

Infection, reinfection, and vaccination under suboptimal immune protection: epidemiological perspectives

M. Gabriela M. Gomes^{a,*}, Lisa J. White^b, Graham F. Medley^b

^a Instituto Gulbenkian de Ciência, Apartado 14, 2781-901 Oeiras, Portugal

^b Ecology and Epidemiology Group, Department of Biological Sciences, University of Warwick, Coventry, CV4 7AL, UK

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Abstract

The *SIR* (susceptible–infectious–resistant) and *SIS* (susceptible–infectious–susceptible) frameworks for infectious disease have been extensively studied and successfully applied. They implicitly assume the upper and lower limits of the range of possibilities for host immune response. However, the majority of infections do not fall into either of these extreme categories. We combine two general avenues that straddle this range: temporary immune protection (immunity wanes over time since infection), and partial immune protection (immunity is not fully protective but reduces the risk of reinfection). We present a systematic analysis of the dynamics and equilibrium properties of these models in comparison to *SIR* and *SIS*, and analyse the outcome of vaccination programmes. We describe how the waning of immunity shortens inter-epidemic periods, and poses major difficulties to disease eradication. We identify a “reinfection threshold” in transmission when partial immunity is included. Below the reinfection threshold primary infection dominates, levels of infection are low, and vaccination is highly effective (approximately an *SIR* model). Above the reinfection threshold reinfection dominates, levels of infection are high, and vaccination fails to protect (approximately an *SIS* situation). This association between high prevalence of infection and vaccine failure emphasizes the problems of controlling recurrent infections in high-burden regions. However, vaccines that induce a better protection than natural infection have the potential to increase the reinfection threshold, and therefore constitute interventions with a surprisingly high capacity to reduce infection where reduction is most needed.

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1. Introduction

Mathematical epidemiological models for the dynamics of microparasite infections that induce lifelong immunity have been extensively developed (Kermack and McKendrick, 1927; Anderson and May, 1991; Grenfell et al., 2001) and used as predictive tools to assist in the design of control programmes (Osborne et al., 2000). This class of infections is viral and usually occurs during childhood (if there is no vaccination occurring), which is indicative of high transmissibility.

Such effective immunity is observed in infections such as measles, mumps and rubella (MMR) but this is unusual. More common is the occurrence of several reinfections throughout life. In similar vein to char-

acterization of a vaccine response (McLean and Blower, 1993), susceptibility to reinfection following a primary infection can be attributed to a combination of two factors: immune protection may wane over time (temporary immunity), or immunity may not be fully protective (partial immunity). Both effects are likely due to a combination of host insufficiency in acquiring and maintaining specific immunity (e.g. tuberculosis—Vynnycky and Fine, 1997; pertussis—Hethcote, 1999; van Boven et al., 2000) and of pathogen ability to generate antigenic diversity thereby avoiding immune recognition (e.g. influenza—Hay et al., 2001; Earn et al., 2002; respiratory syncytial virus—Cane, 2001).

The specific within-host mechanisms allowing recurrent infections are not explicit in this paper. Rather, we construct a series of simple models to investigate the epidemiological consequences of varying the duration and degree of immune protection. The models are systematically analysed and reveal two main outcomes:

*Corresponding author. Tel.: +351-214-464-626;

fax: +351-214-407-970.

E-mail address: ggomes@igc.gulbenkian.pt (M.G.M. Gomes).

the duration of immunity has a crucial impact on potential inter-epidemic periods; the degree of immune protection determines a reinfection threshold in transmission responsible for a steep increase in the prevalence of infection. The models are then extended to analyse the effects of temporary and partial immunity on the global impact of vaccination programmes.

Mathematical models have long been associated with the planning of vaccination programmes. The main contributions have been the prediction of the vaccination coverage necessary to eradicate infections (Nokes and Anderson, 1992, 1993), and the illustration of the post-vaccination dynamics if coverage does not meet the eradication threshold (McLean, 1995a). McLean and Blower (1993) have also investigated the consequences of vaccine failures, but under the assumption that immunity induced by natural infection was fully protective. Here this assumption is relaxed to accommodate the recurrence of infections. In the case of temporary protection, we reach the same basic conclusion that the waning of immunity is a major obstacle to disease eradication even if individuals are protected until very late in their lifetime. In the case of partial protection, our conclusions are fundamentally different due to reinfection threshold. Below the reinfection threshold, levels of infection are low and vaccination impact is high. The transmission dynamics are essentially described by the susceptible–infected–recovered (*SIR*) framework. Populations where transmissibility is above the reinfection threshold are of greater concern. They sustain very high endemicities and vaccination programmes are unable to increase natural immunity further. The transmission dynamics are essentially susceptible–infected–susceptible (*SIS*). The transition between the two regimes happens over a short range in transmission coefficient, indicating high sensitivity to changes.

Throughout this paper we assume that birth and death rates are equal ensuring a constant host population size, the duration of infection is typically two orders of magnitude lower than the host lifetime, and the periods of infection and infectiousness coincide. Vaccination programmes are implemented at birth. These assumptions simplify the model analysis and do not induce unwanted effects.

2. SIR and SIS models

The essence of infectious disease dynamics can be represented by simple systems of ordinary differential equations (Anderson and May, 1991; Diekmann and Heesterbeek, 2000). The host population is assumed homogeneous at birth and differentiation occurs as a result of infection experience. The population is assumed to mix homogeneously and transmission is

according to the mass-action principle. Hosts are divided into three proportions: susceptible (*S*), infectious (*I*), and recovered (*R*). Under the assumption that recovered individuals acquired some immunity that is totally protective and lifelong, we obtain the so-called *SIR* model formalized by the system of differential equations

$$\begin{aligned}\frac{dS}{dT} &= \mu - \beta IS - \mu S, \\ \frac{dI}{dT} &= \beta IS - (\tau + \mu)I, \\ \frac{dR}{dT} &= \tau I - \mu R.\end{aligned}\quad (1)$$

As in Table 1, the parameter μ is the death rate (and equally, the birth rate), τ is the rate of recovery from infection, and β is a transmission coefficient which combines a variety of epidemiological, environmental, and social factors that affect transmission. Susceptible individuals acquire infection at per capita rate βI . If time, T , is measured in years, hosts are born with a life expectancy of

$$L = \frac{1}{\mu} \text{ years}$$

and we assume throughout that $L=70$ years. The average duration of infection is

$$D = \frac{1}{\tau + \mu} \text{ years}$$

and we assume throughout that D is approximately 1 month. The last equation of model (1) can be omitted by recalling that $S + I + R = 1$.

2.1. Non-dimensional *SIR* model

The *SIR* model can be further simplified, resulting in a reduction in the number of parameters. Measuring time in units of duration of infection, $t = T/D$, we get the non-dimensional system (see Table 2)

$$\begin{aligned}\frac{dS}{dt} &= e - R_0 IS - eS, \\ \frac{dI}{dt} &= R_0 IS - I.\end{aligned}\quad (2)$$

The first parameter,

$$e = \frac{D}{L}$$

Table 1
Model parameters when time is measured in years

Symbol	Definition	Value
μ	Death rate and, equally, birth rate	1/70 year ⁻¹
τ	Rate of recovery from infection	12 year ⁻¹
β	Transmission coefficient	Variable

Table 2
Model parameters when time is measured in units of average duration of infection, $D=1/(\tau+\mu)$

Symbol	Definition	Value
e	Death and birth rate, $\mu/(\tau+\mu)$	$0.0012D^{-1}$
R_0	Basic reproduction number, $\beta/(\tau+\mu)$	Variable
α	Factor affecting the rate of loss of acquired immune protection	Between 0 and 1
σ	Factor affecting the degree of partial immunity induced by a previous infection	Between 0 and 1
σ_V	Factor affecting the degree of partial immunity induced by vaccination	Between 0 and 1
v	Vaccination coverage	90%

represents the average duration of an infectious episode, expressed as a proportion of average lifetime. For the values of L and D above we get the approximate value of $e=0.0012$. The second parameter,

$$R_0 = \beta D$$

represents the number of secondary cases expected from a primary case in a completely susceptible population. This is the so-called basic reproduction number, and is a crucial parameter that will be thoroughly explored. The *SIR* model with the new time units is represented diagrammatically in Fig. 1(a). The model has two possible steady states

- (1) Disease-free equilibrium: $S_1 = 1, I_1 = 0$.
 - (2) Endemic equilibrium: $S_2 = 1/R_0, I_2 = e(1-(1/R_0))$.
- The disease-free equilibrium, (S_1, I_1) , is a valid steady state independently of the value of R_0 , but the endemic equilibrium, (S_2, I_2) , requires $R_0 > 1$. The number of steady states changes at $R_0 = 1$, and this is a bifurcation point. The stability of (S_i, I_i) is determined by the eigenvalues of the Jacobian matrix

$$J_i = \begin{pmatrix} -R_0 I_i - e & -R_0 S_i \\ R_0 I_i & R_0 S_i - 1 \end{pmatrix}$$

for $i=1,2$. A straightforward calculation gives the following results:

- (1) Eigenvalues of J_1 : $-e$ and $R_0 - 1$.
- (2) Eigenvalues of J_2 : $\frac{-eR_0 \pm \sqrt{e^2 R_0^2 - 4e(R_0 - 1)}}{2}$.

The eigenvalues of J_1 are always real, and from their signs we infer that the steady state (S_1, I_1) is stable for $R_0 < 1$, and unstable for $R_0 > 1$. The eigenvalues of J_2 have negative real part for the whole range of validity of the steady state (S_2, I_2) , implying that this steady state is always stable. The eigenvalues are real for R_0 just above 1, and become complex for R_0 between $(2/e)(1 - \sqrt{1 - e})$ and $(2/e)(1 + \sqrt{1 - e})$ implying convergence in the form of damped oscillations in this range (for $e=0.0012$, this is $1.0003 < R_0 < 3332$). The vector field is illustrated in Fig. 2(a) for the particular case $R_0 = 3$. Superimposed is a simulation starting near the disease-free equilibrium illustrating how the disease invades and converges to the endemic equilibrium. Fig. 2(b) shows the time-series plot corresponding to the

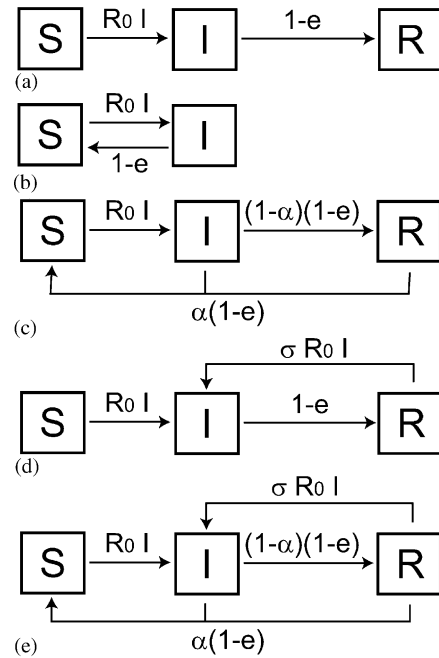


Fig. 1. Flow diagram for all the models analysed in this paper. Time is in units of average duration of infection and the parameters are described in Table 1. For simplicity, births and deaths are not represented.

same simulation. Convergence is in the form of damped oscillations: a series of epidemics of sequentially reducing amplitude occur until the system approaches the endemic equilibrium. A number of mechanisms have been identified as capable of sustaining the oscillations (Liu et al., 1987), enhancing the importance of quantifying the inter-epidemic periods.

2.2. Non-dimensional SIS model

Variations of the *SIR* model have been developed to incorporate features of particular diseases (Anderson and May, 1991). Here, we consider diseases that fail to elicit protective immunity, allowing recurrent infections. In this extreme case, recovered hosts return to the susceptibility class and the recovered class remains empty so that $S+I=1$. The dynamics of this system are driven by a one-dimensional system: the *SIS* model, represented diagrammatically in Fig. 1(b) and more

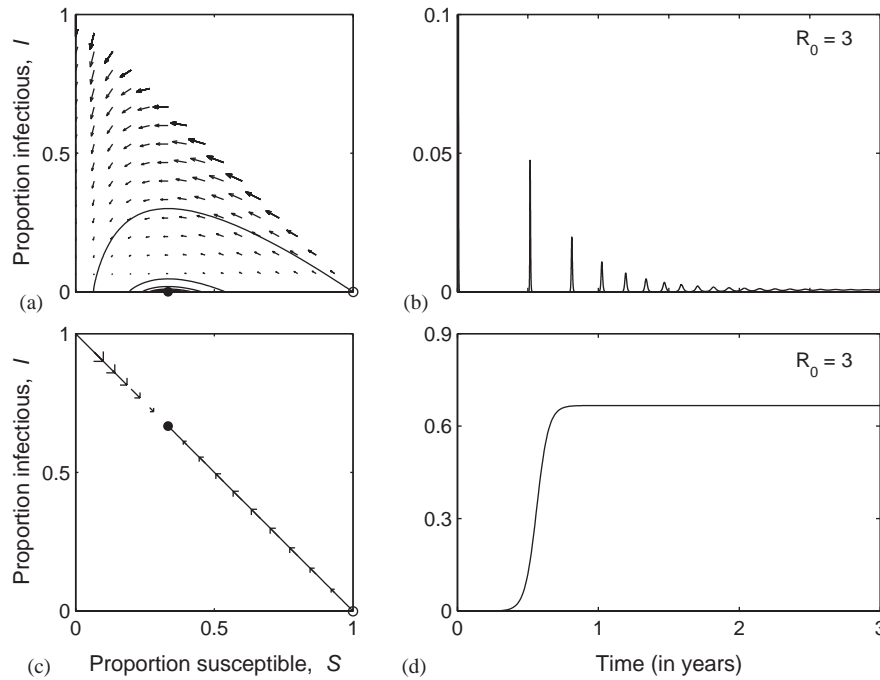


Fig. 2. Vector field and simulated solution corresponding to the *SIR* and *SIS* models for $R_0=3$. (a) two-dimensional vector field of *SIR* model, the two steady states (open circle is unstable, and full circle is stable), and a simulated solution with initial condition near the unstable disease-free state; (b) time series corresponding to the same simulated solution; (c) one-dimensional vector field of *SIS* model, the two steady states, and a simulated solution with initial condition near the unstable disease-free state; (d) time series corresponding to the same simulated solution.

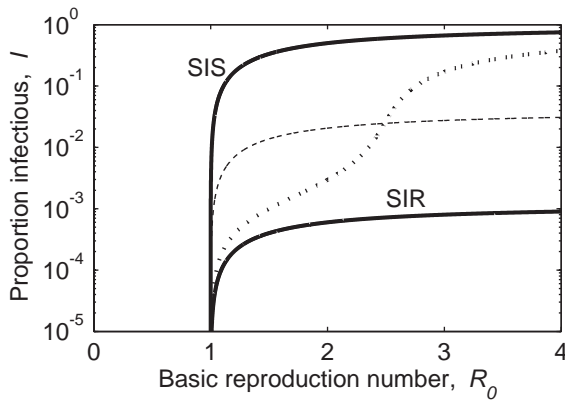


Fig. 3. Bifurcation diagram for the *SIR*, *SIS*, and intermediate models. The extreme *SIR* and *SIS* models are represented by the lower and higher full curves, respectively. Temporary immunity is represented by the dashed line, and partial immunity is represented by the dotted line. Note the logarithmic scale on the vertical axis.

formally by the equation

$$\frac{dI}{dt} = R_0 I(1 - I) - I. \tag{3}$$

Also here, there is a disease-free equilibrium ($S = 1, I = 0$) for all values of R_0 . Stability analysis shows that this steady state is stable for $R_0 < 1$ and unstable for $R_0 > 1$. As R_0 is increased across one, a branch of endemic equilibria bifurcates. At the endemic steady state we have $S = 1/R_0$ and $I = 1 - 1/R_0$. In Fig. 2(c) and (d), we fix $R_0 = 3$ to represent the vector field associated with the

SIS model and a simulation to illustrate convergence to the steady state. The bifurcation diagrams corresponding to both the *SIR* and *SIS* models are shown as the full lines in Fig. 3, setting the lower and upper bounds to the intermediate models.

3. Intermediate models

3.1. Temporary immune protection

Pertussis is a highly contagious infection of the respiratory tract where recurrence has been attributed to the waning of immunity (Hethcote, 1999; van Boven et al., 2000). Control by vaccination is less successful than for other childhood diseases, and recent trends are for overall increase, and occasionally strong epidemic outbreaks.

The temporary immunity model is represented diagrammatically in Fig. 1(c). Our assumption is that, upon infection, individuals develop an immune response that is lost at a certain rate. The parameter α is introduced to control the rate of loss of immunity. The model equations are

$$\begin{aligned} \frac{dS}{dt} &= e - R_0 I S - e S + \alpha(1 - e)(1 - S), \\ \frac{dI}{dt} &= R_0 I S - I. \end{aligned} \tag{4}$$

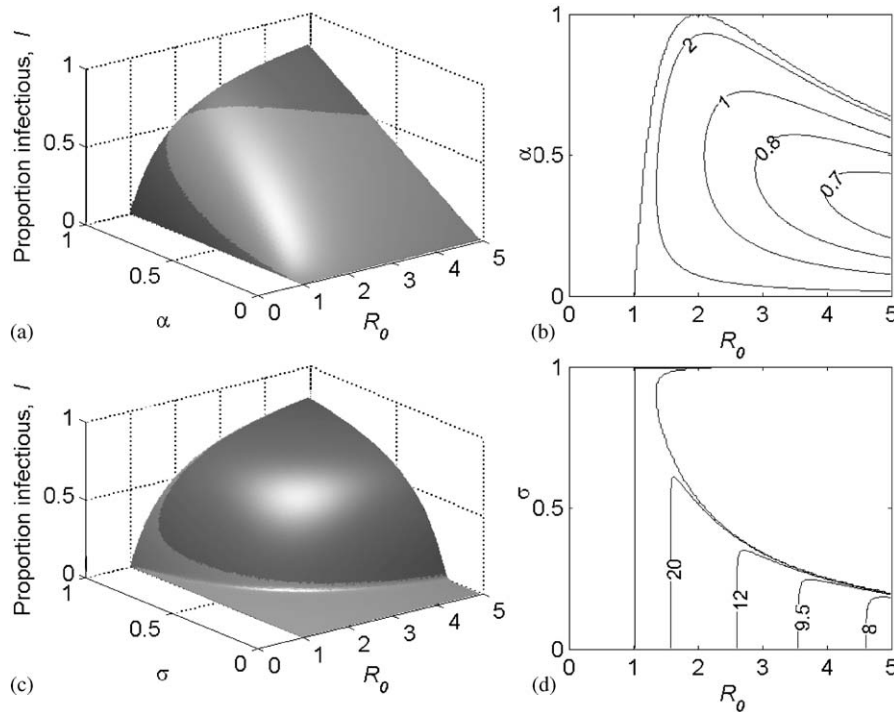


Fig. 4. Equilibria and convergence for the temporary immunity and partial immunity models. (a) and (b) refer to temporary immunity: (a) shows the endemic equilibrium as a function of R_0 and α indicating whether convergence is by damped oscillations (light grey) or linear decay (dark grey); and (b) shows contour plots for the period of the oscillations in the damped oscillatory region (the labels represent years). (c) and (d) provide the same information for partial immunity: (c) shows the endemic equilibrium as a function of R_0 and σ ; and (d) shows contour plots for the period of the oscillations.

The new parameter can take values between 0 and 1. In the limit $\alpha=0$, the rate of loss of immunity equals the death rate (*SIR* limit). In the limit $\alpha=1$, the rate of loss of immunity equals the rate of loss of infectiousness (*SIS* limit). The equilibrium curve for $\alpha=0.04$ is represented as a dashed line in Fig. 3. Fig. 4(a) shows the stable endemic equilibrium as a function of R_0 and α indicating whether convergence is by damped oscillations (light grey) or linear decay (dark grey). Fig. 4(b) shows the contour plots for the period of the oscillations in the damped oscillatory region (with time rescaled back to years). Naturally, waning of immunity has a crucial impact on the time-scale for potential oscillatory dynamics.

3.2. Partial immune protection

There are many infections motivating the study of partial immune protection. Influenza A and B viruses cause the same respiratory disease, but exhibit contrasting evolutionary features (Hay et al., 2001). Partial immunity across antigenic variants is a principal player in models designed to study influenza evolution (Andreasen et al., 1997; Ferguson et al., 2003; Gog and Grenfell, 2002; Gomes et al., 2004b). Other pathogens invoking partial immunity include *Neisseria meningitidis* (Gupta and Maiden, 2001), *Streptococcus pneumoniae*

(Lipsitch, 1997), and *Mycobacterium tuberculosis* (Vynnycky and Fine, 1997; Gomes et al., 2004a).

The partial immunity model is represented diagrammatically in Fig. 1(d) and formalized by the system of equations

$$\begin{aligned} \frac{dS}{dt} &= e - R_0IS - eS, \\ \frac{dI}{dt} &= R_0I(S + \sigma(1 - S - I)) - I. \end{aligned} \tag{5}$$

The new assumption is that individuals are protected while infected but regain some susceptibility upon recovery. This susceptibility is reduced by a factor σ , compared to susceptibility prior to infection. The limit $\sigma=0$ is an *SIR* model, and the limit $\sigma=1$ is an *SIS* model. The equilibrium curve for $\sigma=0.4$ is represented as a dotted line in Fig. 3. Fig. 4(c) shows the stable endemic equilibrium as a function of R_0 and σ indicating whether convergence is by damped oscillations (light grey) or linear decay (dark grey). Fig. 4(d) shows the contour plots for the period of the oscillations in the damped oscillatory region (in years). The figure reveals two clearly distinct types of endemic behaviour: low and potentially oscillatory; and high and steady. Roughly, the first requires that transmissibility is above the threshold for disease persistence ($R_0 > 1$), and the second

relies on reinfection requiring that transmissibility is above a higher threshold ($R_0 > 1/\sigma$).

3.3. Temporary-partial immunity

We used the dynamics of pertussis as an example of temporary immunity, and we evoked a number of infections as examples of partial immunity. However, the transmission dynamics of each of these infections is likely to be associated with a specific combination of both temporary and partial immunity processes, as recently analysed for respiratory syncytial virus (White et al., 2004). We construct a general framework where the contributions of the two mechanisms are combined.

The temporary-partial immunity model is represented by the diagram in Fig. 1(e). As before, α is a parameter that controls the rate of loss of immune protection, and σ controls the degree of protection that individuals acquire upon recovery from infection. The model equations are

$$\begin{aligned} \frac{dS}{dt} &= e - R_0IS - eS + \alpha(1 - e)(1 - S), \\ \frac{dI}{dt} &= R_0I(S + \sigma(1 - S - I)) - I. \end{aligned} \tag{6}$$

Fig. 5 illustrates the position of *SIR* and *SIS* frameworks in the parameter space (α, σ) that characterizes temporary-partial immune protection. The *SIR* model is retrieved at $\alpha=0$ and $\sigma=0$, and the *SIS* model corresponds to $\alpha=1$ or $\sigma=1$. The temporary immunity model is retrieved at $\sigma=0$, and the partial immunity model is obtained at $\alpha=0$. As observed in Fig. 4, a temporary immunity mechanism has a strong impact on potential epidemic dynamics, and a partial immunity mechanism induces a second transmission threshold separating regions of low-oscillating and high-steady endemicity. Here, we analyse the combined effects of these two mechanisms. Fig. 6 illustrates how the behaviour of the system depends on α and σ for three fixed values of R_0 . Fig. 6(a), (c) and (e) show the stable endemic equilibrium, for $R_0=3.5, 4.0, 4.5$, respectively, indicating whether convergence is by damped oscillations (light grey) or linear decay (dark grey). The respective contour plots for the period of the oscillations are shown in Fig. 6(b), (d) and (f).

4. Temporary immune protection and vaccination

We proceed to illustrate the impact of mass vaccination at birth under the temporary immunity model. The model, which assumes that protection induced by the vaccine is equivalent to protection acquired in response to natural infection, is formalized by the system of

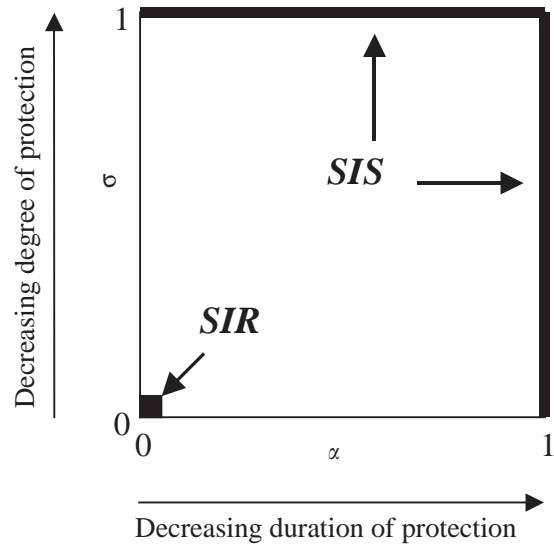


Fig. 5. Schematic diagram illustrating the position of the *SIR* and *SIS* frameworks in the parameter space (α, σ) characterizing temporary-partial immune protection.

equations

$$\begin{aligned} \frac{dS}{dt} &= (1 - v)e - R_0IS - eS + \alpha(1 - e)(1 - S), \\ \frac{dI}{dt} &= R_0IS - I, \end{aligned} \tag{7}$$

where the new parameter, v , represents coverage of the vaccination programme. Fig. 7 illustrates the effect of vaccination in this system when $\alpha=0.0015$. In this case, the average duration of immunity, $1/\alpha(1-e)$, corresponds to approximately 55 years to represent the loss of immunity commonly observed late in life (which is sometimes observed, even for *SIR* type infections). The full curves in Fig. 7(a) represent the endemic equilibrium without vaccination (top curve) and when the system is subject to a mass vaccination programme with 90% coverage (bottom curve). The dashed line represents the unrealistic limit of 100% vaccination coverage. The dashed line meets the R_0 -axis at the critical value:

$$R_{0\alpha} = 1 + \frac{e}{\alpha(1 - e)}$$

implying that eradication is possible only if, in the absence of vaccination, R_0 is lower than this threshold. For the parameter values set here, $R_{0\alpha}$ is approximately 1.84. This appears remarkably low when we note that in the idealized situation of lifelong immunity, $\alpha=0$, the eradication threshold would be pushed to infinity, implying that the infection could always be eradicated by vaccination as long as the coverage was sufficiently high (i.e. greater than $1-1/R_0$). Fig. 7(b) shows the result of simulations with two model populations: population A below the eradication threshold; and population B above the eradication threshold.

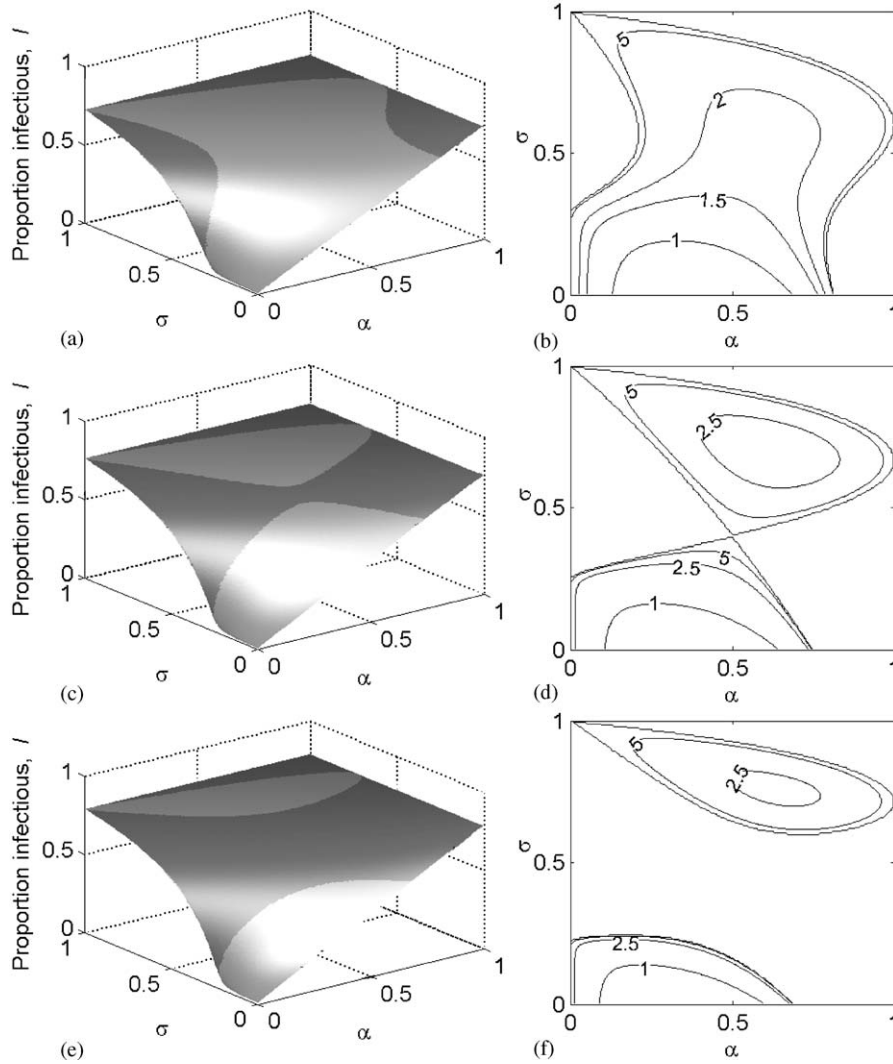


Fig. 6. Equilibria and convergence for the temporary-partial immunity model. (a), (c) and (e) endemic equilibrium as a function of α and σ for three values of R_0 (3.5, \leftrightarrow 4.0, \leftrightarrow 4.5) indicating whether convergence is by damped oscillations (light grey) or linear decay (dark grey); (b), (d) and (f) corresponding contour plots for the period of the oscillations in the damped oscillatory region (the labels represent years).

The main observation from these results is that waning of immunity is a major obstacle to the eradication of infectious diseases. This is illustrated here with a situation where immunity lasts, on average, 55 years and the average lifetime is 70 years. The deterministic models used here predict that, in this scenario, eradication is possible only if R_0 is below 1.84. Further investigations with discrete stochastic models may give more optimistic results, but this is beyond the scope of this paper.

5. Partial immune protection and vaccination

Here, we illustrate the impact of mass vaccination at birth in the case where both infection and vaccination induce partial protection. First, we consider the simplest situation of a vaccine that induces as much protection as

a previous infection. Then we analyse the impact of more efficacious vaccines.

5.1. The reinfection threshold

Before describing vaccination models, we elaborate on a concept that will be crucial to the analysis—the “reinfection threshold”. The full line of Fig. 8 shows how the endemic equilibrium increases with R_0 when $\sigma = 0.25$, revealing a steep increase as transmission crosses

$$R_{0\sigma} = \frac{1}{\sigma}$$

which in this case is $R_0 = 4$. This increase is associated with the reinfection threshold. The dotted and dashed lines are plotted to illustrate the contributions of primary infection and reinfection to the overall infection

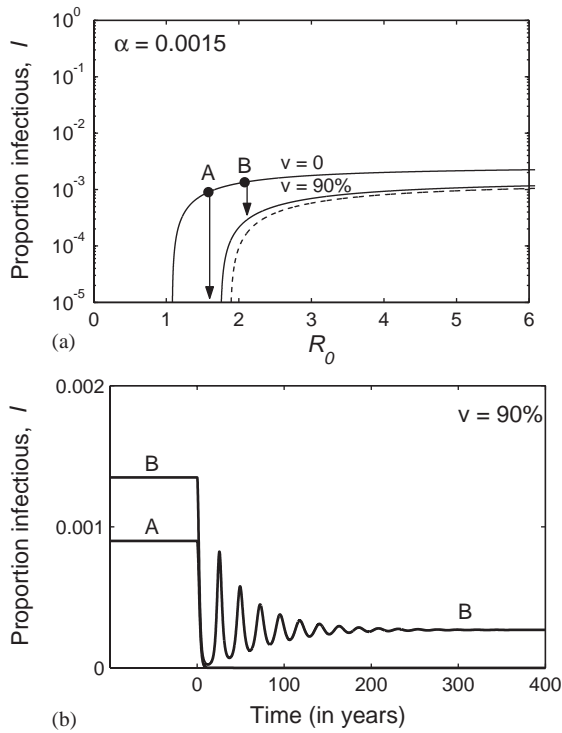


Fig. 7. Temporary protection the global impact of mass vaccination. The rate of loss of immune protection is fixed by setting $\alpha=0.0015$, implying that the average duration of protection of 55 years. (a) shows endemic equilibrium of model (7) as a function of R_0 : without vaccination (higher full curve), with 90% vaccination coverage (lower full curve), and with 100% vaccination coverage (dashed curve). The dots A and B represent two model population with different R_0 : eradication is possible in A, but not in B. (b) simulates the implementation of a vaccination programme (with 90% coverage) on both populations.

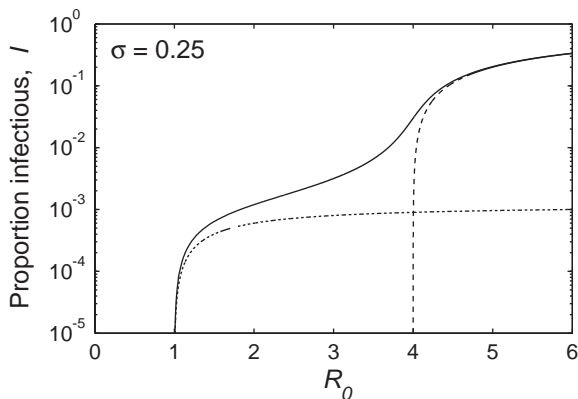


Fig. 8. The reinfection threshold. The full line shows the endemic equilibrium of model (5) as a function of R_0 . The dotted line corresponds to the *SIR* submodel $dS/dt = e - R_0IS - eS$; $dI/dt = R_0IS - I$ and represents primary infection. The submodel sets the invasion threshold at $R_0=1$. The dashed line corresponds to the *SIS* submodel $dI/dt = \sigma R_0I(1-I) - I$ and represents reinfection. By fixing $\sigma=0.25$, the reinfection threshold is set at $R_0=1/\sigma=4$.

levels, and correspond to the equilibria of two sub-models: the dotted line corresponds to the *SIR* model $dS/dt = e - R_0IS - eS$; $dI/dt = R_0IS - I$ and represents pri-

mary infection; and the dashed line corresponds to the *SIS* model $dI/dt = \sigma R_0I(1-I) - I$ and represents reinfection. The invasion threshold ($R_0=1$) is associated with the primary infection submodel, and the reinfection threshold ($R_0=4$) is associated with the reinfection submodel. The full model combines the two dynamical processes, and reveals the potential to convert small baseline variabilities of the transmissibility parameter, R_0 , into large variabilities of infection prevalence.

5.2. Vaccine inducing protection equivalent to natural infection

The first vaccination model corresponds to a vaccine that induces an immune response equivalent to that induced by a previous infection. This is formalized by the system of equations

$$\begin{aligned} \frac{dS}{dt} &= (1-v)e - R_0IS - eS, \\ \frac{dI}{dt} &= R_0I(S + \sigma(1-S-I)) - I, \end{aligned} \tag{8}$$

where v is the vaccination coverage. Fig. 9(a) and (c) illustrates the effect of mass vaccination in this scenario. Note that such a vaccine is protective against primary infection, but not against reinfection. Primary infection is associated with the susceptibility pool S ; and reinfection associated with the susceptibility pool $S+R$. Vaccination moves individuals from S to R and therefore, essentially reduces the resource for primary infections without directly affecting reinfection. Therefore, it is expected that a mass vaccination programme will be effective below the reinfection threshold, but not above. This is evident from Fig. 9(a), where we show the equilibria without vaccination and with 90% vaccination coverage (full lines). The dashed line corresponds the unrealistic limit of 100% vaccination coverage. Three model populations are marked: population A is below the reinfection threshold and infection eradication is expected with the 90% vaccination programme; population B is also below the reinfection threshold and although 90% coverage is not sufficient for eradication, a substantial reduction in prevalence is expected; population C is above the reinfection threshold and the vaccination programme has only a minor effect on the equilibrium prevalence. Fig. 9(c) shows the results of simulating vaccination on the three populations.

The main outcome of this analysis is to demonstrate with a simple model how a vaccination programme can have such variable outcomes. Tuberculosis and the bacille Calmette-Guérin (BCG) vaccine constitute the best documented example of this phenomenon, and this is further discussed in (Gomes et al., 2004a). The variability associated with the reinfection threshold presents serious difficulties to the estimation of vaccine

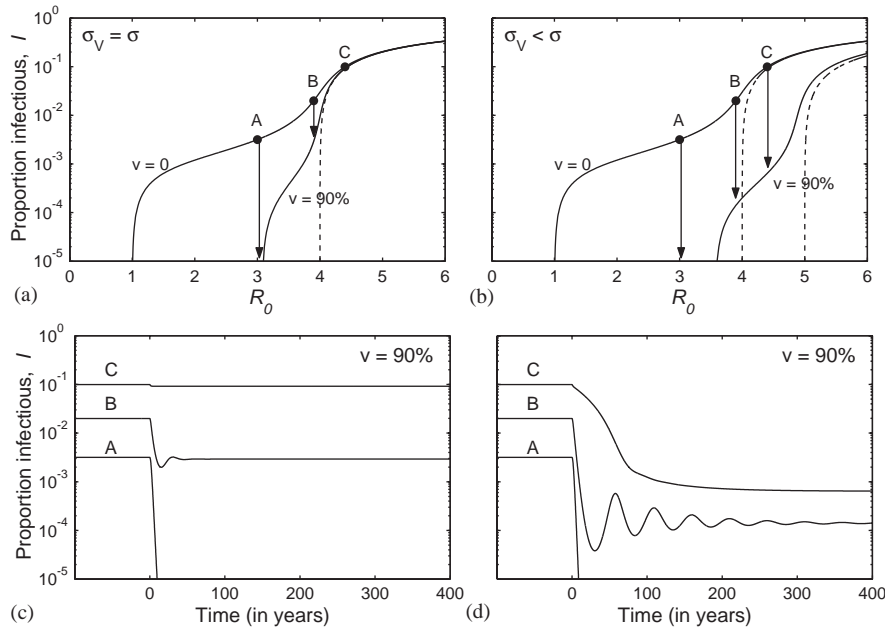


Fig. 9. Partial protection and the global impact of mass vaccination. Using the endemic equilibria of Fig. 8 as baseline pre-vaccination states, we illustrate the impact of two vaccination programmes: (a) and (c) show the result of 90% coverage with a vaccine that confers as much protection as natural infection (model (8)); (b) and (d) show the result of 90% coverage with a vaccine that is more potent than natural infection (model (9) with $\sigma_V = 0.2$). The second vaccine increases the reinfection threshold generating spectacular outcomes in high burden regions that were insensitive to the first vaccine (e.g. population C).

efficacy from field observations, which led to great controversy concerning the use of BCG.

5.3. Vaccine inducing greater protection natural infection

A sound understanding of susceptibilities to infection and reinfection is essential to predict the impact of vaccination programmes when immunity is partially protective. The hope is that, where transmissibility is above the reinfection threshold, vaccines that protect more than a natural infection might prove much more effective. This can be demonstrated with the model

$$\begin{aligned}
 \frac{dS}{dt} &= (1 - v)e - R_0IS - eS, \\
 \frac{dI_S}{dt} &= R_0I(S + \sigma R) - I_S, \\
 \frac{dR}{dt} &= (1 - e)I_S - \sigma R_0IR - eR, \\
 \frac{dV}{dt} &= ve + (1 - e)I_V - \sigma_V R_0IV - eV, \\
 \frac{dI}{dt} &= \sigma_V R_0IV - I_V,
 \end{aligned} \tag{9}$$

where σ_V is the factor by which vaccination reduces susceptibility, and here we consider $\sigma_V < \sigma$. Fig. 9(b) and (d) illustrates the effect of such vaccine when $\sigma = 0.25$ as before and $\sigma_V = 0.2$. The full lines in Fig. 9(b) represent the pre-vaccination state (as before) and the post-vaccination equilibrium with 90% coverage. We see

that the range of R_0 where this vaccine has the potential to be highly effective has increased, going up to $1/\sigma_V$, which is $R_0 = 5$ in this case. In other words, if vaccine protection is higher than naturally acquired protection ($\sigma_V < \sigma$) then vaccination has the power to increase the reinfection threshold to

$$R_{0V} = \frac{1}{\sigma_V}.$$

From the three model populations used before, it is population C (where endemicity is highest) who would benefit the most with the development of such vaccine. Fig. 9(d) shows the simulations of the vaccination programme from the time it is introduced until the new equilibrium is reached.

These results have important implications for public health. A great challenge of vaccine development is to supersede the protection provided by natural infection. The reinfection threshold magnifies epidemiological variabilities, but perhaps the most exciting news is that this threshold can be manipulated by vaccines. Vaccines that protect more than naturally acquired infection increase the threshold for reinfection, and may have greater impact than expected in regions afflicted by high burdens of disease. In the case of tuberculosis, a number of strategies for the development of better vaccines are being followed (Britton and Palendira, 2003; Olsen and Andersen, 2003), and the potential of such interventions cannot be overemphasized (Gomes et al., 2004a).

6. Discussion

Since the majority of pathogen infections involve reinfection, the dynamic consequences of any intervention on rates of (re)infection should be considered. We suggest that most infections can be classified according to the relative significance of partial and temporary immunity in creating susceptibility to reinfection (Fig. 5). Temporary immunity is, to a large extent, determined by the propensity for antigenic change, but also alteration in host status (e.g. immunosuppression due to other infections). Partial immunity is more dependent on the interaction between static host and pathogen variation: high parasite variation decreasing partial immunity, and high host variation increasing partial immunity. As rates of antigenic change will depend on the variability in the population, one might expect that pathogens occupy the diagonal region in Fig. 5.

The exact mechanism by which individuals regain susceptibility following infection has an important influence on the level of infection maintained in the population and on the potential periodicity of epidemics. Fig. 4(a) and (b) suggests that waning of immunity has a crucial impact on the time-scale for potential inter-epidemic periods. Fig. 4(c) and (d) reveals two clearly distinct types of endemic behaviour: low and high. The first requires that transmissibility is just above $R_0 = 1$. The second relies on reinfection and requires that transmissibility is above $R_0 = 1/\sigma$. Furthermore, the different mechanisms of immunity failure impact differentially on control strategies.

Many diseases are subject to seasonal forcing of transmission, and this has motivated extensive investigations for the *SIR* scenario (Keeling et al., 2001). Basically, the resulting inter-epidemic period is inferred from the interplay of the annual cycle and the period of the damped oscillations (Weber et al., 2001). Here we observe that the period for damped oscillations is highly sensitive to the mode of action of immune protection (especially temporary immunity). A combination of these mechanisms with seasonality may be necessary to explain time-scales observed in epidemic cycles.

A more realistic scenario is to consider many levels of susceptibility to infection, or even a continuum. This has been previously implemented (White and Medley, 1998) but analysis has been limited by model complexity. In cases of antigenic diversity, the dynamics of immunity should somewhat correlate with the pathogen evolution, and this has been explored to some extent (Gomes et al., 2002, 2004b; Gog and Grenfell, 2002; Ferguson et al., 2003). In the context of *Trypanosoma* infections, Coen et al. (2001) attempted to estimate the rate of loss of immunity from seroprevalence data. However, unless primary infection can be distinguished from subsequent infection in some manner (e.g. antibody profile), the parameters for infection rate and immunity loss will be

colinear, complicating statistical analysis. A further complication is that the risk of disease on infection might be higher for primary infection (e.g. RSV), or increase with time since last infection or be determined by pathogen (genetic) type. Consequently, the impacts that we show in terms of prevalence and incidence of infection might be different from prevalence and incidence of disease. We have also assumed that subsequent infections are as infectious as primary infections, which is also unlikely to be generally true.

The incidence of pulmonary tuberculosis can vary by two orders of magnitude between different regions of the world, and estimates for the efficacy of the BCG vaccine vary between 0% and 80%. This scenario is aggravated by an association between high prevalence of infection and low vaccine efficacy (Olsen and Andersen, 2003). It is reasonable to expect vaccination programmes to be less successful in highly endemic regions, and all models discussed here show this effect to some extent. However, the extreme variabilities observed in TB and BCG efficacy are a specificity of the partial immune protection mechanism, due to the reinfection threshold. We have recently proposed this mechanism as an explanation for the BCG discrepancies (Gomes et al., 2004a). Here we show that the conclusions are not specific to tuberculosis but rather, they are a general feature of diseases characterized by partial immune protection and the associated reinfection threshold.

The reinfection threshold represents the transmissibility required to promote recurrent infections. Populations that exceed this threshold, sustain high levels of infection and tend to be insensitive to interventions. Furthermore, the reinfection threshold can be potentially manipulated by vaccination. Vaccines that induce more protection than a natural infection increase the reinfection threshold, providing means for the control of recurrent infection in high-burden regions. Consequently, the partial immune protection framework can serve as a basis to assess the impact of specific vaccines, and to set targets for future vaccine performance in terms of their epidemiological impact (McLean, 1995b). While such vaccines are currently unavailable, efforts can be exerted into manipulating transmission. That is, in populations of high endemicity, improvements in hygiene or reduced crowding could reduce the risk of transmission and therefore the basic reproduction number. If the basic reproduction number is reduced below the reinfection threshold, there would be resulting a drop in prevalence of infection and an increase in apparent vaccination effectiveness.

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