,i}] \quad (19)$$

$$\dot{y}_{s,i} = \beta i x_{s,i} - \gamma y_{s,i} + \gamma [(i+1)y_{s,i+1} - i y_{s,i}] + \beta \zeta [(s+1)y_{s+1,i-1} - s y_{s,i}] \quad (20)$$

$$\xi = \frac{\sum_{k_e=1}^M \sum_{s+i=k_e} s i x_{s,i}}{\sum_{k_e=1}^M \sum_{s+i=k_e} s x_{s,i}} \quad (21)$$

$$\zeta = \frac{\sum_{k_e=1}^M \sum_{s+i=k_e} i^2 x_{s,i}}{\sum_{k_e=1}^M \sum_{s+i=k_e} i x_{s,i}} \quad (22)$$

$$S = \sum_{k_e=1}^M \sum_{s+i=k_e} x_{s,i} \quad (23)$$

$$I = \sum_{k_e=1}^M \sum_{s+i=k_e} y_{s,i} \quad (24)$$

$$R = N - S - I. \quad (25)$$

This is a system with $2(M+1)^2$ differential equations where M is the maximum degree. Careful consideration can remove about half of these equations since the $\dot{x}_{s,i}$ equations do not depend on the $y_{s,i}$ terms. So we can sum the $\dot{y}_{s,i}$ equations to arrive at a single equation for $\dot{I} = \beta \sum_{k_e=1}^M \sum_{s+i=k_e} i x_{s,i} - \gamma I$. So we can still calculate S , I , and R if we reduce this system to $(M+1)^2 + 1$ differential equations. The basic effective

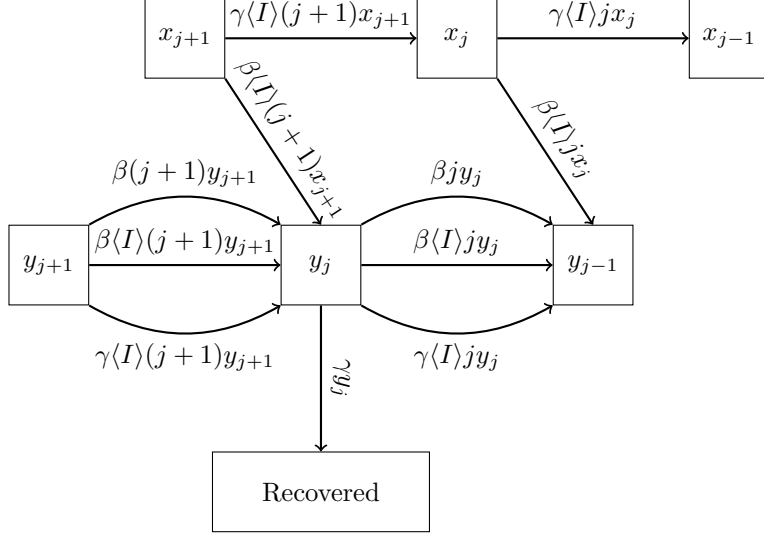


Figure 6: **The flow diagram underlying the model of [2].** Only the fluxes into and out of y_j and x_j are included. Fluxes between other compartments exist, but are not included. An active edge is eliminated if the partner recovers or if it transmits infection in either direction. The quantity $\langle I \rangle$ represents the probability an active edge joins an individual with an infected partner.

degree model was developed by [20] in large part to study SIS disease, and this simplification would not be appropriate in the SIS context.

As before, we may switch to intensive quantities by dividing the x and y variables by N .

2.3.2 The compact effective degree model

A related model [2], uses a different classification to identify “active partnerships”. As above, the model ignores partnerships involving recovered individuals. However, we also assume that we can observe when a transmission occurs in a partnership. So we can observe if a partnership was the original source of infection for either individual and we can observe those partnerships in which one infected individual “transmits” to its partner but fails to cause infection because the partner is already infected. If a partnership has not transmitted infection and neither individual is recovered, then it is an active partnership.

We now add the pairs closure assumption: Given a susceptible individual u the probability that an active partner v is susceptible or infected is independent of any information we have about u or the status of any other partners of u . As a result, the probability that a susceptible individual u has s active susceptible partners and i active infected partners can be deduced by knowing the probability of having $s + i$ active partners and the independent probability that a partner is infected. Thus rather than having $x_{s,i}$ we stratify into groups x_j where j is the number of active partners of a susceptible individual. We refer to this model as “compact” because we have just a single subscript to track. We note that the “reduced” model introduced below is of similar complexity to this one.

To derive the equations of [2] we additionally assume that the probability an active partner of an infected individual u is infected is the same as it would be if u were susceptible. That is, we make the strong pair closure assumption.

We can calculate $\langle I \rangle$, the probability that an active partner is infected. A little consideration will show that this is $\langle I \rangle = \sum_j j y_j / \sum_j j (y_j + x_j)$. Thus on average for susceptible individuals with j active partnerships, we conclude that the rate of becoming infected is $\beta j \langle I \rangle$. Similarly the rate at which such an individual loses an active partner because the partner recovers is $\gamma j \langle I \rangle$. The flow diagram in figure 6 shows

the relevant transitions. The resulting equations are

$$\dot{x}_j = \gamma \langle I \rangle ((j+1)x_{j+1} - jx_j) - \beta \langle I \rangle jx_j \quad (26)$$

$$\dot{y}_j = (\beta + \beta \langle I \rangle + \gamma \langle I \rangle) ((j+1)y_{j+1} - jy_j) + \beta \langle I \rangle (j+1)x_{j+1} - \gamma y_j \quad (27)$$

$$\langle I \rangle = \frac{\sum_j j y_j}{\sum_j j (x_j + y_j)} \quad (28)$$

$$S = \sum_j x_j \quad (29)$$

$$I = \sum_j y_j \quad (30)$$

$$R = N - S - I. \quad (31)$$

This system has significantly fewer equations than the basic effective degree model, just $2(M+1)$ differential equations where M is the maximum degree. These are the equations of [2]. We can eliminate y_j from this system by noting that $\langle I \rangle$ depends on y_j only through the sum $\sum_j j y_j$. If we define $\lambda = \sum_j j y_j$ we have $\langle I \rangle = \lambda / (\lambda + \sum_j j x_j)$ and we find

$$\dot{\lambda} = \beta \langle I \rangle \left(\sum_j j(j+1)x_{j+1} \right) - [\beta + \gamma + (\beta + \gamma) \langle I \rangle] \lambda.$$

So if we know the initial value of λ , we do not need to track the change in each individual y_j . This substantially reduces the number of differential equations. However, because we no longer have an individual equation for each y_j , we no longer have an equation for I . We must add one differential equation back. The simplest form is to take $S = \sum_j x_j$, $I = N - S - R$, and $\dot{R} = \gamma I$. Thus we have $M+3$ differential equations.

The role of λ is to count the number of active partnerships that involve infected individuals (counting double if both are infected). The role of $\sum_j j x_j$ is to count the number of active partnerships that involve susceptible individuals (also counting double if both are susceptible). So $\langle I \rangle = \lambda / (\lambda + \sum_j j x_j)$ measures the probability that an active partner is infected. Thus the probability that a susceptible individual's active partner is infected is $\langle I \rangle$.

2.3.3 Reduced Effective Degree Model

With a small modification of our definition of active partnerships, we can eliminate the need to track the number of partners infected individuals have altogether. We add to our definition the statement that an active partnership must have at least one susceptible individual. With this definition, we will not need to track partnerships between infected individuals. This new approach will not require the strong version of the pairs closure.

We define ν to be the number of active partnerships between susceptible and infected individuals. Using this definition, $\langle I \rangle = \nu / \sum_j j x_j$. There are several ways that ν can change. Infection of a partner in a susceptible-susceptible pair increases ν . The contribution to $\dot{\nu}$ from this is $\sum_j \beta \langle I \rangle j x_j (j-1)(1-\langle I \rangle)$. The $\beta \langle I \rangle j x_j$ gives the rate at which x_j individuals are infected, and $(j-1)(1-\langle I \rangle)$ gives the expected number of new susceptible-infected partnerships resulting. The value of ν decreases when the infected partner within a susceptible-infected partnership recovers or transmits to the susceptible partner. This contributes $-(\beta + \gamma)\nu$ to $\dot{\nu}$. Finally ν decreases when the susceptible member of a susceptible-infected partnership is infected from outside the partnership. This contributes $-\beta \langle I \rangle^2 \sum_j j(j-1)x_j$ to $\dot{\nu}$. We are left

$$\dot{\nu} = -(\beta + \gamma)\nu + \beta \langle I \rangle (1 - 2\langle I \rangle) \sum_j j(j-1)x_j.$$

Note that the derivation of ν does not rely on any independence assumption of partners of infected individuals. This yields the new system

$$\dot{x}_j = \gamma \langle I \rangle ((j+1)x_{j+1} - jx_j) - \beta \langle I \rangle jx_j \quad (32)$$

$$\dot{\nu} = -(\beta + \gamma)\nu + \beta \langle I \rangle (1 - 2\langle I \rangle) \sum_j j(j-1)x_j \quad (33)$$

$$\langle I \rangle = \frac{\nu}{\sum_j jx_j} \quad (34)$$

$$S = \sum_j x_j \quad (35)$$

$$I = N - S - R \quad (36)$$

$$\dot{R} = \gamma I. \quad (37)$$

We again have $M + 3$ equations, but we do not need to assume anything about the partners of infected individuals. The derivation of this requires only the pairs closure.

Again, we may switch to intensive quantities by dividing all x and variables and ν by N .

This system can be derived by adding the pairs closure to the basic effective degree model. We demonstrate this in the appendix.

2.4 Edge-based compartmental modeling

We finally turn to a third approach which has been called the ‘‘Edge-based compartmental modeling’’ (EBCM) approach. This approach relies on creating compartments of partnerships (or edges) rather than compartments of individuals. It is also sometimes referred to as a ‘‘pgf’’ approach because the equations contain probability generating functions. The ‘‘pgf’’ terminology may be more descriptive because the pairwise equations also are based on compartments of edges. Nevertheless, we will stick to the ‘‘EBCM’’ terminology. The method is described in [31]. Some of the definitions of [31] may at first seem unusual, but this is done to simplify the resulting mathematics by eliminating the need to use conditional probability arguments. We begin by discussing the variables used. Then we derive the governing equations.

2.4.1 The EBCM variables

Under the assumption that the population-scale epidemic dynamics are deterministic, the probability a random individual is in a given state will match the proportion of the population in that state. Thus our derivation will focus on calculating the probability a random individual is in each state.

The pairs closure guarantees that as long as an individual u is susceptible, we can treat its partners as independent. Our goal is to calculate the probability that if v is a partner of u , then at time t no individual x reachable from u through v has caused a chain of transmissions that reach u . This is complicated somewhat because if u is infected from another source, it can infect v , which short-circuits the potential chain of transmissions from x . We can get around this by conditioning on the assumption u has not been infected from another source, but the arguments are quickly quite difficult. Instead we modify u to prevent u from transmitting to its partners if infected. This prevents the transmission chains to u from interacting with one another. It does not alter the probability that u has a given status because it only changes things once u is infected, and at that point the recovery time of u is unaffected by any dynamics extrinsic to u . This is discussed in more detail in A.2.

When u is a randomly chosen individual which is prevented from causing infections, we refer to u as a *test individual*. We now are able to define our variables. Our central variable is θ , the probability that a partner v of a test individual u has not yet transmitted to u . We break θ up into three variables $\theta = \phi_S + \phi_I + \phi_R$. Each is the probability that v has not yet transmitted to u and is in the given state.

We can avoid the modification done in defining the test individual by using an alternate definition. We choose u uniformly at random from the population and v as a uniformly chosen random partner of u . Then

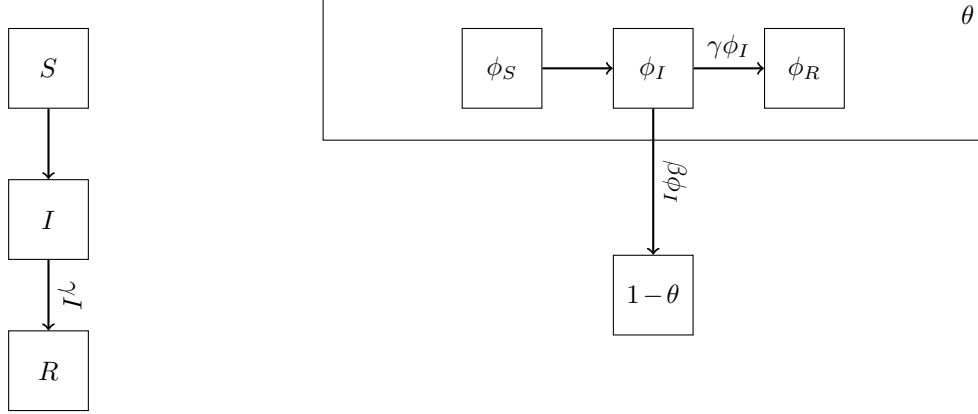


Figure 7: **The flow diagram for the basic Edge-Based Compartmental Model.** We do not calculate the flux from ϕ_S to ϕ_I or from S to I directly because we calculate ϕ_S and S directly. The relevant fluxes can be found by differentiating ϕ_S or S . Using the relative fluxes from ϕ_I , we can calculate ϕ_R in terms of θ .

θ is the probability that no transmission has occurred between the two given that either no transmission has happened or if transmission has happened, the first transmission was v to u . Although this alternative definition would lead to the same equations, for consistency with prior work, we will use the test individual assumption. So u will be prevented from transmitting.

2.4.2 Equation derivation

In deriving the equations, we use S , I , and R to be the number of susceptible, infected, or recovered individuals to be consistent with the models discussed previously. In other work with the EBCM model, these are typically taken to be the proportions.

Let us now consider a randomly chosen individual u in the population. We define $S_k(0)$ to be the number of individuals which are initially susceptible and have degree k . We define $\theta(t)$ to be the probability that an edge to a test individual u has not yet transmitted to u by time t assuming that u was initially susceptible [so $\theta(0) = 1$]. We assume that θ is independent of the degree of u . Averaging over all k , the probability that u is susceptible at time t is thus $\sum_k \frac{S_k(0)}{N} \theta(t)^k$. We define this function to be $\psi(\theta)$. The number of infections is $S(t) = N\psi(\theta)$. To find $R(t)$ and $I(t)$, the number that are recovered or infected respectively, at time t we take $\dot{R} = \gamma I$, and $I = N - S - R$. In the limit that $S_k(0)$ can be approximated as $NP(k)$ (that is, a negligible proportion of each degree initially infected), the function $\psi(\theta)$ is the probability generating function of the degree distribution.

To complete our system, we need to know θ . This is the probability that a partner v of u has not yet transmitted to u given that u was initially susceptible. It is straightforward to see that $\dot{\theta} = -\beta\phi_I$ because θ is decreased by transmissions, transmissions happen along edges at rate β , and ϕ_I is the probability transmission has not yet happened and v is infected. Our goal will be to find ϕ_I in terms of θ to reduce our system to just this single differential equation.

We begin by calculating ϕ_S in terms of θ . If v was initially susceptible, the probability it has degree k is $kS_k(0)/\sum_k kS_k(0)$. Assuming it was initially susceptible with degree k , the probability that it is still susceptible is θ^{k-1} (each partnership other than the one with u has a chance to transmit to v)². So the probability that v is susceptible at time t is $\phi_S(0) \sum_k kS_k(0)\theta^{k-1}/\sum_k kS_k(0) = \phi_S(0)\psi'(\theta)/\psi'(1)$.

²To help make the derivation clearer, we can make a test-individual like assumption so that v can only transmit to u and no other partners.

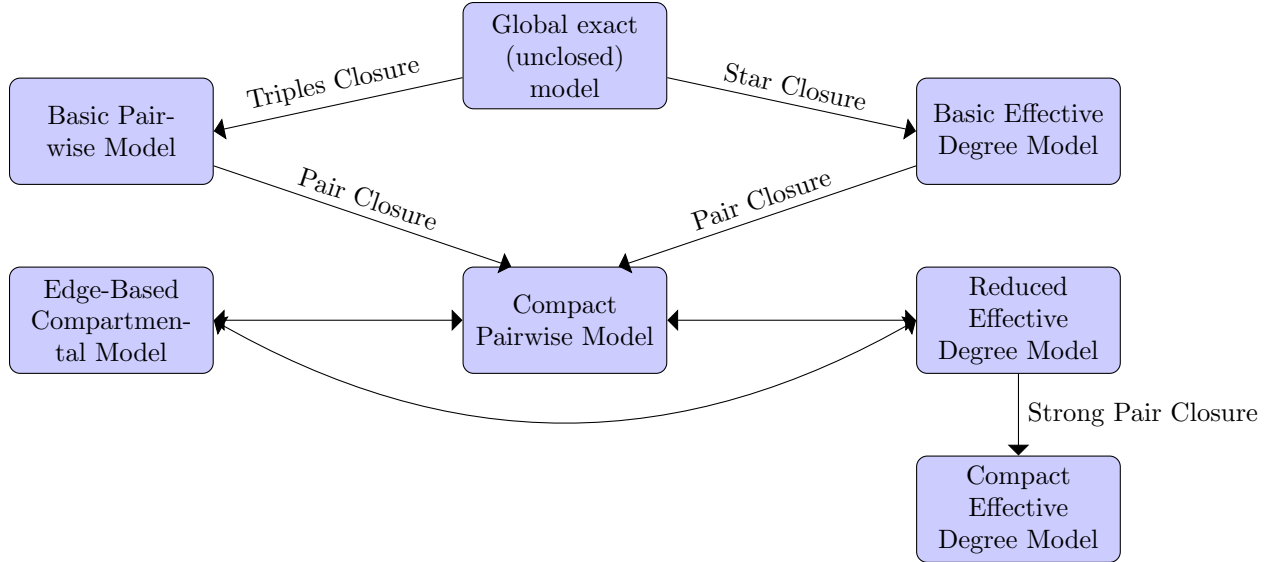


Figure 8: **The hierarchy of models.** Labeled edges denote the additional assumptions required to find the mathematically simpler model. Unlabeled (bidirectional) edges imply that the two models are equivalent and each can be derived from the other. The unclosed exact model will reduce to either the basic effective degree model of [20] or the basic pairwise model [7] depending on the closure used. Each of these then reduces to the compact pairwise model (as modified from [2]). The compact pairwise model, the reduced effective degree Model, and the Edge-based Compartmental model [31] are all equivalent. Using the strong version of the pair closure, we can derive the compact effective degree model which also tracks information about infected individuals and their partners.

We now find ϕ_R . The rate at which an infected neighbor recovers is γ . The rate at which an infected neighbor transmits to u is β . Thus, we find that $\dot{\phi}_R = -\gamma\dot{\theta}/\beta$. Taking $\phi_R(0)$ as given and $\theta(0) = 1$, we conclude $\phi_R(t) = \phi_R(0) + \gamma(1 - \theta)/\beta$.

We finally conclude that $\phi_I = \theta - \phi_S(0)\psi'(\theta)/\psi'(1) - \gamma(1 - \theta)/\beta - \phi_R(0)$. Thus we arrive at

$$\dot{\theta} = -\beta\theta + \beta\phi_S(0)\frac{\psi'(\theta)}{\psi'(1)} + \gamma(1 - \theta) + \beta\phi_R(0) \quad (38)$$

$$S = N\psi(\theta) \quad (39)$$

$$I = N - S - R \quad (40)$$

$$\dot{R} = \gamma I. \quad (41)$$

The single differential equation drives all the dynamics, but if we want S , I , and R as well, we need to include an additional equation. So we have just two differential equations.

Changing to intensive rather than extensive variables has no impact on the equation for θ and simply divides S , I , and R by N .

3 Model Selection

3.1 Relationship of models

The models are based on different assumptions about the independence of partners. Although the assumptions are formally different, the distinction is in a sense unnatural. If the initial infections are randomly

| Model | Minimal number of Differential Equations | Analytic final size calc? | Early growth calc? |
|--------------------------------|--|---------------------------|--|
| Basic pairwise model | $2K^2 + K + 1$ | No | $\mathcal{O}(K^2 \times K^2)$ eigenvalue problem |
| Compact pairwise model | $K + 3$ | No | Analytic calculation |
| Basic effective degree model | $(M + 1)^2 + 1$ | No | $\mathcal{O}(M^2 \times M^2)$ eigenvalue problem |
| Compact effective degree model | $M + 3$ | No | $\mathcal{O}(M \times M)$ eigenvalue problem. |
| Reduced effective degree model | $M + 3$ | No | Analytic calculation. |
| Edge Based Compartmental model | 2 | Yes | Analytic calculation. |

Table 1: **Comparison of the models.** The number of equations needed and computational/analytic tractability for a population having K distinct degrees with maximum degree $M \geq K$.

selected, all the assumptions are satisfied, and the models give identical predictions. If the disease is introduced as a few infections, and allowed to spread until enough infections have occurred that the dynamics are deterministic, then all the assumptions will be satisfied, and the models give identical predictions (technically it is possible that stochastic effects cause one assumption to be valid but not another, but this will be a rare event).

In order for the assumptions of any of the models to be violated, this must be built into the initial condition. Somehow the initial infections must be chosen based on who their neighbors are. Even if the initial condition fails to satisfy assumptions of some of the models, as time progresses information about the initial condition is lost. As the disease spreads, the violated assumptions may become valid (at least approximately) and the corresponding model is accurate.

Formally speaking, some of the models make stronger assumptions than others, and we prove in the appendix that it is possible to derive these models from the models having weaker assumptions. In the case in which we if we take a random susceptible individual u we find that the statuses of partners of u are independent of the degree of u and the status of one another, then we find that all of the models are appropriate. We can reduce all of the models to the EBCM model, having just a single ODE driving the dynamics.

We have discussed six distinct models of epidemic spread in static configuration model networks. These models have varying mathematical and conceptual complexity. A natural question is which model is appropriate for a given application.

3.2 Model robustness to assumptions

All of these models make some form of simplifying assumption (a closure) in the derivation. At the heart of each of the assumptions is some sort of independence assumption. Each assumption can be shown to be valid if there is no information content in certain scales. Since the epidemic is a stochastic process, whatever information may be imposed by the initial conditions is lost over time. We anticipate that all models should behave similarly at later time, and should fit observed dynamics closely.

3.3 Robustness to Configuration Model assumptions

These models have been derived by assuming that the population is a static Configuration Model network. This assumption is unrealistic: individuals change partners, and there may be correlations between the degrees of partners.

3.3.1 Degree correlations

If we allow for correlations to exist between the degrees of individuals, then the effective degree approaches will have to keep extra information. It is not enough to know how many active edges an individual has, we need to know how many partners the individual began with in order to know the risk from those active edges. This significantly increases the dimensionality of these models.

The basic pairwise model approach remains valid if degree correlations exist. The triples closure can hold even if the distribution of partners' degrees depends on an individual's degree. If it holds, then the approach used to derive the basic pairwise model will yield a usable system of equations [7].

The simplifications that go into deriving the compact pairwise model from the basic pairwise model fail if an individual's degree affects its choice of partner. So we will not be able to use this model (though similar simplifications may be possible, with $\langle I \rangle$ becoming a function of degree).

The Edge-Based Compartmental Model presented here was derived assuming that θ , the probability a partner of individual u has not transmitted to it, is independent of the degree of u . Clearly if there are degree correlations, this will not be true. However, depending on how those correlations occur, a modification of the EBCM approach may work. Details of how this can be done are in [32]. It requires allowing θ to be a function of k as well as t . So rather than a single equation for θ , we find K equations where K is the number of distinct degrees.

3.3.2 Dynamic networks

There are multiple ways we can include partner turnover in the model. One of the most obvious is to allow an individual's partnerships to end and be replaced by new partnerships. Another is to allow individuals to have intrinsic rates of creating new partnerships while they end existing partnerships independently. Other models are clearly possible.

Again, the effective degree models would require a significant increase in dimension to account for partnership turnover. We would need to track the number of inactive edges in some way.

The Edge-Based Compartmental Modeling approach is well-suited for many dynamic networks. Examples were developed in [31], and in [33] it was shown that a hierarchy of models could be constructed based on the assumptions about partner turnover. It was shown that a widely-used model [21, 22, 35, 40] that assumes that at every moment partnerships are re-selected randomly arises as a special case of this hierarchy, and indeed the standard mass action model can be recovered in appropriate limits. One particular case that should be noted is that it is possible to model some forms of "serosorting" where individuals actively select partners of similar infection status.

Depending on how the partnership turnover is modeled, the compact pairwise model may be robust enough to accommodate it. The EBCM approach for dynamic networks requires introducing additional variables that are equivalent to $[SS]$, $[SI]$, $[SR]$ and related quantities. So we hypothesize that an attempt to adapt the compact pairwise model to a dynamic network is likely to lead to the same equations found by the EBCM approach.

It is relatively simple to adapt the basic pairwise model to most dynamic networks considered in the generalizations of the EBCM model of [31].

3.4 Final size relations and early growth

3.4.1 Early growth

One of the most important questions to answer about a disease is whether it is capable of causing an epidemic in a given population. To answer this, we typically look at R_0 , the average number of infections caused by an early infectious individual, and try to find conditions under which $R_0 < 1$. However, for models which calculate the dynamic growth of an epidemic, it is difficult to extract R_0 from the model. Instead, we look at an equivalent measure, the early growth rate.

We assume the epidemic is initialized with an infinitesimally small proportion infected and grows proportionally to e^{rt} , and try to calculate the value of r . If $r > 0$, epidemics are possible. Otherwise they are not.

For the basic pairwise model, we take the $[S_k I_{k'}]$ equations from equations (1)–(10). Because we assume the amount of infection is small, the $[IS_k][S_k I_{k'}]$ term is negligibly small. The other terms however are not negligible. The resulting system yields a $K^2 \times K^2$ eigenvalue problem whose dominant eigenvalue determines the stability of the system.

For the compact pairwise model, equations (11)–(18), we can note that so long as the amount of infection is very small, the denominator of $\langle I \rangle$ may be treated as constant, so $\langle I \rangle$ is proportional to $[SI]$. Substituting this into the equation for $[SI]$, we discard the $\beta \langle k \rangle_{\text{ex}} \langle I \rangle [SI]$ term. Assuming that all but an infinitesimal proportion of the population is susceptible (so $[SS] \langle I \rangle = [SI]$), we arrive at $[\dot{SI}] = \beta \langle k \rangle_{\text{ex}}(0) [SI] - (\beta + \gamma) [SI]$, and so we conclude that

$$r = \beta \langle k \rangle_{\text{ex}}(0) - (\beta + \gamma).$$

For our first effective degree model, equations (19)–(25), the original paper [20] showed that the early growth rate could be derived from a $(M + 1)^2 - 1 \times (M + 1)^2 - 1$ eigenvalue problem.

For the compact effective degree model, equations (26)–(31), we can convert the y_j equations into an $M \times M$ eigenvalue problem (neglecting y_0). If we use the reduced effective degree model equations (32)–(37), we get

$$\begin{aligned} \dot{\nu} &= -(\beta + \gamma)\nu + \beta \langle I \rangle (1 - 2\langle I \rangle) \sum_j j(j-1)x_j \\ &= -(\beta + \gamma)\nu + \beta \frac{\nu}{\sum_j jx_j} (1 - 2\langle I \rangle) \sum_j j(j-1)x_j \\ &= -(\beta + \gamma)\nu + \beta (1 - 2\langle I \rangle) \nu \frac{\sum_j j(j-1)x_j}{\sum_j jx_j}. \end{aligned}$$

Assuming ν and $\langle I \rangle$ are both small, we have at leading order

$$\dot{\nu} = \left(-(\beta + \gamma) + \beta \frac{\sum_j j(j-1)x_j}{\sum_j jx_j} \right) \nu.$$

At $t = 0$, the value of $\sum_j j(j-1)x_j / \sum_j jx_j$ corresponds to $\langle k \rangle_{\text{ex}}(0)$, so the resulting growth rate corresponds to the previous value.

Finally, for the EBCM model, we set $\theta = 1 - \epsilon e^{rt}$ and solve for r . The equation for θ becomes

$$-r\epsilon e^{rt} = -\beta + \beta\epsilon e^{rt} + \beta \frac{\psi'(1) - \epsilon e^{rt}\psi''(1)}{\psi'(1)} + \gamma\epsilon e^{rt}.$$

After some rearranging,

$$r = \beta \frac{\psi''(1)}{\psi'(1)} - (\beta + \gamma).$$

We note that $\langle k \rangle_{\text{ex}}(0) = \psi''(1)/\psi'(1)$, so the analytic formulae we can derive are identical.

3.4.2 Final size

To search for a final size relation, we set all derivatives to 0 and look to see if we can find an equation for the final size. The only case in which a clear solution exists is the EBCM model. We can trivially arrive at an equation which gives the fixed point of θ . Using this fixed point, we immediately have $S = \psi(\theta)$, and taking $I = 0$, we conclude that $R = 1 - \psi(\theta)$.

4 Current Challenges

There are a number of questions that remain unanswered for epidemic spread in networks. Two that appear both important and difficult are the spread of disease through non-Configuration Model networks (in particular clustered networks) and the spread of SIS disease, that is diseases in which individuals return to a susceptible state. In both cases the challenge is similar: The independence assumptions made in our earlier models are not valid.

4.1 Clustered networks

Social contact networks violate the assumptions of a Configuration Model networks in a number of ways. One of the clearest violations which is particularly important for the modeling of infectious disease spread is that social contact networks often have clustering: The partners of an individual are likely to be partners of one another.

As a consequence, all of the closure approximations applied above fail. There have been a number of attempts to study disease spread in clustered networks, but most rely on some sort of approximation [23, 24, 28, 44, 3, 6, 8]. A select few are able to avoid such approximation, but only by defining their way out of it: If we restrict our attention to clustered networks which contain very specific motifs, then we can make analytic progress [27, 38, 48, 14]. However, these networks have very specific restrictions. Despite the specific structure, the number of equations in the corresponding ODE systems increases and it becomes difficult to extract analytic results from this.

We foresee that closures at higher structure levels are possible [12] and that the appropriate closures must preserve the structure of cycles within the population. However, this requires knowledge of the motif structure, and may require specialized generation algorithms.

It is possible to generate clustered networks with identical cluster “density” but very different large scale structure, and hence very different epidemic dynamics [9]. Thus clustering alone is not a sufficient metric to describe the network structure. Different motif types (*e.g.*, full connected triangles, squares, and fully connected squares with a missing diagonal) may lead to similar levels of clustering but different overall epidemic dynamics.

Our goal of course is a model which can take an arbitrary social network and accurately predict the dynamics of an epidemic spreading through the network. However, in the immediate future, we foresee that any approximate closure models will generally be developed on a case-by-case basis, lacking the generality of SIR models on Configuration Model networks.

4.2 SIS models

The key simplification underlying the EBCM model was that in calculating the status of individual u , we can ignore the possibility of transmissions from u to its partners. For an SIS epidemic, this assumption fails. In a Configuration Model SIR epidemic, the partners of an individual u are independent until u becomes infected. Then the partners are no longer independent; however, they can no longer influence the status of u because it is infected. So that dependence of partners is irrelevant. In an SIS epidemic, once infected, u can return to a susceptible state and be reinfected. The dependence of partners of u is no longer negligible.

Indeed, the fact that u can alter its future infection probability appears to underlie some initially surprising results [4]. It has been proven that certain networks are able to sustain an epidemic even though simple mean-field models suggested they could not. The apparent reason is that high degree individuals maintain

islands of infection about them by being continually reinfected (and occasionally managing to have infection spread along paths to other high degree individuals).

Our closure assumptions fail once an individual has an opportunity to infect its neighbors, so we must adapt our approaches. One of our challenges in modeling SIS disease is to identify appropriate closures which satisfy the dynamics. It should be noted that although the first effective degree model, equations (19)–(25), does not compare particularly well to the other models, this was in many ways an unfair comparison. It is a modification of a model initially developed for application to SIS epidemics. For SIS epidemics it is likely to outperform the more specialized SIR models because the star closure is expected to be more accurate than the other closures in the SIS case.

5 Discussion

There are multiple approaches used to model the dynamic spread of SIR epidemics in networks. These approaches have been developed under the assumption of a Configuration Model network. They perform well and give identical results under reasonable initial conditions. We have identified the precise conditions under which the models are equivalent. The models have varying complexity, with the simplest analytic model being the Edge-based Compartmental Model of [31, 29]

Some of the approaches can be adapted to population which are assortatively or disassortatively mixed. Other features such as dynamic partnerships or link weights can also be accounted for by some of these approaches.

There are two important restrictions to the approaches developed so far. These models all make some degree of assumption of independence of neighbors of a susceptible individual u . In a network with short cycles, this assumption breaks down. Similarly for an SIS disease, infection could travel from one partner of u to another through u . Even once u returns to susceptible, we can no longer treat the partners as truly independent. We can find very special classes of clustered networks for which our SIR models can be adapted (by assuming very special motifs exist), but the number of equations grows rapidly, and the approach does not appear to be generally applicable. For SIS epidemics, there does not appear to be any analytic exact model, except in the limit of partnerships having negligible duration.

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A Further EBCM details

The EBCM model begins with the pairs closure. We make an observation that if the dynamics are deterministic, then given the initial conditions, the probability a random individual is susceptible, infected, or recovered is equal to the proportion of the population in the corresponding state. This allows us to switch from the question of what proportion of the population is in a given state to an equivalent question of what is the probability an individual is in a given state. This new, equivalent question is more amenable to probabilistic arguments.

By noting that the probability an individual is in a given state is not affected by whether it transmits to its partners once infected, we can shift to yet another equivalent question which allows us to assume the partners of the random individual are independent. This is the key conceptual step in the derivation.

A.1 Variable choice

In this section we motivate the variable choice used in the EBCM approach. We begin by following a few steps from an early version of the EBCM approach [47]. When we consider a susceptible individual u with k partners, the probability p_I that one partner is infected (conditional on u being susceptible) is independent of any other partners and is independent of k . If β is the transmission rate and $S_k(t)$ is the number of susceptibles of degree k , we conclude that $\dot{S}_k = -\beta k p_I$. From this, we can conclude that $S_k = S_k(0)e^{k(-\beta \int_0^t p_I(\tau) d\tau)}$. Setting $\theta(t) = e^{-\beta \int_0^t p_I(\tau) d\tau}$, we conclude that $S_k(t) = S_k(0)\theta(t)^k$. Then $S(t) = \sum_k S_k(0)\theta(t)^k$. The variable θ can be roughly interpreted as the probability that a partner v of u has not transmitted to u . Taking the limit where $S_k(0) \approx NP(k)$, we arrive at $S(t) = N\psi(\theta(t))$ where $\psi(\theta) = \sum_k P(k)\theta^k$ is the probability generating function of the degree distribution.

The fact that $S_k(t)$ is proportional to $\theta(t)^k$ suggests that θ is somehow a measure of the independent probability that one partner of u has not yet transmitted to u . However, it is not immediately obvious how to precisely define the relevant quantity that is θ . If we think of “transmission” as being the first time that u receives an infectious dose, then the probabilities are dependent. Once one individual has transmitted, the others cannot — indeed the definition of p_I used above becomes meaningless as soon as a transmission has occurred. There are multiple ways to get around this. In [47], the approach was to take θ to be the probability an initially susceptible degree 1 individual is still susceptible, and then show that $S_k(t)$ is proportional to $\theta(t)^k$. Then it is possible to derive equations for p_I with (considerable) effort. An alternate approach introduced by [29] and extended in [31] is the one we will use here.

We first define “transmission” to be as we did in the compact effective degree model. An infected individual can still receive a transmission from another infected individual. However, it has no effect in terms of infection. This definition is not enough to create the independence of different partners that we need to proceed. The possibility exists that w will transmit to u which will transmit to v which will transmit back to u . So transmission from v to u is more likely if there has been transmission from w to u . What we actually want is to measure the probability that there is at least one individual which is reachable from u through v that begins infected and sparks a chain of transmissions through v that would reach u by time t . Having a w to u to v back to u chain would short circuit this. So we need to somehow condition on not having this short circuit. There is a lot of conditional probability in the resulting definition, so it is cumbersome to work with. We can eliminate the need for conditional probability arguments by modifying u at the outset. The individual u can receive transmissions, but cannot transmit to its partners. This eliminates the w to u to v chain without affecting the ability of infection to reach u through v , and θ is simply the probability that v has not yet transmitted to u .

Putting these arguments together, we define a *test individual* u to be a randomly chosen individual in the population which is prevented from transmitting to its partners. Then $\theta(t)$ is the probability that a partner of u has not yet transmitted to u .

When u is taken to be a test individual, the definition of p_I above becomes the probability that the partner v is infected given that it has not yet transmitted to u . In other words p_I is the probability that v is infected and has not transmitted given that it has not transmitted, and θ is the probability v has not transmitted. Then p_I and θ are both probabilities, but they are conditional on different events. This makes it difficult to work with these variables. Effectively, p_I is the ratio of two quantities, both of which are changing in time and so there is an implicit quotient rule hiding in the derivation of its derivative — this is a challenge. It is more natural to use $\phi_I = \theta p_I$ to be the probability that v is infected and has not transmitted to u . We similarly define ϕ_S to be the probability that v is susceptible (and obviously has not transmitted to u) and ϕ_R to be the probability that v is recovered and did not transmit to u . With this $\theta = \phi_S + \phi_I + \phi_R$.

A.2 Impact of preventing the test individual from transmitting

One final concern may arise because modifying u to prevent it from causing infection alters the dynamics of the epidemic. Some individuals that would otherwise get infected may now remain susceptible, while others simply have their infection delayed. We present two arguments for why this is not a concern. For both of these arguments, we first note that once u is infected, the time of its recovery is independent of any

transmissions it causes. So the modification of u does not alter the probability that u has a given status.

The first argument is that none of the effects of modifying u are relevant. Modifying u does not affect its probability of being infected. We have already seen that in the original epidemic (before u is modified), the proportion of individuals in each state is equal to the probability u is in each state. We have a series of equivalent questions. The first is, what proportions of the population are in each state in the original population. This is equivalent to our second question, what is the probability a randomly chosen individual u is in each state in the original population. This is equivalent to our third question, what is the probability u is in each state if it is prevented from transmitting. This is not altered by preventing u from transmitting. The changes only come to other individuals. At no point do we need to know anything in the modified population except the status of u . So the impact does not affect any quantities we calculate.

Our second argument is that modifying u has a negligible impact on the proportion infected in the population. Although this is not needed for our argument here, it is relevant for derivation of final sizes [30]. To make this point, we use analogy to the “price taker” assumption of economics. A firm is a price taker if it is too small to influence the price for its product. Consequently, if all firms in a given market are price takers, we can determine how the actions of each firm depends on the price. Then we determine how the price depends on the actions. This will give a system of equations and we have a consistency relation which we can solve to find the strategies and resulting price. We do not need to worry that an individual firm will have to change its strategy in response to the impact its strategy has on the price.

When we assume that a stochastic process is behaving deterministically on some large aggregate scale, we are making a similar assumption. In particular, for a disease spreading through a population, if we can assume that the aggregate dynamics are deterministic, then we are implicitly assuming that whether a particular individual is infected or not (and when that infection occurs) has no influence on the dynamics of the epidemic. Not only does the individual’s infection not have any measurable aggregate-scale impact, but also the infections traced back to that individual have no measurable aggregate-scale impact. Thus when we consider a deterministic epidemic we can make an “infection taker” assumption: We can ignore the impact of modifying u because it has no impact.³

So for the purposes of determining the final proportion infected, we can calculate the probability an individual u is infected given the amount of transmission that happens, but ignoring its impact on transmission. Then we calculate the amount of transmission that will happen given the proportion of the population that is infected. This leads to a consistency relation in which we know the proportion infected as a function of the proportion infected.

B Model hierarchy

In this appendix we show that under reasonable assumptions, the models presented in this paper are in fact equivalent. We have three subtly different closure approximations making slightly different assumptions about the independence of partners. Depending on which assumptions hold, different models result, but all ultimately become identical in appropriate limits. We will show that by making appropriate assumptions, we can derive some of the models from others.

B.1 An example

In several cases, the technique we use is a careful application of integrating factors. We demonstrate this technique with a different physical problem for which most people’s intuition is stronger.

Let us assume there is a single release of a radioactive isotope into the environment. The isotope may be in the air (A), in soil (S), or in biomass (B). It decays in time with rate τ independently of where it is. Assume the fluxes between the compartments are as in figure 9.

³In fact, this explains why final sizes from epidemic simulations in smaller populations are often very similar to larger populations even if the dynamics are still highly stochastic: The timing of an individual’s infection may have a significant impact on the aggregate number infected at any given time and therefore be important dynamically, even if it has little impact on the final size.

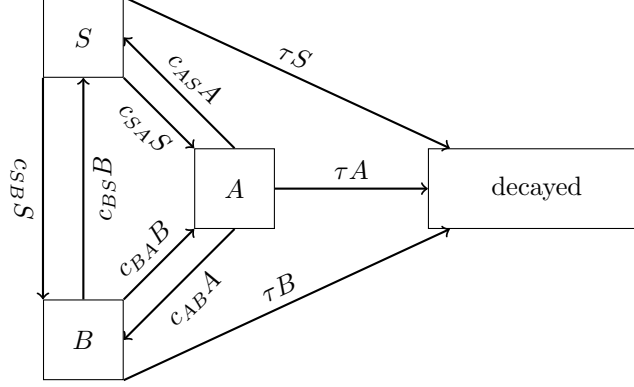


Figure 9: **The flow diagram for the flux of a radioactive isotope between soil, biomass and air.** The isotope decays at rate τ . This model provides a useful example of a technique that will be used in later derivations.

Then the equations are

$$\begin{aligned}\dot{A} &= -\tau A + c_{SA}S - c_{AS}A + c_{BA}B - c_{AB}A \\ \dot{S} &= -\tau S - c_{SA}S + c_{AS}A - c_{SB}S - c_{BS}B \\ \dot{B} &= -\tau B - c_{BA}B + c_{AB}A - c_{SB}S + c_{BS}B.\end{aligned}$$

However, if we define a , s , and b to be the probability that a test atom which does not decay is in each compartment, then the decayed class disappears. We get the new flow diagram shown in figure 10. The new equations are

$$\begin{aligned}\dot{a} &= c_{SA}s - c_{AS}a + c_{BA}b - c_{AB}a \\ \dot{s} &= -c_{SA}s + c_{AS}a - c_{SB}s - c_{BS}b \\ \dot{b} &= -c_{BA}b + c_{AB}a - c_{SB}s + c_{BS}b.\end{aligned}$$

Physically this change of variables is fairly obvious. We are calculating the probability an isotope is in a given compartment conditional on it having not yet decayed.

Mathematically we can get the new system of equations from the original through an integrating factor of $\alpha e^{\tau t}$. We set $a = A\alpha e^{\tau t}$, $b = B\alpha e^{\tau t}$, and $c = C\alpha e^{\tau t}$ with α chosen so that the initial amounts sum to 1. If we multiply the \dot{A} equation by $e^{\tau t}$, we end up with $\frac{d}{dt}(Ae^{\tau t}) = (c_{SA}S - c_{AS}A + c_{BA}B - c_{AB}A)e^{\tau t}$. Using the variable changes, we immediately arrive at the \dot{a} equation. The other equations transform similarly. So using an integrating factor to eliminate the decay term is equivalent to transforming into variables that measure the probability an undecayed isotope is in each compartment.

In general for other systems, so long as all compartments have an identical decay rate and the terms in the equations are homogeneous of order 1, then it is possible to use an integrating factor in this way to define a change of variables that eliminates the decay term. This will be a key step in deriving the EBCM approach from the other models. Here the decay rate corresponding to infection of an individual from outside the pair plays this role.

B.2 Simplifications of basic pairwise model

We begin by showing that the basic pairwise model can be reduced to the compact pairwise model, and that in turn, this is equivalent to the EBCM model.

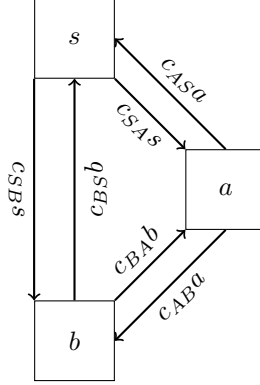


Figure 10: **The flow diagram for the flux of a radioactive isotope between soil, biomass and air.** When we focus on a test particle which does not decay, the number of fluxes and compartments to consider is reduced. This can be achieved mathematically through an integrating factor.

B.2.1 Deriving compact pairwise model from basic pairwise model

We presented two pairwise models. In both, we assumed the triples closure: Nothing we know about one partner of a susceptible individual u gives any information about another partner of u . We showed that the first reduces to the second if we assume that given susceptible u nothing we know about its degree gives any information about whether its partner v is infected or susceptible. Mathematically, this states that $\langle I \rangle_k = [S_k I]/k[S_k]$ and $\langle S \rangle_k = [S_k S]/k[S_k]$ are independent of k . The combination of these two assumptions gives us the pairs closure. So under the pairs closure, we expect the compact pairwise model to hold.

Our derivation of the EBCM model was based on the pairs closure. So we expect some change of variables to show that it is equivalent to the compact pairwise model.

It was previously noted [11] that if we make the generic assumption that $[A_k B] = [AB]k[A_k]/\sum_l l[A_l]$ where $[A_k B]$ represents the number of partnerships between individuals of status A having k partners and individuals of status B , then a pairwise approach can be used to derive an early version of the EBCM model [47]. In general, we expect this assumption to fail if A is either I or R . However, in the particular case where status A is susceptible, the assumption is consistent: Regardless of the degree of an individual, it has no impact on the status of its neighbors so long as it remains susceptible. We do not need the general form of the closure for our derivation, just the particular case with $A = S$.

In the derivation of the compact pairwise model, we claimed that $\langle I \rangle$, the probability a partner of a susceptible individual u is infected, is independent of k . This follows from the pairs closure, but we did not prove that if we start with the basic pairwise model and assume this probability is independent of k , then it remains independent of k at all later times. To address this, we turn to $\langle I \rangle_k = [S_k I]/k[S_k]$ and $\langle S \rangle_k = [S_k S]/k[S_k]$. We will take the derivative of $\langle I \rangle_k$, and show that if these are initially k -independent,

then its derivative is k -independent. We have

$$\begin{aligned}
\langle \dot{I} \rangle_k &= \frac{[\dot{S}_k I]}{k[S_k]} - \frac{[S_k I]k[\dot{S}_k]}{k^2[S_k]^2} \\
&= \frac{\sum_{k'} [S_k \dot{I}_{k'}]}{k[S_k]} - \langle I \rangle_k \frac{k[\dot{S}_k]}{k[S_k]} \\
&= \frac{\sum_{k'} \beta \frac{k'-1}{k'} \frac{[S_k S_{k'}][S_{k'} I]}{[S_{k'}]} - \gamma [S_k I_{k'}] - \beta \frac{k-1}{k} \frac{[I S_k][S_k I_{k'}]}{[S_k]} - \beta [S_k I_{k'}]}{k[S_k]} - \langle I \rangle_k \frac{k(-\beta[S_k I])}{k[S_k]} \\
&= \frac{-\gamma + \beta - \beta(k-1)\langle I \rangle_k [S_k I] + \sum_{k'} \beta(k'-1)[S_k S_{k'}]\langle I \rangle_{k'}}{k[S_k]} - \langle I \rangle_k \frac{k(-\beta[S_k I])}{k[S_k]} \\
&= -(\gamma + \beta)\langle I \rangle_k - \beta(k-1)\langle I \rangle_k^2 + \frac{\sum_{k'} \beta(k'-1)[S_k S_{k'}]\langle I \rangle_{k'}}{k[S_k]} + \beta k \langle I \rangle_k^2 \\
&= -(\gamma + \beta)\langle I \rangle_k - \beta(k-1)\langle I \rangle_k^2 + \frac{\beta[S_k S] \sum_{k'} k'(k'-1)[S_{k'}]\langle I \rangle_{k'}}{k[S_k] \sum_{k''} k''[S_{k''}]} + \beta k \langle I \rangle_k^2 \\
&= -(\gamma + \beta)\langle I \rangle_k - \beta(k-1)\langle I \rangle_k^2 + \frac{[S_k S] \beta \sum_{k'} k'(k'-1)[S_{k'}]\langle I \rangle_{k'}}{k[S_k] \sum_{k''} k''[S_{k''}]} + \beta k \langle I \rangle_k^2 \\
&= -(\gamma + \beta)\langle I \rangle_k - \beta(k-1)\langle I \rangle_k^2 + \langle S \rangle_k \frac{\beta \sum_{k'} k'(k'-1)[S_{k'}]\langle I \rangle_{k'}}{\sum_{k''} k''[S_{k''}]} + \beta k \langle I \rangle_k^2 \\
&= -(\gamma + \beta)\langle I \rangle_k + \beta \langle I \rangle_k^2 + \langle S \rangle_k \frac{\beta \sum_{k'} k'(k'-1)[S_{k'}]\langle I \rangle_{k'}}{\sum_{k''} k''[S_{k''}]} .
\end{aligned}$$

So we see that if $\langle I \rangle_k$ and $\langle S \rangle_k$ are independent of k at a given time, then the derivative of $\langle I \rangle_k$ is also independent of k . A similar calculation shows that the derivative of $\langle S \rangle_k$ is independent of k . Thus we conclude that if at any time all $\langle I \rangle_k$ and all $\langle S \rangle_k$ are independent of k , they remain so for future time.

This combined with the work in the main text shows that if $\langle I \rangle_k$ and $\langle S \rangle_k$ are initially k -independent (equivalently, the pairs closure holds), then the basic pairwise model reduces to the compact pairwise model. We are now ready to derive the EBCM equations from the compact pairwise model.

B.2.2 Deriving EBCM model from compact pairwise model.

We now derive the EBCM model from the compact pairwise model. We later derive the compact pairwise model from the EBCM model.

We begin our derivation with the observation that (for all k) $[\dot{S}_k] = -\beta k \langle I \rangle [S_k]$. So $[S_k](t) = [S_k](0) e^{-\beta k \int_0^t \langle I \rangle(\tau) d\tau}$. We define $\theta(t) = e^{-\beta \int_0^t \langle I \rangle(\tau) d\tau}$. Then $[S_k](t) = [S_k](0) \theta(t)^k$. If we define $\psi(\theta) = \sum_k \frac{[S_k](0)}{N} \theta^k$, then $S(t) = N\psi(\theta)$ and $\langle k \rangle_{\text{ex}} = \theta \psi''(\theta) / \psi'(\theta)$. We have $\dot{\theta} = -\beta \langle I \rangle \theta$.

We return to the equations

$$\begin{aligned}
[\dot{S}S] &= -2\beta \langle k \rangle_{\text{ex}} \langle I \rangle [SS] \\
[\dot{S}I] &= \beta \langle k \rangle_{\text{ex}} \langle I \rangle [SS] - (\beta + \gamma) [SI] - \beta \langle k \rangle_{\text{ex}} \langle I \rangle [SI] \\
[\dot{S}R] &= \gamma [SI] - \beta \langle k \rangle_{\text{ex}} \langle I \rangle [SR].
\end{aligned}$$

We observe that each of these equations has a term which represents infection of the first individual in the partnership from a source outside the partnership (with the same coefficient, $\beta \langle k \rangle_{\text{ex}} \langle I \rangle$, each time). So we anticipate that an integrating factor could eliminate this term. In the EBCM approach, the test individual is prevented from causing infection. This allows us to ignore infections to the test node from individuals other than the partner along a given edge. Mathematically, this shows up as ignoring infections of the first individual in the partnership from outside the partnership.

We define $F(t)$ such that $F'(t) = \beta \langle k \rangle_{\text{ex}} \langle I \rangle$. There is an arbitrary constant in F which we will choose later. Then multiplying through by $e^{F(t)}$ we find

$$\begin{aligned}\frac{d}{dt}([SS]e^{F(t)}) &= -\beta \langle k \rangle_{\text{ex}} \langle I \rangle [SS]e^{F(t)} \\ \frac{d}{dt}([SI]e^{F(t)}) &= \beta \langle k \rangle_{\text{ex}} \langle I \rangle [SS]e^{F(t)} - (\beta + \gamma)[SI]e^{F(t)} \\ \frac{d}{dt}([SR]e^{F(t)}) &= \gamma [SI]e^{F(t)}.\end{aligned}$$

We define $\phi_S = [SS]e^{F(t)}$, $\phi_I = [SI]e^{F(t)}$, and $\phi_R = [SR]e^{F(t)}$. Substituting these in and using $\langle k \rangle_{\text{ex}} = \theta \psi''(\theta)/\psi'(\theta)$, the equations become

$$\begin{aligned}\dot{\phi}_S &= -\beta \langle k \rangle_{\text{ex}} \langle I \rangle \phi_S = -\dot{\theta} \frac{\psi''(\theta)}{\psi'(\theta)} \phi_S \\ \dot{\phi}_I &= \beta \langle k \rangle_{\text{ex}} \langle I \rangle \phi_S - (\beta + \gamma) \phi_I = -\dot{\phi}_S - (\beta + \gamma) \phi_I \\ \dot{\phi}_R &= \gamma \phi_I.\end{aligned}$$

By directly substituting in, we can check that $\phi_S = \frac{\psi'(\theta)}{\psi'(1)} \phi_S(0)$.

We choose the arbitrary constant in $F(t)$ such that $\phi_S(0) + \phi_I(0) + \phi_R(0) = 1$. We note that the equations suggest that there is some quantity being moved between compartments. We have $\frac{d}{dt} \phi_S + \phi_I + \phi_R = -\beta \phi_I$. If we introduce compartments, we have flux from ϕ_S to ϕ_I , from ϕ_I to ϕ_R , and from ϕ_I to an as-yet-undefined compartment (which will become $1 - \theta$).

Since $\langle I \rangle = [SI]/([SS] + [SI] + [SR]) = \phi_I/(\phi_S + \phi_I + \phi_R)$, we conclude that

$$\begin{aligned}\frac{d\theta}{dt} &= -\beta \langle I \rangle \theta \\ &= -\beta \phi_I \frac{\theta}{\phi_S + \phi_I + \phi_R} \\ &= \frac{d\phi_S + \phi_I + \phi_R}{dt} \theta.\end{aligned}$$

Since our initial condition has $\theta(0) = 1 = \phi_S(0) + \phi_I(0) + \phi_R(0)$, direct substitution into this equation shows that $\theta = \phi_S + \phi_I + \phi_R$ is the solution. So $\dot{\theta} = -\beta \phi_I$.

Because $\dot{\phi}_R = -\gamma \dot{\theta}/\beta$, we can show that $\phi_R = \phi_R(0) + \frac{\gamma}{\beta}(1 - \theta)$. Using our equations for $\dot{\theta}$, our solutions for ϕ_S and ϕ_R , and our relation $\theta = \phi_S + \phi_I + \phi_R$, we finally arrive at the EBCM equations (38)–(41).

Deriving compact pairwise model from EBCM model Our goal is to show that using the EBCM model we can use a change of variables to arrive at the compact pairwise equations. Guided by the physical interpretation of our variables, our substitutions are:

$$\begin{aligned}[S] &= S \\ [I] &= I \\ [R] &= R \\ [S_k] &= S_k(0)\theta^k \\ [SS] &= \psi'(\theta)\phi_S \\ [SI] &= \psi'(\theta)\phi_I \\ \langle I \rangle &= \frac{\phi_I}{\theta} \\ \langle k \rangle_{\text{ex}} &= \theta \frac{\psi''(\theta)}{\psi'(\theta)}.\end{aligned}$$

Under these assumptions, we have

$$[\dot{S}_k] = S_k(0)k\theta^{k-1}\dot{\theta} = -\beta S_k(0)k\theta^{k-1}\phi_I = -\beta k S_k(0)\theta^k \langle I \rangle = -\beta k [S_k]$$

which is the compact pairwise model equation for $[\dot{S}_k]$. So the equation for $[\dot{S}_k]$ is as expected.

Since $[S] = S$ and $S = \sum_k S_k(0)\theta^k$, the equation for $[S]$ is $[S] = \sum_k S_k(0)\theta^k = \sum_k [S_k]$ as expected. The equations for $[I]$ and $[R]$ are immediately as anticipated.

We now look at $[SS] = \psi'(\theta)\phi_S$. We differentiate this to get

$$\begin{aligned} [\dot{SS}] &= \dot{\theta}\psi''(\theta)\phi_S + \psi'(\theta)\dot{\phi}_S \\ &= -\beta\phi_I\psi''(\theta)\phi_S + \psi'(\theta)\frac{d}{dt}\frac{\phi_S(0)\psi'(\theta)}{\psi'(1)} \\ &= -\beta\frac{\phi_I}{\theta}\theta\psi''(\theta)\phi_S + \psi'(\theta)\frac{\phi_S(0)\dot{\theta}\psi''(\theta)}{\psi'(1)} \\ &= -\beta\langle I \rangle\frac{\theta\psi''(\theta)}{\psi'(\theta)}\psi'(\theta)\phi_S - \beta\phi_I\psi'(\theta)\frac{\phi_S(0)\psi''(\theta)}{\psi'(1)} \\ &= -\beta\langle I \rangle\langle k \rangle_{\text{ex}}[SS] - \beta\frac{\phi_I}{\theta}\frac{\psi'(\theta)\phi_S(0)}{\psi'(1)}\theta\psi''(\theta) \\ &= -\beta\langle I \rangle\langle k \rangle_{\text{ex}}[SS] - \beta\langle I \rangle\phi_S\psi'(\theta)\frac{\theta\psi''(\theta)}{\psi'(\theta)} \\ &= -\beta\langle I \rangle\langle k \rangle_{\text{ex}}[SS] - \beta\langle I \rangle[SS]\langle k \rangle_{\text{ex}} \\ &= -2\beta\langle I \rangle\langle k \rangle_{\text{ex}}[SS]. \end{aligned}$$

Which is the compact pairwise equation for $[\dot{SS}]$.

We next look at $[SI] = \psi'(\theta)\phi_I$. We differentiate it to get

$$\begin{aligned} [\dot{SI}] &= \frac{d}{dt}[\psi'(\theta)\phi_I] \\ &= \dot{\theta}\psi''(\theta)\phi_I + \psi'(\theta)\dot{\phi}_I \\ &= \dot{\theta}\psi''(\theta)\phi_I + \psi'(\theta)\frac{d}{dt}(\theta - \phi_S - \phi_R) \\ &= \dot{\theta}\psi''(\theta)\phi_I + \psi'(\theta)(\dot{\theta} - \dot{\phi}_S - \dot{\phi}_R) \\ &= \dot{\theta}\psi''(\theta)\phi_I + \psi'(\theta)(-\beta\phi_I - \dot{\phi}_S - \gamma\phi_I) \\ &= \dot{\theta}\psi''(\theta)\phi_I - \psi'(\theta)\dot{\phi}_S - (\beta + \gamma)[SI] \\ &= \dot{\theta}\frac{\psi'(\theta)\langle k \rangle_{\text{ex}}}{\theta}\phi_I - \psi'(\theta)\dot{\phi}_S - (\beta + \gamma)[SI] \\ &= (-\beta\phi_I)\frac{[SI]\langle k \rangle_{\text{ex}}}{\theta} - \psi'(\theta)\dot{\phi}_S - (\beta + \gamma)[SI] \\ &= -\beta\langle I \rangle[SI]\langle k \rangle_{\text{ex}} - \psi'(\theta)\dot{\phi}_S - (\beta + \gamma)[SI] \\ &= -\beta\langle I \rangle[SI]\langle k \rangle_{\text{ex}} - (\beta + \gamma)[SI] - \psi'(\theta)\dot{\phi}_S. \end{aligned}$$

The term $\psi'(\theta)\dot{\phi}_S$ was shown above to be $-\beta\langle I \rangle\langle k \rangle_{\text{ex}}[SS]$. So this becomes

$$[\dot{SI}] = -\beta\langle I \rangle[SI]\langle k \rangle_{\text{ex}} - (\beta + \gamma)[SI] + \beta\langle I \rangle\langle k \rangle_{\text{ex}}[SS]$$

which is the compact pairwise equation for $[\dot{SI}]$.

We now look at $\langle I \rangle$. We have assumed $\langle I \rangle = \phi_I/\theta$. This becomes

$$\begin{aligned}
\langle I \rangle &= \frac{\phi_I}{\theta} \\
&= \frac{[SI]}{\psi'(\theta)\theta} \\
&= \frac{[SI]}{\theta \sum_k S_k(0)k\theta^{k-1}} \\
&= \frac{[SI]}{\sum_k S_k(0)k\theta^k} \\
&= \frac{[SI]}{\sum_k k[S_k]}
\end{aligned}$$

which is the compact pairwise equation for $\langle I \rangle$.

We are finally left with $\langle k \rangle_{\text{ex}}$. We have

$$\begin{aligned}
\langle k \rangle_{\text{ex}} &= \theta \frac{\psi''(\theta)}{\psi'(\theta)} \\
&= \frac{\theta \sum_k k(k-1)S_k(0)\theta^{k-2}}{\sum_k kS_k(0)\theta^{k-1}} \\
&= \frac{\theta \theta \sum_k k(k-1)S_k(0)\theta^{k-2}}{\theta \sum_k kS_k(0)\theta^{k-1}} \\
&= \frac{\sum_k k(k-1)S_k(0)\theta^k}{\sum_k kS_k(0)\theta^k} \\
&= \frac{\sum_k k(k-1)[S_k]}{\sum_k k[S_k]}
\end{aligned}$$

which is the compact pairwise equation for $\langle k \rangle_{\text{ex}}$.

Thus we have derived all of the equations for the compact pairwise model from the EBCM model. Since each model can be derived from the other, the models are equivalent.

B.3 Simplification of the effective degree models

Because of the order that different parts of this paper were developed, we find that rather than deriving the reduced effective degree model from the basic effective degree model, it is more practical to derive the basic pairwise model from the basic effective degree by adding the pairs closure. Then we show that the reduced effective degree model is equivalent to the EBCM model, which we have already shown is equivalent to the basic effective degree model. This suffices to prove that the reduced effective degree model can be derived from the basic effective degree model using the pairs closure. For completeness, we later sketch the derivation of the reduced effective degree model from the basic effective degree model.

It should be noted that although we can recover the compact pairwise model from both the basic effective degree model and the basic pairwise model using the pair closure, we cannot recover the basic pairwise model from the basic effective degree model (or *vice versa*). The two basic models have slightly different underlying assumptions (the effective degree model makes the “star closure” while the basic pairwise model makes the “triples closure”. Neither is a special case of the other.

B.3.1 Deriving compact pairwise model from basic effective degree model

The effective degree model of [20] can be used to derive a pairwise model closely related to the compact pairwise model we used. Once the appropriate additional closure assumption is made, it becomes the compact pairwise model.

To derive the compact pairwise model starting from the basic effective degree model, we begin by defining

$$[ss] = \sum_{s,i} sx_{s,i}$$

$$[si] = \sum_{s,i} ix_{s,i}.$$

These will represent the number of susceptible-susceptible edges (counted twice, once with each partner as the first individual) and the number of susceptible-infected edges.

We define

$$[ssi] = \sum_{s,i} siox_{s,i}$$

$$[isi] = \sum_{s,i} i(i-1)x_{s,i}.$$

These will represent the number of triples of the corresponding types. We define

$$\hat{\xi} = \frac{\sum_{s,i} i s x_{s,i}}{\sum_{s,i} s x_{s,i}}$$

$$= \frac{[ssi]}{[ss]}$$

and

$$\hat{\zeta} = \frac{\sum_{s,i} i^2 x_{s,i}}{\sum_{s,i} i x_{s,i}}$$

$$= \frac{\sum_{s,i} i(i-1)x_{s,i} + ix_{s,i}}{\sum_{s,i} i x_{s,i}}$$

$$= \frac{[isi]}{[si]} + 1.$$

These correspond to the ξ and ζ of the compact effective degree model.

We have

$$\begin{aligned} [\dot{ss}] &= \sum_{s,i} s\dot{x}_{s,i} \\ &= \sum_{s,i} s(-\beta ix_{s,i}) + \sum_{s,i} s\gamma(i+1)x_{s,i+1} - \sum_{s,i} s\gamma ix_{s,i} + \beta\xi \sum_{s,i} s[(s+1)x_{s+1,i-1} - sx_{s,i}] \\ &= -\beta \sum_{s,i} siox_{s,i} + \beta\xi \sum_{s,i} s[(s+1)x_{s+1,i-1} - sx_{s,i}] \\ &= -\beta \sum_{s,i} siox_{s,i} + \beta\xi \sum_{s,i} (s+1)^2 x_{s+1,i-1} - \beta\xi \sum_{s,i} (s+1)x_{s+1,i-1} - \beta\xi \sum_{s,i} s^2 x_{s,i} \\ &= -\beta \sum_{s,i} siox_{s,i} - \beta\xi \sum_{s,i} (s+1)x_{s+1,i-1} \\ &= -\beta\xi \sum_{s,i} sx_{s,i} - \beta\xi [ss] \\ &= -2\beta\xi [ss] \\ &= -2\beta [ssi]. \end{aligned}$$

Going from the third to the fourth line, we used the fact that $\sum_{s,i} s(i+1)x_{s,i+1} = \sum_{s,i} s i x_{s,i}$, and going from the fourth to the fifth line we used $\sum_{s,i} (s+1)^2 x_{s+1,i-1} = \sum_{s,i} s^2 x_{s,i}$. We further have

$$\begin{aligned}
\dot{[si]} &= \sum_{s,i} i \dot{x}_{s,i} \\
&= \sum_{s,i} i(-\beta i x_{s,i}) + \gamma \sum_{s,i} i(i+1)x_{s,i+1} - \gamma \sum_{s,i} i^2 x_{s,i} + \beta \xi \sum_{s,i} i[(s+1)x_{s+1,i-1} - s x_{s,i}] \\
&= -\beta \sum_{s,i} i^2 x_{s,i} + \gamma \sum_{s,i} (i+1)^2 x_{s,i+1} - \gamma \sum_{s,i} i^2 x_{s,i} - \gamma \sum_{s,i} (i+1)x_{s,i+1} + \beta \xi \sum_{s,i} i[(s+1)x_{s+1,i-1} - s x_{s,i}] \\
&= -\gamma[si] - \beta \sum_{s,i} i^2 x_{s,i} + \beta \xi \sum_{s,i} (i-1)(s+1)x_{s+1,i-1} + \beta \xi \sum_{s,i} (s+1)x_{s+1,i-1} - \beta \xi \sum_{s,i} s i x_{s,i} \\
&= -\gamma[si] - \beta \sum_{s,i} i^2 x_{s,i} + \beta \xi \sum_{s,i} (s+1)x_{s+1,i-1} \\
&= -\beta \zeta[si] - \gamma[si] + \beta \xi[ss] \\
&= -\beta[isi] - \beta[si] - \gamma[si] + \beta[ssi].
\end{aligned}$$

These are the equations of the global (unclosed) pairwise model at the level of pairs without using either the triples or the pairs closure.

Rather than using the triples closure to simplify the triples terms in these equations, we use the star closure, or more precisely, we use the basic effective degree model to simplify the triples terms. We can derive equations for the rate of change of $[isi] = \sum_{s,i} i(i-1)x_{s,i}$ and the rate of change of $[ssi] = \sum_{s,i} s i x_{s,i}$ simply by using the derivative of $x_{s,i}$, equation (19). When we use this, we are introducing the star closure to the global (unclosed) pairwise model. Although in general the rate of change of $[isi]$ and $[ssi]$ should depend on groupings of four individuals either as 3 individuals connected to a central individual or 4 individuals in a path, when we assume the derivative of $x_{s,i}$ from equation (19), we are assuming that we can safely average out the case of 4 individuals in a path. This is the simplification of the star closure. So we can express the basic effective degree model in terms of pairs and triples rather than effective degree.

We now add the pairs closure to this model. Let us choose a random ss partnership and label one individual u and the other v . Because the partnership is randomly chosen, we do not know the effective degree of u . Note that u is chosen with probability proportional to the number of ss partnerships it is in. The expected number of additional partners of u is $\langle k \rangle_{\text{ex}} = \sum_{s,i} s(i+s-1)x_{s,i} / \sum_{s,i} s x_{s,i}$. The probability that a partner $w \neq v$ is infected is $\langle I \rangle = \sum_{s,i} s i x_{s,i} / \sum_{s,i} s(i+s-1)x_{s,i}$ (the numerator gives the number of ways to choose v , u , and w such that v and u are susceptible and w infected while the denominator gives the number of ways to choose v and u to be susceptible and w to be either susceptible or infected). Thus the number of ssi triples is $[ssi] = [ss]\langle k \rangle_{\text{ex}} \langle I \rangle$. If we repeat this argument with an si partnership having u susceptible and v infected, we will get a new expression for the probability w is infected: $\langle I \rangle = \sum_{s,i} i(i-1)x_{s,i} / \sum_{s,i} i(i+s-1)x_{s,i}$. However, the pairs closure means both values of $\langle I \rangle$ are the same and both values of $\langle k \rangle_{\text{ex}}$ are the same. So we find $[isi] = [is]\langle k \rangle_{\text{ex}} \langle I \rangle$. Plugging this in we have

$$\begin{aligned}
\dot{[ss]} &= -2\beta[ss]\langle k \rangle_{\text{ex}} \langle I \rangle \\
\dot{[si]} &= -\beta[is]\langle k \rangle_{\text{ex}} \langle I \rangle - (\beta + \gamma)[si] + \beta[ss]\langle k \rangle_{\text{ex}} \langle I \rangle.
\end{aligned}$$

These are the equations we need for the pairs in the compact pairwise model, equations (11)–(18). The remaining equations follow easily.

B.3.2 Equivalence of the reduced effective degree model and the EBCM model

Transforming reduced effective degree model to EBCM model We begin with the reduced effective degree model, equations (32)–(37). We will look at the equation for $\dot{\nu}$ and an equation for $\frac{d}{dt} \sum_j j x_j$. Our goal is to derive the EBCM equations. We anticipate that ν should be proportional to ϕ_I and $\sum_j j x_j$ should

be proportional to $\phi_S + \phi_I$. When we consider an individual u and a partner v in the EBCM approach, we never have to consider the effect of u being infected by a partner other than v . Guided by this, we expect an integrating factor to allow us to throw out terms that represent infection coming from outside the partnership. We begin with $\sum_j jx_j$. This sums all of the partnerships from the perspective of each susceptible individual. We look for a term that corresponds to eliminating an edge from consideration because the focal individual is infected along a different edge.

$$\begin{aligned}
\frac{d}{dt} \sum_j jx_j &= \sum_j j\dot{x}_j \\
&= \gamma\langle I \rangle \sum_j ((j+1)jx_{j+1} - j^2x_j) - \beta\langle I \rangle \sum_j j^2x_j \\
&= \gamma\langle I \rangle \sum_j ((j+1)jx_{j+1} - j^2x_j) - \beta\langle I \rangle \sum_j (j(j-1)x_j + jx_j) \\
&= \gamma\langle I \rangle \sum_j ((j+1)jx_{j+1} - j^2x_j) - \beta\langle I \rangle \sum_j j(j-1)x_j - \beta\langle I \rangle \sum_j jx_j \\
&= \gamma\langle I \rangle \sum_j ((j+1)jx_{j+1} - j^2x_j) - \beta\langle I \rangle \frac{\sum_j j(j-1)x_j}{\sum_j jx_j} \sum_j jx_j - \beta\langle I \rangle \sum_j jx_j.
\end{aligned}$$

The term $-\beta\langle I \rangle \frac{\sum_j j(j-1)x_j}{\sum_j jx_j} \sum_j jx_j$ is exactly the term of interest. To remove it, we set $F'(t) = \beta\langle I \rangle \frac{\sum_j j(j-1)x_j}{\sum_j jx_j}$ and multiply by $e^{F(t)}$ (there is an arbitrary constant of integration which we will set later). We define $\phi_S + \phi_I = e^{F(t)} \sum_j jx_j$.

$$\begin{aligned}
\frac{d}{dt} (\phi_S + \phi_I) &= \gamma\langle I \rangle \sum_j ((j+1)jx_{j+1} - j^2x_j) e^{F(t)} - \beta\langle I \rangle \sum_j jx_j e^{F(t)} \\
&= \gamma\langle I \rangle \left(\sum_j ((j+1)^2 - (j+1))x_{j+1} - \sum_j j^2x_j \right) e^{F(t)} - \beta\langle I \rangle (\phi_S + \phi_I) \\
&= -\gamma\langle I \rangle \sum_j (j+1)x_{j+1} e^{F(t)} - \beta\langle I \rangle (\phi_S + \phi_I) \\
&= -\gamma\langle I \rangle (\phi_S + \phi_I) - \beta\langle I \rangle (\phi_S + \phi_I).
\end{aligned}$$

We now move to ν . This counts the number of susceptible-infected partnerships. We look for the term that corresponds to infection of the susceptible individual from outside the partnership.

$$\begin{aligned}
\dot{\nu} &= -(\beta + \gamma)\nu + \beta\langle I \rangle (1 - 2\langle I \rangle) \sum_j j(j-1)x_j \\
&= -(\beta + \gamma)\nu + \beta\langle I \rangle (1 - \langle I \rangle) \sum_j j(j-1)x_j - \beta\langle I \rangle^2 \sum_j j(j-1)x_j \\
&= -(\beta + \gamma)\nu + \beta\langle I \rangle (1 - \langle I \rangle) \sum_j j(j-1)x_j - \beta\langle I \rangle \frac{\sum_j j(j-1)x_j}{\sum_j jx_j} \nu.
\end{aligned}$$

The final term is the term we want. Not coincidentally, the coefficient is the same as we saw above for $\sum_j jx_j$. We use the same integrating factor and define $\phi_I = \nu e^{F(t)}$. We have

$$\begin{aligned}
\dot{\phi}_I &= -(\beta + \gamma)\nu e^{F(t)} + \beta \langle I \rangle (1 - \langle I \rangle) \sum_j j(j-1)x_j e^{F(t)} \\
&= -(\beta + \gamma)\phi_I + \beta \langle I \rangle \left(1 - \frac{\nu}{\sum_j jx_j}\right) \sum_j j(j-1)x_j e^{F(t)} \\
&= -(\beta + \gamma)\phi_I + \beta \langle I \rangle \left(\frac{\sum_j jx_j - \nu}{\sum_j jx_j}\right) \sum_j j(j-1)x_j \\
&= -(\beta + \gamma)\phi_I + \beta \langle I \rangle \phi_S \frac{\sum_j j(j-1)x_j}{\sum_j jx_j} \\
&= -(\beta + \gamma)\phi_I + \phi_S F'(t).
\end{aligned}$$

Note that it is now clear that $\langle I \rangle = \phi_I / (\phi_S + \phi_I)$. So we can simplify somewhat. Our expression for $\text{diff}(\phi_S + \phi_I)$ becomes $-(\beta + \gamma)\phi_I$. Using the equations

$$\begin{aligned}
\frac{d}{dt}(\phi_S + \phi_I) &= -(\beta + \gamma)\phi_I \\
\frac{d}{dt}\phi_I &= -(\beta + \gamma)\phi_I + \phi_S F'(t)
\end{aligned}$$

we finally have

$$\begin{aligned}
\dot{\phi}_S &= -\phi_S F'(t) \\
\dot{\phi}_I &= -(\beta + \gamma)\phi_I + \phi_S F'(t).
\end{aligned}$$

We now introduce two variables θ and ϕ_R . We take $\theta(0) = 1$, $\dot{\theta} = -\beta\phi_I$, and $\phi_R = \theta - \phi_S - \phi_I$. It follows that $\dot{\phi}_R = \gamma\phi_I$. Note that as yet, ϕ_S and ϕ_I are only known up to a multiplicative constant (due to the constant of integration in F). We can arbitrarily set $\phi_S(0) + \phi_I(0) = 1$, so that $\phi_R(0) = 0$. In terms of the interpretation of the EBCM variables, this corresponds to saying that we can ignore individuals that are already recovered at $t = 0$. We can choose another constant in $(0, 1)$ if we wish to account for the already recovered partners of random individuals.

We turn our attention to finding x_j in terms of our new variables, which will allow us to determine $F'(t)$ in terms of our new variables. Guided by our expectations, we assume $x_j = \sum_{k \geq j} x_k(0) \binom{k}{j} (\phi_S + \phi_I)^j \phi_R^{k-j}$. At $t = 0$, this is correct. We now check that the evolution of x_j is correctly captured by this equation. We find

$$\begin{aligned}
\dot{x}_j &= \sum_{k \geq j} x_k(0) \binom{k}{j} \frac{d}{dt} ((\phi_S + \phi_I)^j \phi_R^{k-j}) \\
&= \sum_{k \geq j} x_k(0) \binom{k}{j} (-j(\beta + \gamma)\phi_I (\phi_S + \phi_I)^{j-1} \phi_R^{k-j} + \gamma(k-j)\phi_I (\phi_S + \phi_I)^j \phi_R^{k-j-1}) \\
&= \frac{\phi_I}{\phi_S + \phi_I} \sum_{k \geq j} x_k(0) \binom{k}{j} (-j(\beta + \gamma)(\phi_S + \phi_I)^{j-1} \phi_R^{k-j} + \gamma(k-j)(\phi_S + \phi_I)^j \phi_R^{k-j-1}) \\
&= \frac{\phi_I}{\phi_S + \phi_I} \sum_{k \geq j} x_k(0) \left(-\binom{k}{j} j(\beta + \gamma)(\phi_S + \phi_I)^{j-1} \phi_R^{k-j} + \binom{k}{j} (k-j)\gamma(\phi_S + \phi_I)^j \phi_R^{k-j-1} \right) \\
&= \langle I \rangle (-(\beta + \gamma)jx_j + \gamma x_{j+1}).
\end{aligned}$$

This is the equation for \dot{x}_j . So our ansatz is correct.

We now define $\psi(\theta) = \sum_k x_k(0)\theta^k$ so that

$$\sum_j x_j = \sum_k \sum_j x_k(0) \binom{k}{j} \phi_R^{k-j} (\phi_S + \phi_I)^j = \sum_k x_k(0) (\phi_S + \phi_I + \phi_R)^k = \psi(\theta).$$

We will use this to express F' in terms of the EBCM variables. We first derive two identities using the “derivative trick”:

$$\begin{aligned} \sum_{j=0}^{\infty} \binom{k}{j} j a^j b^{k-j} &= a \sum_{j=0}^{\infty} \binom{k}{j} j a^{j-1} b^{k-j} \\ &= a \frac{d}{da} \sum_{j=0}^{\infty} \binom{k}{j} a^j b^{k-j} \\ &= a \frac{d}{da} (a+b)^k \\ &= ka(a+b)^{k-1}, \end{aligned}$$

and

$$\begin{aligned} \sum_{j=0}^{\infty} \binom{k}{j} j(j-1) a^j b^{k-j} &= a^2 \sum_{j=0}^{\infty} \binom{k}{j} j(j-1) a^{j-2} b^{k-j} \\ &= a^2 \frac{d^2}{da^2} \sum_{j=0}^{\infty} \binom{k}{j} a^j b^{k-j} \\ &= a^2 \frac{d^2}{da^2} (a+b)^k \\ &= k(k-1)a^2(a+b)^{k-2}. \end{aligned}$$

So

$$\begin{aligned} F'(t) &= \beta \langle I \rangle \frac{\sum_j j(j-1)x_j}{\sum_j jx_j} \\ &= \beta \langle I \rangle \frac{\sum_k \sum_j j(j-1)x_k(0) \binom{k}{j} (\phi_S + \phi_I)^j \phi_R^{k-j}}{\sum_k \sum_j jx_k(0) \binom{k}{j} (\phi_S + \phi_I)^j \phi_R^{k-j}} \\ &= \beta \langle I \rangle \frac{(\phi_S + \phi_I)^2 \sum_k k(k-1)x_k(0)\theta^{k-2}}{(\phi_S + \phi_I) \sum_k kx_k(0)\theta^{k-1}} \\ &= \beta \langle I \rangle (\phi_S + \phi_I) \frac{\psi''(\theta)}{\psi'(\theta)} \\ &= \beta \phi_I \frac{\psi''(\theta)}{\psi'(\theta)}. \end{aligned}$$

So we now have

$$\dot{\phi}_S = -\phi_S F'(t) = -\phi_S \beta \phi_I \frac{\psi''(\theta)}{\psi'(\theta)}.$$

We make the observation that $\phi_S = \phi_S(0)\psi'(\theta)/\psi'(1)$ solves the ODE for ϕ_S . We have $\dot{\phi}_R = \gamma\phi_I$ and $\phi_I = \theta - \phi_S - \phi_R$. This combined with $\dot{\theta} = -\beta\phi_I$ completes our system and gets us back to the EBCM equations.

So we have shown that starting from the reduced effective degree equations, we can derive the EBCM equations with $\phi_R(0) = 0$. The assumption that $\phi_R(0) = 0$ forced us to use $x_k(0)$ rather than $S_k(0)$ for our definition of ψ . We could have used $S_k(0)$, but this would have required a different value of $\phi_R(0)$. The derivation would have been more complicated, but not significantly altered.

EBCM to reduced effective degree We now go in the opposite direction, we derive the reduced effective degree equations from the EBCM equations. This is less strenuous. We simply determine what the reduced effective degree variables should be in terms of the EBCM variables and then test that they satisfy the appropriate equations. We can drop the assumption that $\phi_R(0) = 0$. We start with the EBCM equations and write

$$x_j(t) = \sum_k S_k(0) \binom{k}{j} (\phi_S(t) + \phi_I(t))^j \phi_R(t)^{k-j}$$

[this is similar to the ansatz we used above for x_j , except we take $S_k(0)$ rather than $x_k(0)$]. We also anticipate that $\nu = \phi_I \psi'(\theta)$. This is expected because $\psi'(\theta)$ gives the sum of degrees of susceptible individuals and ϕ_I gives the probability an edge to a susceptible individual connects to an infected individual.

To test the equation for x_j we follow the steps we did above for the ansatz of x_j . These steps arrive at the expected ODE for x_j .

To test the equation for ν , we first recall that $\phi_S = \phi_S(0)\psi'(\theta)/\psi'(1)$. This will be used below. We have

$$\begin{aligned} \dot{\nu} &= \frac{d}{dt} \phi_I \psi'(\theta) \\ &= \left(-(\beta + \gamma)\phi_I + \frac{\phi_S(0)}{\psi'(1)} \beta \phi_I \psi''(\theta) \right) \psi'(\theta) + \phi_I (-\beta \phi_I) \psi''(\theta) \\ &= -(\beta + \gamma)\phi_I \psi'(\theta) + \frac{\phi_S(0)}{\psi'(1)} \psi'(\theta) \beta \phi_I \psi''(\theta) - \beta \phi_I^2 \psi''(\theta) \\ &= -(\beta + \gamma)\nu + \phi_S \beta \phi_I \psi''(\theta) - \beta \phi_I^2 \psi''(\theta) \\ &= -(\beta + \gamma)\nu + \beta \phi_I \psi''(\theta) (\phi_S - \phi_I) \\ &= -(\beta + \gamma)\nu + \beta \frac{\phi_I}{\phi_S + \phi_I} (\phi_S + \phi_I) \psi''(\theta) (\phi_S + \phi_I - 2\phi_I) \\ &= -(\beta + \gamma)\nu + \beta \langle I \rangle (\phi_S + \phi_I)^2 \psi''(\theta) (1 - 2\langle I \rangle) \\ &= -(\beta + \gamma)\nu + \beta \langle I \rangle (1 - 2\langle I \rangle) (\phi_S + \phi_I)^2 \sum_k k(k-1) S_k(0) \theta^{k-2}, \end{aligned}$$

Using $\theta = (\phi_S + \phi_I) + \phi_R$ and the derivative trick used above, we know $S_k(0)k(k-1)(\phi_S + \phi_I)^2 \theta^{k-2}$ is $\sum_j \binom{k}{j} j(j-1) S_k(0) (\phi_S + \phi_I)^j \phi_R^{k-j}$. So the final term above becomes $\beta \langle I \rangle (1 - 2\langle I \rangle) \sum_j j(j-1) \sum_k S_k(0) (\phi_S + \phi_I)^j \phi_R^{k-j} = \beta \langle I \rangle (1 - 2\langle I \rangle) \sum_j j(j-1) x_j$. Thus we have

$$\dot{\nu} = -(\beta + \gamma)\nu + \beta \langle I \rangle (1 - 2\langle I \rangle) \sum_j j(j-1) x_j.$$

So we can derive the reduced pairwise equations from the EBCM equations. The two models are equivalent.

B.3.3 Deriving reduced effective degree model from basic effective degree model

We now address the direct relation between the basic effective degree model and the reduced effective degree model. The two models use slightly different definitions of active partnerships, but these definitions coincide in the case of susceptible individuals. This allows us to derive the simplified equations from the basic effective degree model. We have already shown that it is possible to derive the reduced effective degree model from the basic effective degree model by a series of indirect steps deriving other models, so we only sketch this derivation.

We can define $x_{k_e} = \sum_{i+s=k_e} x_{s,i}$ where $x_{s,i}$ is the variable from the basic effective degree model. We find

$$\dot{x}_{k_e} = \sum_{s+i=k_e} \gamma(i+1)x_{s,i+1} - \gamma i x_{s,i} - \beta i x_{s,i} = \sum_{s+j=k_e+1} \gamma j x_{s,j} - \sum_{s+i=k_e} \gamma i x_{s,i} - \sum_{s+i=k_e} \beta i x_{s,i}.$$

We set $\langle I \rangle_{k_e} = \sum_{s+i=k_e} ix_{s,i} / \sum_{s+i=k_e} k_e x_{s,i}$ and make the assumption that $\langle I \rangle_{k_e}$ is in fact independent of k_e (that is, the probability a partner of a susceptible individual u is infected is independent of the number of active edges u has which follows from the pairs closure), then our equation reduces to

$$\dot{x}_{k_e} = \gamma \langle I \rangle (k_e + 1) x_{k_e+1} - \gamma k_e \langle I \rangle x_{k_e} - \beta k_e \langle I \rangle x_{k_e}$$

which is equation (32) with k_e replacing j .

If we define $\nu = \sum_s \sum_i ix_{s,i}$ we get

$$\begin{aligned} \dot{\nu} &= \sum_s \sum_i i \dot{x}_{s,i} \\ &= \sum_s \sum_i \gamma i [(i+1)x_{s,i+1} - ix_{s,i}] - \beta i^2 x_{s,i} + \beta i \xi [(s+1)x_{s+1,i-1} - sx_{s,i}]. \end{aligned}$$

After taking advantage of the fact that, for example $\sum_s \sum_i ix_{s,i} = \sum_s \sum_i (i+1)x_{s,i+1}$ and repeatedly reindexing some sums, and the fact that $\xi = \sum_s \sum_i ix_{s,i} / \sum_s \sum_i sx_{s,i}$ we can convert this to

$$\dot{\nu} = -(\gamma + \beta)\nu + \beta \sum_s \sum_i -i(i-1)x_{s,i} + \beta \sum_s \sum_i si x_{s,i}.$$

The sum $\sum_s \sum_i i(i-1)x_{s,i}$ counts the total number of (ordered) triples (v, u, w) where u has partnerships with both v and w , u is susceptible, and v and w are both infected. We find this value through a different accounting based on the pairs closure: We count the total number of triples having a susceptible central individual. The probability that both peripheral individuals are infected is $\langle I \rangle^2$. We get $\langle I \rangle^2 \sum_{k_e} k_e (k_e - 1) x_{k_e}$ where x_{k_e} is as above. Similarly, $\sum_s \sum_i si x_{s,i}$ becomes $\langle I \rangle (1 - \langle I \rangle) \sum_{k_e} k_e (k_e - 1) x_{k_e}$. Thus we finally have

$$\dot{\nu} = -(\beta + \gamma)\nu + \beta \langle I \rangle (1 - 2\langle I \rangle) \sum_{k_e} k_e (k_e - 1) x_{k_e}$$

which is the equation for ν from before.

To complete this derivation, we would need to additionally show that $\langle I \rangle_{k_e}$ remains independent of k_e if the initial conditions satisfy the pairs closure. This will require expanding the derivative of $\langle I \rangle_{k_e}$ and showing that if initially these are independent of k_e , then the derivatives are independent of k_e .

B.3.4 Deriving the compact effective degree model from the reduced effective degree model

To derive the compact effective degree model, we start with $\nu = \langle I \rangle \sum_j j x_j$. Anticipating that the two models to have the same value for $\langle I \rangle$, we note that in the compact effective degree model, $\lambda = \lambda \langle I \rangle + \langle I \rangle \sum_j j x_j$. Solving this, we find $\lambda = \nu / (1 - \langle I \rangle)$. So starting from the reduced model equations, we define $\lambda = \nu / (1 - \langle I \rangle)$. Taking the derivative of this and using the same sort of changes done above will show that the derivative of λ is indeed as found for the compact effective degree model.

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