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Measuring infection transmission in a stochastic SIV model with infection reintroduction and imperfect vaccine

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Abstract An additional compartment of vaccinated individuals is considered in a SIS stochastic epidemic model with infection reintroduction. The quantification of the spread of the disease is modeled by a continuous time Markov chain. A well-known measure of the initial transmission potential is the basic reproduction number R_0 , which determines the herd immunity threshold or the critical proportion of immune individuals required to stop the spread of a disease when a vaccine offers a complete protection. Due to repeated contacts between the typical infective and previously infected individuals, R_0 overestimates the average number of secondary infections and leads to, perhaps unnecessary, high immunization coverage. Assuming that the vaccine is imperfect, alternative measures to R_0 are defined in order to study the influence of the initial coverage and vaccine efficacy on the transmission of the epidemic.

 $\mathbf{Keywords}$ Stochastic Markovian epidemic · Imperfect vaccine · Basic reproduction number

Mathematics Subject Classification (2000) 92D30 · 60J28 · 60J22

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1 Introduction

Mathematical modeling is an essential tool to represent the progress of an epidemic through a population. It is commonly accepted that the work of Kermack and Mckendrick (1927) is the prototype of almost all epidemiological models based on a classification of individuals by their epidemic status. Since then, many other compartmental models have been developed to investigate a diverse range of communicable diseases to obtain a better knowledge of their transmission mechanisms (Anderson and Britton 2000; Kretzschmar et al. 2001; Aguiar et al. 2010; Artalejo and Lopez-Herrero 2014; Liu et al. 2018).

A common assumption is that the communicable disease spreads in a community of constant size. The population can be closed in the sense that infectious individuals can infect only other individuals within the population under study during the epidemic's time span. However, assuming reintroduction of the disease through contact with infected individuals from other areas could represent a more realistic scenario (Marchette and Wierman 2004; Stone et al. 2007; Amador 2016).

After infection, it is assumed that patients recover due to their own immune system which acts as a body's defense force against germs and other invading substances. Hence, to reduce the incidence of an infectious disease requires improving sanitary and living conditions. In that sense, we cannot ignore the impact of vaccination on this reduction, especially in the developing world. Vaccines activate the immune system's capacity of producing antibodies to fight diseases without exposing it to diseases-producing pathogens. If a vaccinated person comes into contact with the disease for which she/he has been vaccinated, her/his immune system recognizes the invading germs and immediately produces the antibodies that will kill foreign invaders. Generally, vaccines provide immunity similar to that acquired from the natural infection, and duration of protection varies depending on diseases and also on vaccine strains. Lifelong immunity is not always provided by vaccination and usually several doses of vaccine may be required. Furthermore, the immune response may wane over time and it is necessary to administrate new doses of vaccine to increase or restore immunity.

As it is an effective method of disease control, recent epidemiological models (Kribs-Zaleta and Martcheva 2002; Arino et al. 2003; Arino et al. 2010; Ball and Sirl 2013; Samanta 2015; Ball and Sirl 2018; Li and Zhang 2019) have added a vaccination compartment and vaccination strategies into their mathematical model. Some papers discussing the impact of vaccination on the spread of an epidemic assume complete protection (Iannelli et al. 2005; Alexander et al. 2006; Ball et al. 2007; Lin et al. 2014; Eckalbar and Eckalbar 2015; Guo 2017) but even vaccine efficacy, as measured by observational studies, is not 100% (e.g., measles: 90-95%, mumps: 72-88% or rubella: 95-98%) and depends on internal or individual factors, as well as on the dose and strain of the vaccine virus (Demicheli et al. 2012). In scientific literature, vaccine efficacy and effectiveness are often used interchangeably. Vaccine efficacy represents the reduction in the risk of infection at individual level under optimal condi-

tion (e.g., randomized controlled trials) while vaccine effectiveness compares rates of the transmission of the disease between vaccinated and unvaccinated individuals once the vaccine is approved for use in the general population.

The SIS model with imperfect vaccine has been studied from a deterministic point of view in the papers by Moghadas (2004) and Safan and Rihan (2014), under nonlinear incidence in Xiao and Tang (2010) and Yang et al. (2015), and with inclusion of a latency period as well as psychological effects in both susceptible and vaccinated individuals in Cheng et al. (2015). A SIV model with stochastic perturbations is discussed in Liu et al. (2018) and López-García (2016).

The above-mentioned studies deal with large populations, but transmission patterns become quantitatively different when a small population is involved. Firstly, individual variations of infectivity, recovery periods and vaccine protection, even for a homogeneous social group, should be taken into account. Moreover, when the susceptible population becomes depleted the extent of the epidemic is interrupted and, for closed populations, the extinction of the disease is possible (Keeling and Ross 2008).

The aim of this paper is to quantify the spread of an infectious disease that does not confer immunity, within a population that is partially protected against the disease by a vaccine. A continuous time Markov chain (CTMC) model represents changes in the composition of infected and vaccinated classes. Two random variables will quantify the transmission of the epidemic process with reintroduction: R_{e0} , the number of infectious cases caused directly by the first infected individual, and R_p , which is the number of infectious cases caused by any infectious spreader. These random variables act as stochastic counterparts to the basic reproduction number, R_0 , and more specifically the control reproduction number, R_c , when there is an available vaccine.

In the literature, the term herd-immunity threshold refers to the critical proportion of immune individuals that is needed to interrupt epidemic transmission in a population. There is a simple relationship between herd-immunity coverage and the basic reproduction number. If a perfect vaccine is available and a fraction f of the population is vaccinated, then the disease will not spread if $(1-f)R_0 < 1$. In general terms, we can represent the quality of a vaccine by a measure of the vaccine imperfection $h \in [0,1]$, with h=0 indicating a perfect vaccine, and h=1 a useless vaccine. For imperfect but not useless vaccines, the critical vaccination coverage level to eradicate the infection is related to the proportion $(1-1/R_0)/(1-h)$. In any case, herd immunity depends on estimates of R_0 , and is the result of a reduction in viral transmission caused by removing vaccinated individuals from the susceptible class. We will investigate the spread of the disease directly, at any time, by updating the current population state, and we will apply our alternative measures to control disease spread by fixing an adequate vaccine coverage level.

The structure of the paper is as follows. In Section 2, we describe the SIV stochastic model with infection reintroduction and summarize results for its long term behavior in terms of the reintroduction parameter. Section 3 introduces the proposed random variables, R_{e0} and R_p , as measures of the in-

fection transmission in a partially vaccinated population. Theoretical and algorithmic results will provide probability mass functions and moments for R_{e0} and R_p , depending on the initial vaccine coverage and effectiveness. Section 4 illustrates the applicability of the algorithmic results and presents several qualitative facts regarding the infection transmission. Finally, we present some concluding remarks and tentative future research lines.

2 Model description

We consider a homogeneous and uniformly mixing population of constant size, N, where individuals are affected by a contagious disease. This disease is transmitted by direct contact with an infected individual. The population is not isolated, so we assume that there is an additional source of infection due to external contacts. We suppose that some individuals in the population have been protected against the disease with an available vaccine that confers immunity, but it is not a perfect vaccine and some contacts between vaccinated and infectious individuals produce an effective contagion. Once a vaccinated individual gets the infection, he no longer belongs to the class of individuals that have been vaccinated and he belongs to the infective class while he is infectious. Recovered individuals become susceptible to the disease, no matter if they were previously vaccinated or not. Consequently, individuals in the population are classified into three separate classes, namely susceptible, vaccinated and infected. Vaccination was implemented at t=0 and no vaccination will take place after this epoch.

In general terms, the underlying mathematical model involves a SIV model, where movement of individuals among the three epidemiological classes is shown in Figure 1, with S denoting the class of susceptible individuals, I denoting the class of infected individuals, and V denoting the class of vaccinated individuals.

$$\begin{array}{c|c}
\hline
S & \hline
\hline
 & Infection \\
\hline
\hline
 & Recovery \\
\hline
\end{array} & I & \hline
 & Vaccine \\
\hline
 & failure \\
\hline
\end{array} & V$$

Fig. 1 SIV compartmental diagram

At any time t > 0, the state of the epidemic is described by random variables S(t), V(t), I(t), that record the number of susceptible, vaccinated and infective individuals, respectively, at time t. According to the constant size hypothesis we have

$$S(t) + V(t) + I(t) = N.$$

We represent the evolution of the disease in terms of a two-dimensional CTMC: $X = \{(V(t), I(t)), t \geq 0\}$. Assuming that initially the population contains v_0 vaccinated individuals, with $0 \leq v_0 \leq N$, the state space of X is $S = \{(v, i) : 0 \leq v \leq v_0, 0 \leq i \leq N, 0 \leq v + i \leq N\}$ that contains $(v_0 + 1)(N + 1 - v_0/2)$ states. Notice that once v = 0 the vaccination compartment is empty and the underlying model is the standard SIS epidemic one.

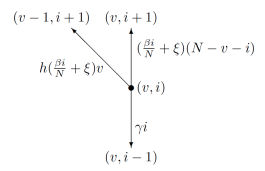
Next, we describe the dynamics of the Markov chain, X, in full details. See Table 1 for a summary of parameters used. The exhaustive description of the events and their transition rates is presented in Table 2. Finally, a diagram showing transitions from a general state $(v,i) \in S$ is depicted in Figure 2.

Table 1 Parameters of the model

Population size	N
Disease internal transmission rate	β
Disease external transmission rate	ξ
Probability of vaccine failure	h
Recovery rate	γ

Table 2 Effective events and their transition rates

Effective event	Transition	Rate
Susceptible-Infected internal contagion Susceptible-Infected external contagion Vaccinated-Infected internal contagion Vaccinated-Infected external contagion Recovery and loss of immunity	$ \begin{aligned} &(v,i) \to (v,i+1) \\ &(v,i) \to (v,i+1) \\ &(v,i) \to (v-1,i+1) \\ &(v,i) \to (v-1,i+1) \\ &(v,i) \to (v,i-1) \end{aligned} $	$\beta i(N-v-i)/N \\ \xi(N-v-i) \\ h\beta iv/N \\ h\xi v \\ \gamma i$



 ${\bf Fig.~2} \ \ {\bf Transitions~among~states}$

To understand the dynamics of the model it is necessary to describe the events that cause a change in the current model state (v, i). There are five possible effective events that are listed in Table 2.

 E_1 . A susceptible individual gets the infection from an infective individual within the population.

 E_2 . A susceptible individual gets the infection from an external source of infection.

 E_3 . Due to vaccine failure, a vaccinated individual becomes infected from an infective individual within the population.

 E_4 . Due to vaccine failure, a vaccinated individual becomes infected from an external source of infection.

 E_5 . An infective individual is recovered and becomes susceptible.

Sojourn times at each state in ${\sf S}$ are independent and exponential random variables, with rate

$$q_{v,i} = \left(\frac{\beta i}{N} + \xi\right) (N - v - i) + h \left(\frac{\beta i}{N} + \xi\right) v + \gamma i. \tag{1}$$

To describe \mathbf{Q} , the infinitesimal generator of the Markov chain X, we partition the state space in levels regarding the number of vaccinated individuals, that is $\mathsf{S} = \cup_{v=0}^{v_0} L(v)$, where level $L(v) = \{(v,i) \in \mathsf{S} : 0 \le i \le N-v\}$, for $0 \le v \le v_0$, which contains (N+1-v) states. Then, we can express the infinitesimal generator of X in the following non-null block form

where $\mathbf{Q_{v,v^*}}$, for $0 \le v, v^* \le v_0$, are matrices of dimension $(N+1-v) \times (N+1-v^*)$. The blocks $\mathbf{Q_{v,v-1}}$, for $1 \le v \le v_0$, correspond to transitions due to vaccine failures, and the blocks $\mathbf{Q_{v,v}}$, for $0 \le v \le v_0$, correspond to transitions within level L(v) that are due either to infections of susceptible or recoveries. Non-null sub-matrices are described as follows

$$\mathbf{Q_{v,v-1}} = \begin{pmatrix} 0 & h\xi v \\ 0 & h\left(\frac{\beta}{N} + \xi\right)v \\ & \ddots & \ddots \\ & & 0 & h\left(\frac{\beta(N-v)}{N} + \xi\right)v \end{pmatrix}$$

and

$$\mathbf{Q_{v,v}} = \begin{pmatrix} -q_{v,0} & \xi(N-v) \\ \gamma & -q_{v,1} & \left(\frac{\beta}{N} + \xi\right)(N-v-1) \\ & \ddots & \ddots & \ddots \\ (N-v-1)\gamma & -q_{v,N-v-1} & \frac{\beta(N-v-1)}{N} + \xi \\ & (N-v)\gamma & -q_{v,N-v} \end{pmatrix}$$

The above block bidiagonal structure of \mathbf{Q} guarantees that square matrices appearing in the forthcoming theoretical results are, at most, tridiagonal and diagonally dominant. Thus, they are non-singular, and linear systems involving these matrices can be solved efficiently by a simplified form of the Gaussian elimination algorithm (Golub and van Loan 1996).

2.1 Stationary behavior

As we are dealing with a finite state CTMC, the long-term behavior of X depends on the structure of communicating classes of absorbing states. In that sense, we notice that the reintroduction parameter (that is, the external transmission rate ξ) plays an important role in the classification of the states in S. If we assume that the population is isolated and contagions are produced only by internal contacts; i.e., $\xi=0$, then states with 0 infective individuals are absorbing. This fact and the finiteness of the state space guarantee that the process will become absorbed into any of the non-communicating classes of absorbing states with probability one. Hence, the epidemic extinction is certain and outbreaks involve a single epidemic episode that will last a finite expected time.

On the other hand, when $\xi > 0$, the state space of the finite CTMC X contains a single absorbing set given by $L(0) = \{(0,i) : 0 \le i \le N\}$. Thus, once the process enters into L(0) it can move across these states but it can not leave the absorbing set. Hence the stationary distribution assigns mass to every state with no vaccinated individuals. Which means that, in the model with external source of infection, occasionally the disease is faded away (i.e., I=0) for a short time, but the infection is reintroduced at a later time. More specifically, the theoretical long-term distribution of the number of infective individuals agrees with the stationary distribution provided in Stone et al. (2007), Section 2.6.

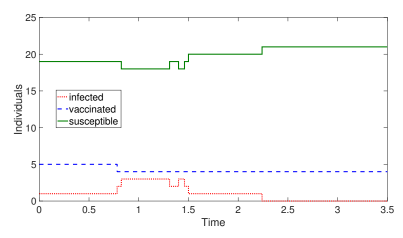


Fig. 3 Simulated trajectories of X when $N=25, \beta=2.5, \xi=0, h=0.3$ and $\gamma=1.0$

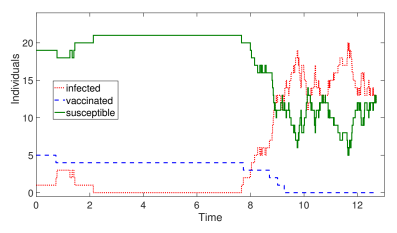


Fig. 4 Simulated trajectories of X when $N=25,\,\beta=2.5,\,\xi=0.01,\,h=0.3$ and $\gamma=1.0$

Figures 3 and 4 show typical trajectories for the CTMC X, starting from a single individual infected in a population of 25 individuals, 5 of which are vaccinated. We simulate 500 transitions among states and keep track of the time at which a transition occurs. Each line in Figures 3 and 4 represents the sequence of time points and the number of individuals (infected, vaccinated or susceptible) recorded at these time epochs. The paths in Figure 3 correspond to an isolated population and we observe that, approximately, after 2 time units the epidemic transmission will cease. In Figure 4, the reintroduction parameter

is $\xi = 0.01$. The path for vaccinated individuals shows that vaccine protection has faded away in about 10 time units, while the paths for the infected and susceptible populations settle towards the stationary endemic scenario.

3 Disease spread

This section deals with a probabilistic characterization of the offspring distribution of secondary infections of an epidemic process, at a particular time. We will consider two random variables to measure the spread of the infection, when a first case of disease is identified in a population.

In epidemiology, the individual who first brings the disease into a group is called the *index case* and we will start focusing on the number of infections coming from the index case during its entire infectious period. Secondly, we will study all of the secondary cases produced by the whole set of currently infectious individuals prior to the first recovery. The above mentioned measures are the stochastic analogues to the well-known basic reproduction number R_0 and, more specifically, to the control reproduction number R_c , defined as the average number of secondary cases due to each infective individual in the presence of control measures. In the case of vaccination R_0 and R_c satisfy the simple expression $R_c = R_0(1 - (1 - h)f)$, where h is the proportion of vaccinated individuals for which the vaccine has no effect (hence, 1 - h quantifies the vaccine effectiveness), and f represents the vaccine coverage as the initial fraction of the target population that has received the vaccine (Alexander et al. 2004; Magpantay 2014).

The Markovian chain, describing the evolution of the epidemic in the compartmental model, will play an essential role to develop theoretical results. The evaluation of the probability mass distribution of the number of secondary infections produced by a selected individual was firstly introduced by Ross (2011) and, in an independent way, by Artalejo and Lopez-Herrero (2013). Both papers consider finite SIS and SIR compartmental models and present probabilities of the offspring distribution as the solution of systems of linear equations. Their results generalize the study of the infection transmission from the instant of the invasion to any stage of the disease progression and show that, for populations of moderate size, the basic reproduction number can overestimate the potential transmission of the epidemic process.

The analysis of the transmission of an epidemic process has been extended to epidemic models with vector-borne infections (Artalejo 2014), heterogeneous contacts (Economou et al. 2015; López-García 2016), infective and susceptible immigrants (Almaraz et al. 2016), latency periods (Lopez-Herrero 2017), generally distributed infectious periods (Gómez-Corral and López-García 2017), Markov-modulated interactions (Almaraz and Gómez-Corral 2018) or to models for two-species competition (Gómez-Corral and López-García 2015).

Our results will reveal the influence of the vaccine, coverage and effectiveness, and the reintroduction parameter on the transmission potential of the

disease. Hence, in the subsequent sections we proceed with theoretical discussions leading to probabilistic results.

3.1 The exact transmission variable R_{e0}

In this section we study the potential transmission of an infective process by studying the random variable R_{e0} , defined as the number of infective individuals that arise from contagions caused directly by the index case, that is, the first individual in the population able to spread the disease. The objective is to characterize the distribution of the random variable R_{e0} and to observe the influence of the initial group of vaccinated individuals (i.e., vaccine coverage) on the transmission.

This analysis is directly related to the infectious period of the index case. During this period, the underlying Markov chain X evolves in \widehat{S} , the subset of states showing at least one infectious individual. For practical purposes \widehat{S} will be partitioned in levels, according to the current number of vaccinated individuals, as follows:

$$\widehat{S} = \bigcup_{v=0}^{v_0} \widehat{L}(v) = \bigcup_{v=0}^{v_0} \{(v,i) : 1 \leq i \leq N-v\}.$$
 First, we introduce appropriate notations for the generating and probability

First, we introduce appropriate notations for the generating and probability mass functions, and for the factorial moments of R_{e0} conditioned to a specific state $(v, i) \in \hat{S}$.

$$\varphi_{v,i}(z) = E\left[z^{R_{e0}} | (V(0) = v, I(0) = i)\right]$$

$$= \sum_{k=0}^{\infty} z^k P\left(\left\{R_{e0} = k | (V(0) = v, I(0) = i)\right\}\right), \text{ for } |z| \le 1,$$

$$x_{v,i}^k = P\left(\left\{R_{e0} = k | (V(0) = v, I(0) = i)\right\}\right), \text{ for } k \ge 0,$$

$$m_{v,i}^k = \begin{cases} 1 & \text{if } k = 0, \\ E\left[R_{e0} (R_{e0} - 1) \cdots (R_{e0} - k + 1) | (V(0) = v, I(0) = i)\right] & \text{if } k > 0. \end{cases}$$

In the rest of the section we will develop algorithmic schemes for determining $\varphi_{v,i}(z)$, $x_{v,i}^k$ and the moments $m_{v,i}^k$. Firstly, we mark the index case. The homogeneous mixing assumption guarantees all-to-all interactions with no preferences in relationship among individuals. Therefore, individuals are all the same and we split transitions in Table 2, associated to contagions, by distinguishing whether or not a new infection comes from the index case. That is, we partition contagion rates in the following way

$$\left(\frac{\beta i}{N} + \xi\right)(N - v - i) = \frac{\beta}{N}(N - v - i) + \left(\frac{\beta(i - 1)}{N} + \xi\right)(N - v - i),$$
$$h\left(\frac{\beta i}{N} + \xi\right)v = h\frac{\beta}{N}v + h\left(\frac{\beta(i - 1)}{N} + \xi\right)v.$$

Then, for $(v,i) \in \widehat{S}$, we define new rates $\alpha_{v,i}^*$, $\widetilde{\alpha}_{v,i}$, $\widetilde{\beta}_v^*$ and $\widetilde{\beta}_{v,i}$ as follows.

$$\alpha_{v,i}^* = \frac{\beta}{N}(N - v - i),\tag{2}$$

$$\widetilde{\alpha}_{v,i} = \left(\frac{\beta(i-1)}{N} + \xi\right)(N - v - i),\tag{3}$$

$$\beta_v^* = h \frac{\beta}{N} v, \tag{4}$$

$$\widetilde{\beta}_{v,i} = h\left(\frac{\beta(i-1)}{N} + \xi\right)v.$$
 (5)

A first-step argument, conditioning on the possible transitions out of a fixed state $(v,i) \in \widehat{S}$, shows that generating functions $\varphi_{v,i}(z)$ satisfy the following set of linear equations:

$$\varphi_{v,i}(z) = \frac{\gamma}{q_{v,i}} + (1 - \delta_{v,0}) \left(\frac{\beta_v^*}{q_{v,i}} z \varphi_{v-1,i+1}(z) + \frac{\widetilde{\beta}_{v,i}}{q_{v,i}} \varphi_{v-1,i+1}(z) \right)
+ (1 - \delta_{i,1}) \frac{\gamma(i-1)}{q_{v,i}} \varphi_{v,i-1}(z)
+ (1 - \delta_{i,N-v}) \left(\frac{\alpha_{v,i}^*}{q_{v,i}} z \varphi_{v,i+1}(z) + \frac{\widetilde{\alpha}_{v,i}}{q_{v,i}} \varphi_{v,i+1}(z) \right),$$
(6)

where $\delta_{i,j}$ represents the Kronecker's delta function, defined as 1 when i = j, and 0 otherwise.

Equation (6) is the basis to get the mass function of the conditional random variable $R_{e0} | (V(0) = v, I(0) = i)$ by numerical inversion, which can be done with the help of a Fast Fourier Transform (FFT) algorithm (Tijms 2003). As mass functions will be obtained through a direct recursive scheme, we do not go further on this point and we proceed to get factorial moments $m_{v,i}^k$, for $k \geq 1$.

By differentiating equation (6) with respect to z repeatedly k times $(k \ge 1)$ and evaluating at z = 1, we obtain the equations involving factorial moments conditioned to states $(v, i) \in \hat{S}$.

$$m_{v,i}^{k} = (1 - \delta_{v,0}) \left(\frac{\beta_{v}^{*}}{q_{v,i}} m_{v-1,i+1}^{k} + \frac{\widetilde{\beta}_{v,i}}{q_{v,i}} m_{v-1,i+1}^{k} \right) + (1 - \delta_{i,1}) \frac{\gamma(i-1)}{q_{v,i}} m_{v,i-1}^{k}$$

$$+ (1 - \delta_{i,N-v}) \left(\frac{\alpha_{v,i}^{*}}{q_{v,i}} m_{v,i+1}^{k} + \frac{\widetilde{\alpha}_{v,i}}{q_{v,i}} m_{v,i+1}^{k} \right)$$

$$+ (1 - \delta_{v,0}) k \frac{\beta_{v}^{*}}{q_{v,i}} m_{v-1,i+1}^{k-1} + (1 - \delta_{i,N-v}) k \frac{\alpha_{v,i}^{*}}{q_{v,i}} m_{v,i+1}^{k-1}.$$

$$(7)$$

Using relationships appearing in (2)-(5), we can write a useful and simplified version of (7),

$$q_{v,i}m_{v,i}^k = (1 - \delta_{v,0})h\left(\frac{\beta i}{N} + \xi\right)vm_{v-1,i+1}^k + (1 - \delta_{i,1})\gamma(i-1)m_{v,i-1}^k$$

$$+(1 - \delta_{i,N-v}) \left(\frac{\beta i}{N} + \xi\right) (N - v - i) m_{v,i+1}^{k}$$

$$+(1 - \delta_{v,0}) k \beta_{v}^{*} m_{v-1,i+1}^{k-1} + (1 - \delta_{i,N-v}) k \alpha_{v,i}^{*} m_{v,i+1}^{k-1}.$$
(8)

We notice that equation (8) provides conditional moments of order k based on conditional moments of one order less.

Moreover, at any level $0 \le v \le v_0$, the system of equations described in (8), for $1 \le i \le N - v$ and $k \ge 0$, can be written in matrix form as follows

$$\mathbf{m}_{\mathbf{v}}^{\mathbf{0}} = \mathbf{e}_{\mathbf{v}},\tag{9}$$

$$\mathbf{A}_{\mathbf{v}}\mathbf{m}_{\mathbf{v}}^{\mathbf{k}} = -\left(\left(1 - \delta_{v,0}\right)\left(\mathbf{D}_{\mathbf{v}}\widehat{\mathbf{m}}_{\mathbf{v}-1}^{\mathbf{k}} + \mathbf{D}_{\beta\mathbf{v}}\widehat{\mathbf{m}}_{\mathbf{v}-1}^{\mathbf{k}-1}\right) + k\mathbf{D}_{\alpha\mathbf{v}}\widetilde{\mathbf{m}}_{\mathbf{v}}^{\mathbf{k}-1}\right), \quad (10)$$

where the auxiliary matrices and vectors involved in (9)-(10) are defined below. For $0 \le v \le v_0$, **e.**, is an all-ones vector of dimension (N-v) and **A**, will

For $0 \le v \le v_0$, $\mathbf{e_v}$ is an all-ones vector of dimension (N-v) and $\mathbf{A_v}$ will denote the square (N-v) matrix with non null entries

$$A_{v}(i,j) = \begin{cases} (i-1)\gamma & \text{if } j = i-1, 2 \le i \le N-v, \\ -q_{v,i} & \text{if } j = i, 1 \le i \le N-v, \\ \left(\frac{\beta i}{N} + \xi\right)(N-v-i) & \text{if } j = i+1, 1 \le i \le N-v-1. \end{cases}$$

In addition, $\mathbf{D_v}$, $\mathbf{D_{\beta v}}$ and $\mathbf{D_{\alpha v}}$ represent (N-v) diagonal matrices, defined as follows:

$$\begin{aligned} \mathbf{D_{v}} &= Diag\left(h\left(\frac{\beta i}{N} + \xi\right)v : 1 \leq i \leq N - v\right), \\ \mathbf{D_{\beta v}} &= Diag\left(\beta_{v}^{*} : 1 \leq i \leq N - v\right), \\ \mathbf{D_{\alpha v}} &= Diag\left(\alpha_{v,i}^{*} : 1 \leq i \leq N - v\right). \end{aligned}$$

Finally, for $k \geq 0$ and $0 \leq v \leq v_0$, $\mathbf{m}_{\mathbf{v}}^{\mathbf{k}}$ and $\widetilde{\mathbf{m}}_{\mathbf{v}}^{\mathbf{k}}$ are (N-v)-dimensional vectors defined as

$$\mathbf{m_{v}^{k}} = (m_{v,1}^{k}, ..., m_{v,N-v}^{k})^{T},$$
$$\widetilde{\mathbf{m}_{v}^{k}} = (m_{v,2}^{k}, ..., m_{v,N-v}^{k}, 0)^{T},$$

and $\widehat{\mathbf{m}}_{\mathbf{v}}^{\mathbf{k}}$ is an (N-v-1)-dimensional vector given by $\widehat{\mathbf{m}}_{\mathbf{v}}^{\mathbf{k}} = \left(m_{v,2}^{k},...,m_{v,N-v}^{k}\right)^{T}$, where notation T denotes transposition.

Given an integer k, factorial moments are recursively determined with the help of the algorithmic scheme shown in Algorithm 1.

Remark 1 Notice that the random variable R_{e0} measures the exact number of secondary infective individuals arising directly from the index case and, in contradistinction with R_0 or R_c that are defined at the time of the invasion, R_{e0} can be checked at all times. If we set t=0 for the time at which the invasion starts the initial situation is $V(0)=v_0$ and I(0)=1, and $\overline{R}_{e0}=E\left[R_{e0}\left|V(0)=v_0,I(0)=1\right]\right]$ provides the exact amount of expected disease transmission, considering vaccine characteristics such as coverage and effectiveness.

Algorithm 1 R_{e0} factorial moments

For any k>0, factorial moments $\mathbf{m}_{\mathbf{v}}^{\mathbf{k}}=\left(m_{v,1}^{k},...,m_{v,N-v}^{k}\right)^{T}$, with $0\leq v\leq v_{0}$, are computed as follows:

Step 1. Set j = 0.

Step 1a. For $0 \le v \le v_0$, set $\mathbf{m_v^0} = \mathbf{e_v}$.

Step 2. Set j = 1.

Step 2a. Set v = 0, and

$$\mathbf{w_0^j} = -j\mathbf{D}_{\alpha \mathbf{0}}\widetilde{\mathbf{m}_0^{j-1}}.$$

Step 2b. Compute $\mathbf{m_0^j} = \mathbf{A_0^{-1} w_0^j}$. Set v = 1. If $v_0 = 0$, go to Step 3.

Step 2c. Set

$$\mathbf{w_v^j} = -\left(\mathbf{D_v}\widehat{\mathbf{m}_{\mathbf{v-1}}^j} + \mathbf{D_{\beta v}}\widehat{\mathbf{m}_{\mathbf{v-1}}^{j-1}} + k\mathbf{D_{\alpha v}}\widetilde{\mathbf{m}_{\mathbf{v}}^{j-1}}\right).$$

Step 2d. Compute $\mathbf{m}_{\mathbf{v}}^{\mathbf{j}} = \mathbf{A}_{\mathbf{v}}^{-1} \mathbf{w}_{\mathbf{v}}^{\mathbf{j}}$. Set v = v + 1. If $v \leq v_0$, go to Step 2c. Step 3. Set j = j + 1. If $j \leq k$, go to Step 2a.

Let us proceed to the analytic derivation of the probabilities $x_{v,i}^k$ that the index case will originate $k \geq 0$ new infections, given that the current situation is $(v,i) \in \widehat{S}$. A new appeal to the first-step methodology, by observing transitions out of the state (v,i), provides the following set of recursive equations:

$$q_{v,i}x_{v,i}^{k} = \delta_{k,0}\gamma + (1 - \delta_{k,0})\left((1 - \delta_{v,0})\beta_{v}^{*}x_{v-1,i+1}^{k-1} + (1 - \delta_{i,N-v})\alpha_{v,i}^{*}x_{v,i+1}^{k-1}\right) + (1 - \delta_{v,0})\widetilde{\beta}_{v,i}x_{v-1,i+1}^{k} + (i - 1)\gamma x_{v,i-1}^{k} + (1 - \delta_{i,N-v})\widetilde{\alpha}_{v,i}x_{v,i+1}^{k}.$$

$$(11)$$

For $k \geq 0$ and $0 \leq v \leq v_0$, equation (11) for states in L(v) can be written in matrix form in the following way:

$$\mathbf{B}_{\mathbf{v}}\mathbf{x}_{\mathbf{v}}^{\mathbf{0}} = -\gamma \mathbf{e}_{\mathbf{v}} - (1 - \delta_{v,0}) \mathbf{C}_{\mathbf{v}} \hat{\mathbf{x}}_{\mathbf{v}-1}^{\mathbf{0}},$$

$$\mathbf{B}_{\mathbf{v}}\mathbf{x}_{\mathbf{v}}^{\mathbf{k}} = -(1 - \delta_{v,0}) \left(\mathbf{C}_{\mathbf{v}} \hat{\mathbf{x}}_{\mathbf{v}-1}^{\mathbf{k}} + \mathbf{D}_{\beta \mathbf{v}} \hat{\mathbf{x}}_{\mathbf{v}-1}^{\mathbf{k}-1} \right) - \mathbf{D}_{\alpha \mathbf{v}} \tilde{\mathbf{x}}_{\mathbf{v}}^{\mathbf{k}-1}, k > 0,$$
(12)

where notations $\mathbf{e}_{\mathbf{v}}$, $\mathbf{D}_{\beta\mathbf{v}}$ and $\mathbf{D}_{\alpha\mathbf{v}}$ represent the same algebraic structures as defined in the lines following equations (9)-(10).

The remaining algebraic structures appearing in matrix expressions (12) are described next: $\mathbf{B_v}$ and $\mathbf{C_v}$ are (N-v) square matrices with entries

$$B_{v}(i,j) = \begin{cases} (i-1)\gamma & \text{if } j = i-1, 2 \leq i \leq N-v, \\ -q_{v,i} & \text{if } j = i, 1 \leq i \leq N-v, \\ \widetilde{\alpha}_{v,i} & \text{if } j = i+1, 1 \leq i \leq N-v-1, \\ 0 & \text{otherwise,} \end{cases}$$

$$\mathbf{C}_{\mathbf{v}} = Diag\left(\widetilde{\beta}_{v,i} : 1 \leq i \leq N-v\right),$$

and the vectors $\mathbf{x}_{\mathbf{v}}^{\mathbf{k}}$, $\widehat{\mathbf{x}}_{\mathbf{v}}^{\mathbf{k}}$ and $\widetilde{\mathbf{x}}_{\mathbf{v}}^{\mathbf{k}}$, containing probabilities $x_{v,i}^k$, are

$$\mathbf{x}_{\mathbf{v}}^{\mathbf{k}} = \left(x_{v,1}^{k}, ..., x_{v,N-v}^{k}\right)^{T},$$

$$\begin{split} \widehat{\mathbf{x}}_{\mathbf{v}}^{\mathbf{k}} &= \left(x_{v,2}^{k}, ..., x_{v,N-v}^{k}\right)^{T}, \\ \widetilde{\mathbf{x}}_{\mathbf{v}}^{\mathbf{k}} &= \left(x_{v,2}^{k}, ..., x_{v,N-v}^{k}, 0\right)^{T}. \end{split}$$

Notice that $P(\lbrace R_{e0} < \infty | V(0) = v, I(0) = i \rbrace) = 1$, for $(v, i) \in \widehat{S}$, because \widehat{S} is a finite union of disjoint finite sets, amd in a finite population the number of infective individuals arising from contagions caused by the index case is

necessarily finite. Consequently, $\sum_{k=0}^{\infty} x_{v,i}^{k} = 1$, for $(v,i) \in \widehat{S}$. For every number of contagions $k \geq 0$ the equations in (12) are solved recursively with the help of Algorithm 2. In order to determine mass distribution functions, a stopping criteria should be provided to avoid longer computation runs. In fact, numerical results appearing in Section 4 come from applying both recursive Algorithms 1 and 2. More specifically, those results dealing with R_{e0} -distributions for a given initial coverage value, v_0 , are obtained from iterating Algorithm 2 until 95% of the values of the distribution of $(R_{e0} | V(0) = v_0, I(0) = 1)$ is accumulated.

Algorithm 2 R_{e0} probabilities

For a fixed integer $k \geq 0$, the set of conditional probabilities $x_{v,i}^k$, for $(v,i) \in \widehat{S}$, can be computed according to the following scheme:

Step 1. Set v = 0.

Step 1a. Set j = 0 and compute $\mathbf{x_0^0} = -\gamma \mathbf{B_0^{-1}} \mathbf{e_0}$.

Step 1b. Set j = j + 1. While $j \le k$, compute $\mathbf{x_0^j} = -\mathbf{B_0^{-1}} \mathbf{D}_{\alpha 0} \widetilde{\mathbf{x}_0^{j-1}}$.

Step 2. Set v = 1. If $v_0 = 0$, stop.

Step 2a. Set j = 0 and compute $\mathbf{x}_{\mathbf{v}}^{\mathbf{0}} = -\mathbf{B}_{\mathbf{v}}^{-1} \left(\gamma \mathbf{e}_{\mathbf{v}} + \mathbf{C}_{\mathbf{v}} \widehat{\mathbf{x}}_{\mathbf{v}-1}^{\mathbf{0}} \right)$.

Step 2b. Set j = j + 1. While $j \le k$, compute

$$\mathbf{x}_{\mathbf{v}}^{j} = -\mathbf{B}_{\mathbf{v}}^{-1} \left(\mathbf{C}_{\mathbf{v}} \widehat{\mathbf{x}}_{\mathbf{v}-1}^{j} + \mathbf{D}_{\beta \mathbf{v}} \widehat{\mathbf{x}}_{\mathbf{v}-1}^{j-1} + \mathbf{D}_{\alpha \mathbf{v}} \widetilde{\mathbf{x}}_{\mathbf{v}}^{j-1} \right).$$

Step 3. Set v = v + 1. If $v \le v_0$, go to step 2a.

3.2 The population transmission variable

Another measure of the expansion of a contagious disease is R_p , which provides the global spread of the disease by counting all the infections that take place within the population, no matter who is the infectious spreader, before the first recovery occurs. R_p is a random variable that can be analyzed either at the beginning of the outbreak or at any later time, by updating the population situation in terms of the current state of the CTMC X.

The aim of this section is to describe the probabilistic behavior of R_p and to study the influence of the model parameters on global infection transmission. In particular, we are interested in comparing R_c , the control reproduction number of this model, with the expected value $\overline{R}_p = E[R_p | (V(0) = v_0, I(0) = v_0)]$ 1) for different scenarios.

First, we derive theoretical results involving the probability distribution and factorial moments of R_p . As in the preceding section, the central tool for our results will be the first-step methodology. But to avoid repetitive arguments we provide results in a comprehensive manner, leaving out unnecessary details.

Let $\psi_{v,i}(z)$ be the generating function of R_p , given that the current state of X is $(v,i) \in \widehat{S}$, with factorial moments $M_{v,i}^k$ for $k \geq 1$ ($M_{v,i}^0 = 1$). At any point z, with $|z| \leq 1$, generating functions are the solution of the following tridiagonal set of linear equations:

$$\psi_{v,i}(z) = \frac{\gamma i}{q_{v,i}} + (1 - \delta_{v,0}) \frac{h\left(\frac{\beta i}{N} + \xi\right) v}{q_{v,i}} z \psi_{v-1,i+1}(z) + (1 - \delta_{i,N-v}) \frac{\left(\frac{\beta i}{N} + \xi\right) (N - v - i)}{q_{v,i}} z \psi_{v,i+1}(z),$$
(13)

for $0 \le v \le v_0$, $1 \le i \le N - v$.

The k-th factorial moments of states $(v, i) \in \widehat{S}$ are the solution of a system of linear equations, arising from (13) as usual by taking derivatives with respect to z followed by an evaluation for z = 1, that is expressed in matrix form as follows:

$$\mathbf{H}_{\mathbf{v}}\mathbf{M}_{\mathbf{v}}^{\mathbf{k}} = -(1 - \delta_{v,0})\mathbf{D}_{\mathbf{v}}\left(k\widehat{\mathbf{M}}_{\mathbf{v}-\mathbf{1}}^{\mathbf{k}-\mathbf{1}} + \widehat{\mathbf{M}}_{\mathbf{v}-\mathbf{1}}^{\mathbf{k}}\right) - k\widetilde{\mathbf{D}}_{\mathbf{v}}\widetilde{\mathbf{M}}_{\mathbf{v}}^{\mathbf{k}-\mathbf{1}},\tag{14}$$

for $k \geq 1$ and $0 \leq v \leq v_0$. Here $\mathbf{H_v}$ and $\widetilde{\mathbf{D}_v}$ are (N-v) square matrices defined by:

$$\begin{split} H_v(i,j) &= \begin{cases} -q_{v,i} & \text{if } j=i, 1 \leq i \leq N-v, \\ \left(\frac{\beta i}{N} + \xi\right) \left(N-v-i\right) & \text{if } j=i+1, 1 \leq i \leq N-v-1, \\ 0 & \text{otherwise.} \end{cases} \\ \widetilde{\mathbf{D}}_{\mathbf{v}} &= Diag\left(\left(\frac{\beta i}{N} + \xi\right) \left(N-v-i\right) : 1 \leq i \leq N-v\right). \end{split}$$

Vectors M_v^k , \widehat{M}_v^k and \widetilde{M}_v^k related to factorial moments are defined as follows

$$\begin{aligned} \mathbf{M_{v}^{k}} &= \left(M_{v,1}^{k}, ..., M_{v,N-v}^{k}\right)^{T}, \\ \widehat{\mathbf{M_{v}^{k}}} &= \left(M_{v,2}^{k}, ..., M_{v,N-v}^{k}\right)^{T}, \\ \widehat{\mathbf{M_{v}^{k}}} &= \left(M_{v,2}^{k}, ..., M_{v,N-v}^{k}, 0\right)^{T}. \end{aligned}$$

The distribution of the random variable R_p , conditioned to any state $(v,i) \in \widehat{S}$, can be obtained by inverting transforms with the help of the recursive equations (13) and an FFT algorithm. But, as it was stated in the preceding section, it is possible to find a set of equations whose solution provides directly the point mass function of R_p : $z_{v,i}^k = P(\{R_p = k | V(0) = v, I(0) = i\})$,

when $(v,i) \in \widehat{S}$, for $0 \le k \le N-1$. Notice that when we observe i infected individuals in the population, then the number of secondary cases taking place before the first recovery is at most N-i. So, for any integer k, with $0 \le k \le N-1$, $z_{v,i}^k = 0$, whenever i > N-k.

Next, we introduce appropriate vectors to derive probabilities $z_{v,i}^k$ when $(v,i) \in \widehat{S}$, that is:

$$\begin{split} \mathbf{z_{v}^{k}} &= \left(z_{v,1}^{k}, ..., z_{v,N-v}^{k}\right)^{T}, \\ \widehat{\mathbf{z}_{v}^{k}} &= \left(z_{v,2}^{k}, ..., z_{v,N-v}^{k}\right)^{T}, \\ \widetilde{\mathbf{z}_{v}^{k}} &= \left(z_{v,2}^{k}, ..., z_{v,N-v}^{k}, 0\right)^{T}. \end{split}$$

Finally, for each level v, with $0 \le v \le v_0$, R_p point probabilities come from the following recursive equations:

$$\mathbf{D}_{\mathbf{Q}}\mathbf{z}_{\mathbf{v}}^{\mathbf{0}} = \mathbf{d}_{\gamma},\tag{15}$$

$$\mathbf{D}_{\mathbf{Q}}\mathbf{z}_{\mathbf{v}}^{\mathbf{k}} = (1 - \delta_{v,0})\mathbf{D}_{\mathbf{v}}\widehat{\mathbf{z}}_{\mathbf{v}-1}^{\mathbf{k}-1} + \widetilde{\mathbf{D}}_{\mathbf{v}}\widetilde{\mathbf{z}}_{\mathbf{v}}^{\mathbf{k}-1}, \text{ for } 1 \le k \le N - 1,$$
 (16)

where $\mathbf{D}_{\mathbf{Q}}$ and \mathbf{d}_{γ} are a diagonal matrix and a vector, with respective entries $q_{v,i}$ and γi , for $1 \leq i \leq N - v$.

As in Section 3.1, R_p factorial moments and probabilities can be computed from recursive schemes based on equations (14) and (15 - 16), respectively. Their algorithmic descriptions are similar to Algorithms 1 - 2 and are not stated in the text, but they are the basis for obtaining R_p 's numerical results appearing in Section 4.

Remark 2 In the paper by Artalejo and Lopez-Herrero (2014), the authors presented closed form expressions for probabilities dealing with the population transmission random variable in stochastic SIS and SIR models. For our SIV model, the mathematics is more involved due to the external transmission parameter and, unfortunately, it is not possible to derive general closed expressions for conditional probabilities $z_{v,i}^k$. However, after some algebra we obtained the closed values of point probabilities in a few specific situations.

$$z_{0,N}^k = P(\{R_p = k | V(0) = 0, I(0) = N\}) = \delta_{k,0}, \text{ for } 0 \le k \le N - 1.$$

For $0 \le v \le v_0$ and $1 \le i \le N - v$ we get:

$$\begin{split} z_{v,i}^0 &= P(\{R_p = 0 \,| V(0) = v, I(0) = i\}) = \frac{\gamma i}{q_{v,i}}, \\ z_{v,i}^1 &= P(\{R_p = 1 \,| V(0) = v, I(0) = i\}) \\ &= (1 - \delta_{i,N}) \frac{\left(\frac{\beta}{N}i + \xi\right)\gamma(i+1)}{q_{v,i}} \left(\frac{hv}{q_{v-1,i+1}} + \frac{(N-v-i)}{q_{v,i+1}}\right). \end{split}$$

4 Numerical illustrations

This section illustrates theoretical and algorithmic results derived in previous sections.

We fix the recovery rate as $\gamma = 1.0$ in all the experiments, so the time unit is taken as to be the expected time that an infected individual takes to recover to become susceptible again.

In the first scenario, we consider a population of N=100 individuals, 20% of which is partially protected against the infection by a vaccine with effectiveness of 97%. We are interested in the random variable R_{e0} , that is the number of secondary infections produced by the index case. Hence, the initial number of infective individuals is $I(0) = i_0 = 1$.

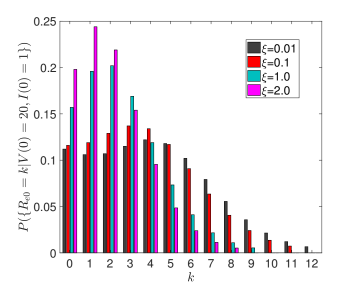


Fig. 5 R_{e0} distribution for several values of ξ , when N = 100, $\beta = 10.0$ and h = 0.03

Figures 5 and 6 represent histograms for the distribution of R_{e0} when we vary external or internal contact rates, respectively. Heights indicate the value of the probabilities $x_{20,1}^k = P(\{R_{e0} = k | V(0) = 20, I(0) = 1\})$ and colors appearing in both figures are depicted for a better distinction among considered situations. In more detail, Figure 5 shows mass functions of R_{e0} when the external transmission rate ξ is 0.01, 0.1, 1.0 and 2.0, for a fixed internal transmission rate $\beta = 10.0$. For each rate ξ , mean values of R_{e0} are 4.2035, 3.7601, 2.5342 and 2.0592, respectively. Moreover, distributions present a strictly positive mode that occurs with a probability that decreases for increasing values of the external rate. This remark is according to the intuition because for a fixed internal contact transmission rate β , when the external transmission rate in-

creases the index case has less opportunities to spread the disease compared with outsider infection sources.

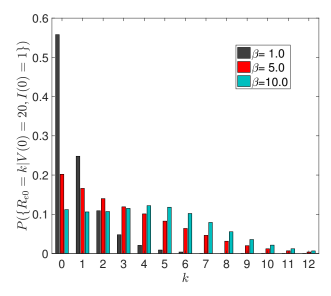


Fig. 6 R_{e0} distribution for several values of β , when N = 100, $\xi = 0.01$ and h = 0.03

Figure 6 displays mass functions corresponding to $\beta=1.0, 5.0$ and 10.0, for an external transmission rate $\xi=0.01$. We get decreasing shape functions for $\beta=1.0$ or $\beta=5.0$ and the chance that the index case produces no secondary infections is 55% and 20%, respectively. However, for $\beta=10.0$, the R_{e0} distribution is bimodal, with an 11.2% chance that the index case recovers before spreading the infection. In general terms, long internal transmission rates contribute a higher number of secondary cases.

Next we focus on the effect of the vaccine coverage on the expansion of the infection. Figure 7 shows the probability that the index case produces two or more secondary cases of infection, as a function of the external rate β . This quantity can give an idea of what is, for a given infective process, the chance of invading a susceptible-vaccinated population. Each curve corresponds to a different initial vaccine coverage. The remaining parameters of the model are $N=100,\,\xi=0.01,\,$ and the vaccine is efficient in 97% of the vaccinated individuals. Probability increases with β , no matter how large the initial coverage v_0 is. For a fixed external transmission rate, the chance of having at least two infections increases when initial coverage decreases. In particular for a small population of 100 individuals affected by a hard measles outbreak, with $R_0=\beta/\gamma=18.0,\,$ numerical results evince that a massive vaccination policy with two doses of MMR vaccine (measles, mumps, rubella) guarantees that the probability of having two or more measles infections from the index case

is 0.12, while in an unprotected population $(v_0 = 0)$ this probability grows up to 0.9.

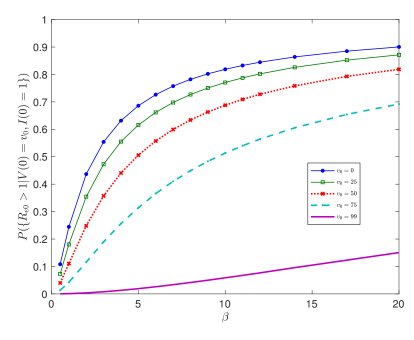


Fig. 7 $P(\{R_{e0} > 1 | V(0) = v_0, I(0) = 1\})$ as a function of the internal transmission rate when N = 100, $\xi = 0.01$ and h = 0.03

Now we deal with the expected number of secondary cases produced by the index case since their introduction in a population where individuals are either susceptible or vaccinated. We observe the influence of the model parameters β , ξ , h and v_0 on the expected value of R_{e0} . Results correspond to a population of N=100 individuals.

Figures 8-10 are contour graphs for $\overline{R}_{e0} = E[R_{e0} | V(0) = v_0, I(0) = 1]$, arising when we combine the influence of two parameters of the model. Different colors represent different values for \overline{R}_{e0} , as it is indicated by the color code given in each of the Figures.

Figure 8 shows the influence on \overline{R}_{e0} of the internal and external transmission rates, when 20% of the population has received a vaccine which is effective among 97% of the vaccinated individuals. The average of secondary infections, produced by the index case, increases with the internal transmission rate and it decreases when ξ increases, which is in agreement with the comment for Figure 5. These behaviors are more noticeable for large transmission intensities.

Figure 9 displays \overline{R}_{e0} as a function of the internal transmission rate, β , and the potential risk of vaccine failure, h. We assume in addition that 20% of the

population is vaccinated and that the rate of external transmission is $\xi = 0.01$. The expected number of infections caused directly by the index case increases with β . In the early spread of the epidemic, the influence of the vaccine failure risk is relatively small for outbreaks showing internal rates β smaller than 6 compared to those with higher values for β .

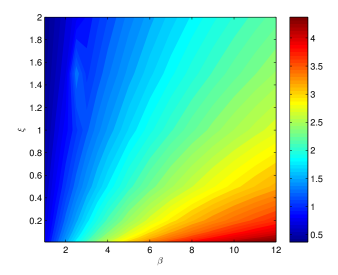


Fig. 8 \overline{R}_{e0} as a function of β and ξ when $N=100,\,h=0.03$ and $v_0=20$

Finally, the contour graph shown in Figure 10 presents the relationship of the internal transmission and vaccine coverage on \overline{R}_{e0} . The additional parameters are $N=100,\,\xi=0.01$ and 97% for vaccine effectiveness. An increase of vaccine coverage produces a decrease on the initial spread of the infection. For instance, administration of two doses of MMR vaccine is 97% effective against rubella (basic reproduction number $R_0\approx 6$). So, assuming that $R_0=\beta/\gamma$, results in Figure 10 show that the index case transmits an infection like rubella to an average of 3.5 people in an unprotected population, and to less than one person when a massive vaccination policy was launched in the population. To achieve that the infection will not propagate in the population, that is $R_{e0}<1$, higher immunization coverage is needed as β increases.

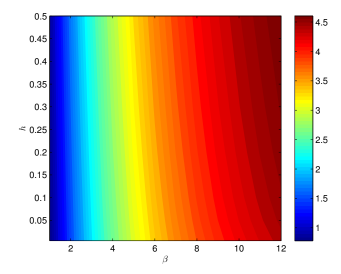


Fig. 9 \overline{R}_{e0} as a function of β and h when $N=100,\,\xi=0.01$ and $v_0=20$

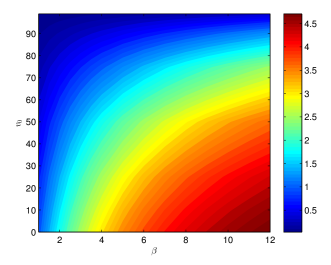


Fig. 10 \overline{R}_{e0} as a function of β and v_0 when N = 100, $\xi = 0.01$ and h = 0.03

The boxplot appearing in Figure 11 corresponds to the distribution of the number of secondary infections produced by the index case. The box encloses the middle central part of the distribution, lower and upper edges of the box

correspond to the lower and upper quartile, respectively, and the line drawn across the box indicates the median of the distribution; finally, whiskers above and below the box cover 95% of the distribution. The objective is to compare the patterns of the epidemic when we increase the vaccination coverage, in a population of N=100 individuals, assuming that a 97%-effective vaccine is available to control an epidemic process with internal and external transmission rates $\beta=10.0$ and $\xi=0.1$, respectively. This choice for model parameters corresponds to an infection by varicella-zoster virus (VZV) and the administration of varicella vaccine that is 97% effective in the first year after vaccination. A low vaccine coverage leads to a large number of secondary infections. In general terms, the number of secondary cases produced by the index case decreases when v_0 increases. Notice that changes on the number of secondary infections are not significant while the vaccination coverage is under 25. This fact shows the importance of the random variable R_{e0} to fix an appropriate vaccination level.

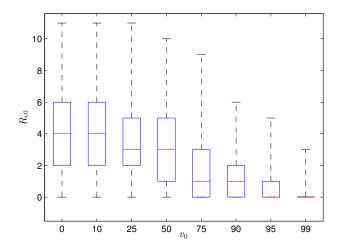


Fig. 11 Boxplot for R_{e0} under several vaccine coverage values when $N=100,\,\beta=10.0,\,\xi=0.1$ and h=0.03

Next we present some results relative to the global spread of the disease. In Figure 12, we compare R_c , \overline{R}_{e0} and \overline{R}_p when the internal transmission rate β varies over the interval (0, 20.0). As in previous scenarios, the recovery rate is $\gamma=1.0$. Vaccine coverage reaches half of the population and presents an effectiveness of 97%. We compare results for populations of N=100 and 1000 individuals, with external transmission rate $\xi=1/N$. For our parameter selection $R_0=\beta$ and $R_c=R_0(1-(1-h)/2)=0.515\beta$. Hence, as a function of β , the control reproduction number corresponds to the top line in Figure

12. We notice that differences between R_c and R_{e0} increase with increasing transmission rates. When $\beta=20.0$, we find $R_c=10.30$ and $\overline{R}_{e0}=4.36$ for N=100, and $\overline{R}_{e0}=5.67$ for N=1000. These magnitudes show the overestimation of the number of secondary cases of infection produced by the index case, as provided by R_c , and how the expected exact reproduction number \overline{R}_{e0} corrects the effect of the linearization hypothesis commonly assumed in the deterministic literature. Regarding the expected population transmission, \overline{R}_p , we conclude that it converges to the control reproduction number as the population size increases. In fact, for N=1000 differences between R_c and \overline{R}_p are smaller than 0.4 when $\beta \in (0,20.0)$. Hence, the line for \overline{R}_p is graphically indistinguishable from R_c and it is not plotted for N=1000.

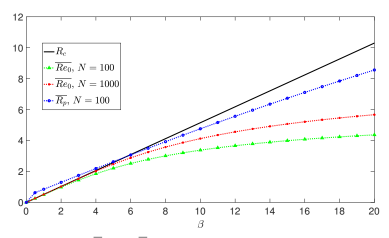


Fig. 12 Comparing R_c , \overline{R}_{e0} and \overline{R}_p versus the internal transmission rate when $\xi = 1/N$, h = 0.03 and $v_0 = N/2$

Finally, Table 3 shows the minimal vaccination coverage needed to interrupt epidemic transmission; that is, in order to get values smaller than one for the expected number of secondary cases coming from the index case. Notice that if the expected transmission (either from the whole set of infectious or limited to the index case) is less than 1, each infected individual transmits the disease to less than one person, which means that not every case will result in a new individual infection and epidemic transmission will cease. We display results for a population of N=1000 individuals, where the external transmission rate is taken as $\xi=1/N$ and the internal rate β is assumed to be at least 1. We compare two scenarios by considering that the effectiveness of a vaccine is either 97% or 90%. Values represented by v_c give the herd-immunity threshold based on the control reproduction number R_c (i.e., starting from $R_0=\beta/\gamma$, coverage is chosen as $v_c/N>(R_0-1)/(R_0(1-h))$ in order that $R_c<1$). Additionally, v_{e0} and \overline{R}_p , respectively, are less than one.

Numerical values come from an iterative application of Algorithm 1 for R_{e0} and its variant for R_p , by raising initial coverage until expected values fulfill the required condition. Any dash symbol in Table 3 means that even a 100% vaccine coverage does not guarantee that the expected number of secondary cases (measured in terms of R_c , \overline{R}_{e0} or \overline{R}_p) are less than one. Note that, as \overline{R}_p includes also secondary cases arising from external infectious individuals, the minimal critical level v_p presents higher values, at a fixed transmission rate β , when it is compared with v_c or v_{e0} . On the other hand, when the internal transmission rate β increases, higher immunization coverage is needed to keep the expansion of the epidemic under control.

Table 3	Minimum	level of	vaccination	with	parameters	N =	1000	and §	= 1	/N
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	h = 0.03			h = 0.1			
β	v_c	v_{e0}	v_p	v_c	v_{e0}	v_p	
1	0	0	413	0	0	445	
1.5	344	341	556	370	368	695	
2	515	513	645	556	554	808	
3	687	686	750	741	739	873	
4	773	772	809	833	832	915	
5	825	823	848	889	888	944	
6	859	858	875	926	925	966	
7	884	882	896	952	952	982	
8	902	902	911	972	972	996	
9	916	915	923	988	988	_	
10	928	927	933	_	_	_	
11	937	936	942	_	_	_	
12	945	944	949	_	_	_	

5 Conclusions

This paper studies infectious disease dynamics in a stochastic framework, where a Markov chain is used to model disease transmission. The continuous time Markov chain models changes in the state of the process defined as the number of individuals that are susceptible, infected or vaccinated. Assuming that susceptible and vaccinated individuals (due to vaccine failures) can get the infection through both internal and external contacts makes the model more realistic than those with only internal contacts.

Deterministic models are very useful in understanding the dynamics of infectious diseases and estimating important epidemiological descriptors as, for instance, the basic reproduction number. In this context, populations are relatively large, and the effect of depletion of susceptibles is minor but leads to reproductive numbers that overestimate the transmission potential of the disease. Our research involves a pair of random variables, R_{e0} and R_p , as alternative measures of the control reproduction number R_c . The stochastic

Markovian framework allows us to identify the effective event leading to a new contagion. Consequently, we exclude repeated contacts established between the index case and already infected individuals, thus correcting for the effect of the linearization assumption commonly assumed in the deterministic framework and which produces the overestimation of the reproductive potential of a disease.

Concerning speed, our numerical algorithms provide results depending strongly on the population size and the other parameters of the model. For instance, the derivation of \overline{R}_{e0} through Algorithm 1 involves factorial moments $m_{v,i}^1$ of $(N-v_0/2)(v_0+1)$ transient states, which for the particular choice of N=1000 and $v_0=N/2$ implies 375,750 such states and requires 118 sec CPU time, in a personal computer of 2.31GHz and 6GB RAM.

Numerical results regarding vaccination coverage levels should be seen as a first approach to investigate optimal policies for controlling the spread of an infectious disease. The aim for future research is to develop health policies based on time horizons and warning levels related to the remaining vaccinated individuals.

This research can be generalized to different models for imperfect vaccine response by considering vaccines that reduce the probability of infection or that confer protection that wanes over time (see, for instance, Ball et al. 2008, Ball and Sirl 2018). Additionally, more sophisticated models could be treated by introducing population structures such as households, and also by including assumptions concerning to epidemics showing latent infectious periods or lifetime immunity after recovery.

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