

Intentional infection as a method of population-level disease control

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June 16th 2018

Abstract

In this paper, we study the possible advantages of intentional infection, as a method of population-level disease control. Intentional infection is a generalization of variolation, which was invented in 15th century, and widely used around the world in 17th and 18th century. People believed that variolation would result in a mild but protective infection, giving them a higher chance of survival than by being naturally infected. This paper aims to provide mathematical models describing the dynamics of infected classes, when intentional infection is introduced on a population level, and to perform predictions based on the models.

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1 Introduction

Disease control has drawn people's attention since the early stage of human civilization. Other than curing diseases by medication, people have also discovered and attempted a vast range of method for disease prevention. Nowadays, people have vaccination as a primary method to prevent diseases [2]. However, before vaccination became a developed technology, people had limited power when encountering fatal diseases such as smallpox.

A practice called variolation was invented as an attempt to contain smallpox [5]. The earliest record of such method can be found in the 15th century, from an ancient Chinese documentation [6]. It is a method used to immunize individuals against smallpox with materials taken from other smallpox patient, or from recently variolated individuals. People have found that, by variolation, a mild, but protective infection will likely occur [6]. Instead of having a considerable chance of becoming gravely ill, a vast majority of variolated cases will survive after an infection. As a result, variolated individuals have a higher chance of survival.

More generally speaking, a similar method may be applied to other diseases for disease control. Fundamentally, since individuals are deliberately exposed to living virus, variolation is an example of intentional infection. Although an individual benefits from intentional infection when there is threat of being transmitted with fatal disease [6], people did not understand whether intentional infection can also bring positive effects when applied on a population level.

Similar to vaccination, there are different strategies for variolation. One of strategies is to intentional infect newborn individuals [3]. In fact, newborn infection does not mean instant infection when people are born, but rather an infection after their maternal immunity wanes [4]. Another strategy considered in this paper is to intentionally infect susceptible individuals in the population, at a fixed rate [9].

2 Models: Intentional infect proportion of Newborn

2.1 Model: Modification to SIR model

2.1.1 System of differential equations

We begin our analysis by modifying the SIR model. Our first strategy is to intentionally infect newborn individuals, with a certain proportion. The following assumptions are made to simplify the model:

- No difference between intentionally infected and naturally infected individuals.
- No disease induced mortality (disease induced mortality will be introduced in the later models).
- Birth and natural death rate are the same (total population N remains constant).
- The latent period (time from infection to becoming infectious) is short enough to be ignored.
- All susceptible individuals are equally likely to be infected, and all infected individuals are equally infectious.

Equipped with the assumptions above, we now setup our system of differential equations.

Just like in SIR model, S , I and R represent the proportion of susceptible, infected and recovered with respect to total population.

$$\begin{aligned}\frac{dS}{dt} &= \mu(1 - p) - \beta SI - \mu S, \\ \frac{dI}{dt} &= \beta SI + \mu p - \gamma I - \mu I, \\ \frac{dR}{dt} &= \gamma I - \mu R.\end{aligned}\tag{1}$$

Here, β is the transmission rate, γ is the recovery rate, μ is the *per capita* rate of birth and death, p is the proportion of newborns that are intentionally infected.

We non-dimensionalize Equation 1 by scaling time, using

$$\tau = (\gamma + \mu)t, \quad (2)$$

so the time unit now is “mean time infected”.

The dimensionless system is:

$$\frac{dS}{d\tau} = \epsilon(1 - p) - \mathcal{R}_0 SI - \epsilon S, \quad (3a)$$

$$\frac{dI}{d\tau} = \mathcal{R}_0 SI + \epsilon p - I, \quad (3b)$$

where $\epsilon = \frac{\mu}{\gamma + \mu}$, $\mathcal{R}_0 = \frac{\beta}{\gamma + \mu}$.

We are not considering $\frac{dR}{d\tau}$ since it does not impact the dynamics of S and I , and we do not need to track the proportion of recovered individuals in this model.

2.1.2 Equilibria

By setting Equation 3 equal to 0, we solve for equilibria. The only equilibrium for this model is,

$$\hat{S} = \frac{1}{\mathcal{R}_0} - \frac{2p}{(\mathcal{R}_0 - 1) + \sqrt{(\mathcal{R}_0 - 1)^2 + 4\mathcal{R}_0 p}}, \quad (4a)$$

$$\hat{I} = \frac{\epsilon(\mathcal{R}_0 - 1) + \epsilon\sqrt{(\mathcal{R}_0 - 1)^2 + 4\mathcal{R}_0 p}}{2\mathcal{R}_0}. \quad (4b)$$

Notice, Equation 4b does not return 0 for any p value between 0 and 1, meaning there is always infected individuals present in the population. Therefore, we can claim that this is not a disease free equilibrium. It follows that the equilibrium above is an endemic equilibrium (EE).

We would like to know if the EE is stable, therefore we need the Jacobian matrix of Equation 3. The Jacobian is,

$$\mathcal{J} = \begin{bmatrix} -\mathcal{R}_0 I - \epsilon & -\mathcal{R}_0 S \\ \mathcal{R}_0 I & \mathcal{R}_0 S - 1 \end{bmatrix}. \quad (5)$$

Now for convenience, let

$$K = (\mathcal{R}_0 - 1) + \sqrt{(\mathcal{R}_0 - 1)^2 + 4\mathcal{R}_0 p}. \quad (6)$$

For the purpose of our discussion, we are interested in disease that transmits fast enough to result in an epidemic. Therefore, the \mathcal{R}_0 value for the disease has to be greater than 1.

Notice, $K > 0$ if $p \neq 0$.

Thus, the Jacobian evaluated at endemic equilibrium is,

$$\mathcal{J}|_{EE} = \begin{bmatrix} -\frac{\epsilon K}{2} - \epsilon & -1 + \frac{2p\mathcal{R}_0}{K} \\ \frac{\epsilon K}{2} & -\frac{2p\mathcal{R}_0}{K} \end{bmatrix}, \quad (7)$$

The eigenvalues of Equation 7 are,

$$\lambda_{1,2} = \frac{-(\epsilon K^2 + 2\epsilon K + 4p\mathcal{R}_0) \pm \sqrt{(\epsilon K^2 + 2\epsilon K + 4p\mathcal{R}_0)^2 - 4(2\epsilon K^3 + 8\epsilon K p \mathcal{R}_0)}}{4K} \quad (8)$$

Since $(2\epsilon K^3 + 8\epsilon K p \mathcal{R}_0) > 0$, if the discriminant is positive, we have

$$\sqrt{(\epsilon K^2 + 2\epsilon K + 4p\mathcal{R}_0)^2 - 4(2\epsilon K^3 + 8\epsilon K p \mathcal{R}_0)} \leq |\epsilon K^2 + 2\epsilon K + 4p\mathcal{R}_0|, \quad (9)$$

thus, $\Re(\lambda_{1,2}) < 0$.

But if the discriminant is negative, we have

$$\Re(\lambda_1) = \Re(\lambda_2) = -(\epsilon K^2 + 2\epsilon K + 4p\mathcal{R}_0) < 0, \quad (10)$$

As a result, we can conclude that EE is stable.

To fully investigate the dynamics of the system, we're also interested in whether the eigenvalues could be complex, which will lead to damped oscillation.

Although it is difficult to determine the sign of the discriminant analytically, we can plot the value of the discriminant as a function of other parameter, i.e. p or \mathcal{R}_0 .

134 We begin our analysis with specific values of each parameter. Given the variolation history
 135 of smallpox, it is reasonable to adopt its parameter values as an example. The values are
 136 listed in Table 1.

Table 1: Model parameters and smallpox values.

Symbol	Meaning	Value
μ	Natural <i>per capita</i> death rate	$\frac{1}{50 \times 365}$ per day
γ	Recovery rate	$\frac{1}{22}$ per day
\mathcal{R}_0	Basic reproductive number	4.5

137 2.1.3 Region of (\mathcal{R}_0, p) plane where there are damped oscillations (fixed ϵ)

138 p is the proportion of intentional infection and \mathcal{R}_0 is the basic reproduction number.

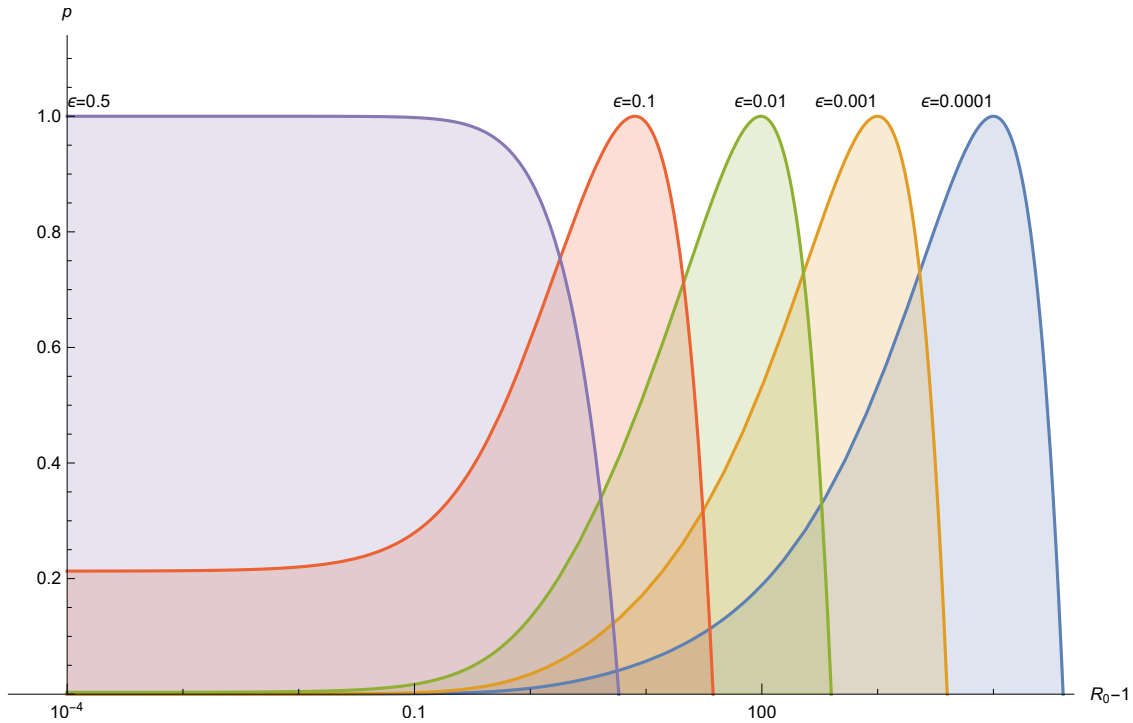


Figure 1: The areas under each curve, shaded in different colors, represent the region of damped oscillations with respect to different ϵ values.

2.1.4 Region of $(\mathcal{R}_0, \epsilon)$ plane where there are damped oscillations (fixed p)

The plots below were made with different p in increasing order, the shaded area between the blue curve and the orange curve represents the region where system has damped oscillation.

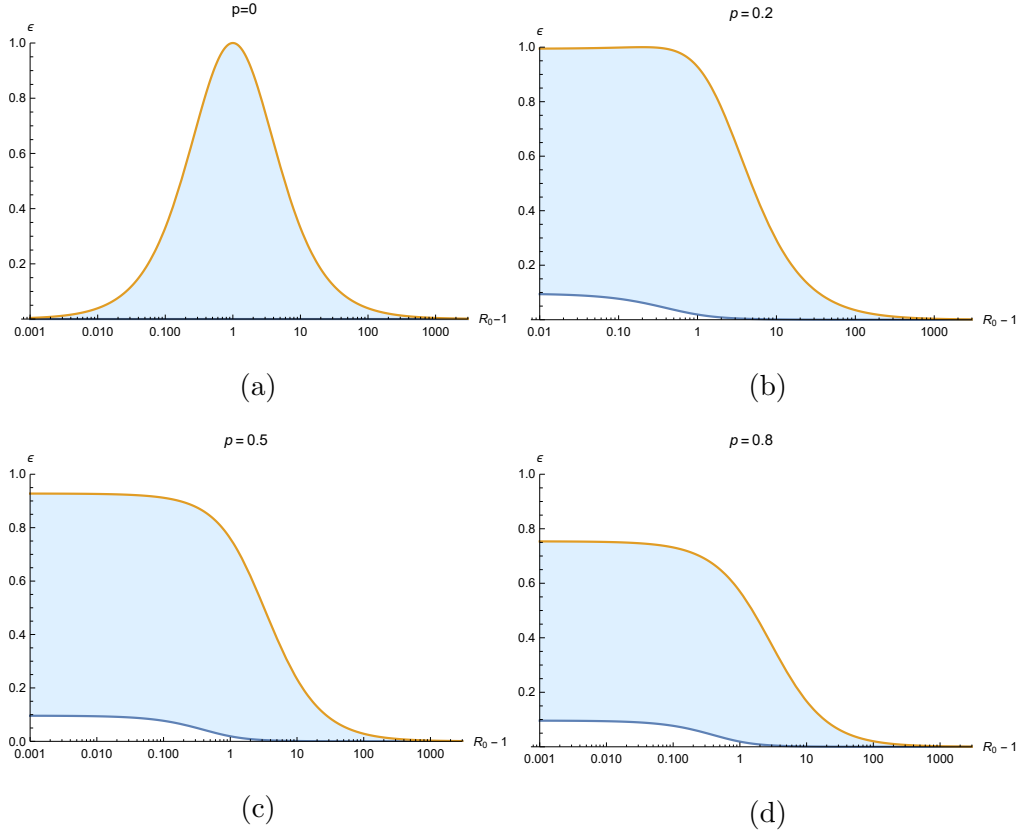


Figure 2: Panel (a)-(d) illustrate $(\mathcal{R}_0, \epsilon)$ planes with different p values. The shaded regions represent the range of conditions for damped oscillation to occur.

a) $p = 0$, b) $p = 0.2$, c) $p = 0.5$, d) $p = 0.8$

2.1.5 Comments and discussion on this model

This is the initial model of intentional infection on newborns, which we obtained by directly modifying standard SIR model. Notice, the main difference between intentional infection model and vaccination model is the direction of flow of individuals being treated with intentional infection or vaccination. Intentionally infected individuals are capable of transmitting the disease to susceptible individuals, whereas the vaccinated individuals will enter R (recovered) directly, therefore not being able to impact the dynamics of the system anymore.

In the past, for the case of smallpox, people believed that individuals that are variolated have a much better chance of survival, compared with naturally infected cases. Therefore, we should divide the infected compartment into two separate infective classes, namely, intentionally infected and naturally infected. To investigate the benefit of intentional infection, here we define “advantageous” as fewer total death. More specifically, we will be comparing total disease induced mortality. To achieve this, we need to involve a new parameter, namely, case fatality proportion.

2.2 Model: Addition of disease induced mortality

2.2.1 System of differential equations

Disease induced mortality is created by deaths of infected population, with a ratio. This ratio is also known as case fatality proportion.

Historic application of variolation have shown a lower case fatality proportion for those being variolated, compared to those being naturally infected. Thus, we need to assign different case fatality proportion to each class. This also requires us to divide I from our previous model into two distinct infective classes, “Intentionally infected” (V) and “Naturally infected” (I).

In this model, we still assume no latent period, and similar to the last assumption in our

previous model, we assume that all infected individuals(including V and I) are equally infectious.

After applying all assumptions, our model becomes,

$$\begin{aligned}
\frac{dS}{dt} &= \mu(1-p) - \beta S(V+I) - \mu S, \\
\frac{dV}{dt} &= \beta SV + \mu p - \gamma V - \mu V, \\
\frac{dI}{dt} &= \beta SI - \gamma I - \mu I, \\
\frac{dM}{dt} &= p_V \gamma V + p_I \gamma I, \\
\frac{dR}{dt} &= (1-p_V) \gamma V + (1-p_I) \gamma I - \mu R.
\end{aligned} \tag{11}$$

In addition to the previous model, p_V and p_I represent the case fatality proportion for intentionally infected and naturally infected cases, respectively.

Again, we non-dimensionalize Equation 11 by time, using

$$\tau = (\gamma + \mu)t, \tag{12}$$

which yields,

$$\frac{dS}{d\tau} = \epsilon(1-p) - \mathcal{R}_0 S(V+I) - \epsilon S, \tag{13a}$$

$$\frac{dV}{d\tau} = \mathcal{R}_0 SV + \epsilon p - V, \tag{13b}$$

$$\frac{dI}{d\tau} = \mathcal{R}_0 SI - I, \tag{13c}$$

$$\frac{dM}{d\tau} = p_V(1-\epsilon)V + p_I(1-\epsilon)I, \tag{13d}$$

$$\frac{dR}{d\tau} = (1-p_V)(1-\epsilon)V + (1-p_I)(1-\epsilon)I - \epsilon R. \tag{13e}$$

Notice here that \mathcal{R}_0 , which is equal to $\frac{\beta}{\gamma+\mu}$, is again the basic reproduction number of the naturally infected cases. Since we assume that the transmission rate of naturally infected and intentionally infected cases are identical, \mathcal{R}_0 also plays a role as the basic reproduction number of intentionally infected cases.

Our construction of the model suggests, once an individual is intentionally infected, either directly or indirectly, this individual will not be naturally infected anymore.

2.2.2 Equilibria

By setting Equation 13a, Equation 13b and Equation 13c equal to 0, we solve for equilibria.

If $p = 0$, meaning there is no intentional infection, our system reduces to the standard SIR model.

There are two equilibria for the standard SIR model, an Endemic equilibrium,

$$\hat{S} = \frac{1}{\mathcal{R}_0}, \quad (14a)$$

$$\hat{V} = 0, \quad (14b)$$

$$\hat{I} = \epsilon(1 - \frac{1}{\mathcal{R}_0}). \quad (14c)$$

And a disease free equilibrium(with both \hat{V} and \hat{I} are 0),

$$\hat{S} = 1, \quad (15a)$$

$$\hat{V} = 0, \quad (15b)$$

$$\hat{I} = 0. \quad (15c)$$

However, if $p \neq 0$, there is only one equilibrium,

$$\hat{S} = \frac{1}{\mathcal{R}_0} - \frac{2p}{(\mathcal{R}_0 - 1) + \sqrt{(\mathcal{R}_0 - 1)^2 + 4\mathcal{R}_0p}}, \quad (16a)$$

$$\hat{V} = \frac{\epsilon(\mathcal{R}_0 - 1) + \epsilon\sqrt{(\mathcal{R}_0 - 1)^2 + 4\mathcal{R}_0p}}{2\mathcal{R}_0}, \quad (16b)$$

$$\hat{I} = 0. \quad (16c)$$

It is worthwhile mentioning that, since $\hat{V} \neq 0$, for any p between 0 and 1, this equilibrium is not disease free. It follows that this equilibrium is the endemic equilibrium.

Stability analysis relies on Jacobian matrix, which is,

$$\mathcal{J} = \begin{bmatrix} -\mathcal{R}_0(V + I) - \epsilon & -\mathcal{R}_0S & -\mathcal{R}_0S \\ \mathcal{R}_0V & \mathcal{R}_0S - 1 & 0 \\ \mathcal{R}_0I & 0 & \mathcal{R}_0S - 1 \end{bmatrix}. \quad (17)$$

210 Eigenvalues of Jacobian are given as follow. Here in our derivations, we are considering
 211 $S = \hat{S}$, $V = \hat{V}$.

$$212 \quad \lambda_1 = -1 + \mathcal{R}_0 S \quad (18a)$$

$$213 \quad \lambda_2 = \frac{-1 + \mathcal{R}_0 S - \epsilon - \mathcal{R}_0 V - \sqrt{(-1 + \mathcal{R}_0 S - \epsilon - \mathcal{R}_0 V)^2 - 4(\mathcal{R}_0 + \epsilon - \mathcal{R}_0 S \epsilon)}}{2} \quad (18b)$$

$$214 \quad \lambda_3 = \frac{-1 + \mathcal{R}_0 S - \epsilon - \mathcal{R}_0 V + \sqrt{(-1 + \mathcal{R}_0 S - \epsilon - \mathcal{R}_0 V)^2 - 4(\mathcal{R}_0 + \epsilon - \mathcal{R}_0 S \epsilon)}}{2} \quad (18c)$$

216 By using Equation 16a and Equation 18a, we obtain

$$217 \quad -1 + \mathcal{R}_0 S = -\frac{2p\mathcal{R}_0}{(\mathcal{R}_0 - 1) + \sqrt{(\mathcal{R}_0 - 1)^2 + 4\mathcal{R}_0 p}} < 0 \quad (19)$$

218 Therefore,

$$219 \quad \Re(\lambda_1) = -1 + \mathcal{R}_0 S < 0. \quad (20)$$

220 To determine the real part of λ_2 and λ_3 , we need to determine the sign of the quantity under
 221 the square root in equation Equation 18b and Equation 18c.

222 By using Equation 16a again, we have

$$223 \quad \mathcal{R}_0 S \epsilon < \epsilon. \quad (21)$$

224 Therefore,

$$225 \quad (\mathcal{R}_0 + \epsilon - \mathcal{R}_0 S \epsilon) > \mathcal{R}_0 > 0, \quad (22)$$

226 which means, if the sign of the quantity under the square root is positive, we have

$$227 \quad \sqrt{(-1 + \mathcal{R}_0 S - \epsilon - \mathcal{R}_0 V)^2 - 4(\mathcal{R}_0 + \epsilon - \mathcal{R}_0 S \epsilon)} < |(-1 + \mathcal{R}_0 S - \epsilon - \mathcal{R}_0 V)| \quad (23)$$

228 Therefore, $\Re(\lambda_2) < \Re(\lambda_3) < 0$.

229 Certainly, if the sign of the quantity under the square root is negative,

$$230 \quad \Re(\lambda_2) = \Re(\lambda_3) = -1 + \mathcal{R}_0 S - \epsilon - \mathcal{R}_0 V < 0 \quad (24)$$

231 We are able to conclude that EE is locally asymptotically stable.

232 We would also like to investigate the global stability of the EE. A Lyapunov function is
 233 usually required to conclude on the global stability of an equilibrium, but such function may
 234 not always be easily found. Therefore, instead of searching for a Lyapunov function, we
 235 will show global stability by simulating the system with random initial conditions. If the
 236 EE is globally asymptotically stable (GAS), the dynamics of the system should eventually
 237 converge to EE, regardless of the selection of the initial condition.

238 Here, we define a function that describes the distance between any point in the system to
 239 the EE. Knowing that EE is $(\hat{S}, \hat{V}, \hat{I})$, the distance between any point (S, V, I) to the EE is
 240 given by:

$$d((S, V, I)) = ((\hat{S} - S)^2 + (\hat{V} - V)^2 + (\hat{I} - I)^2)^{\frac{1}{2}} \quad (25)$$

241 If EE is GAS, the output of the distance function with any initial condition will eventually
 242 approach 0. If not, we will see the distance function unable to approach 0 for some initial
 243 conditions.

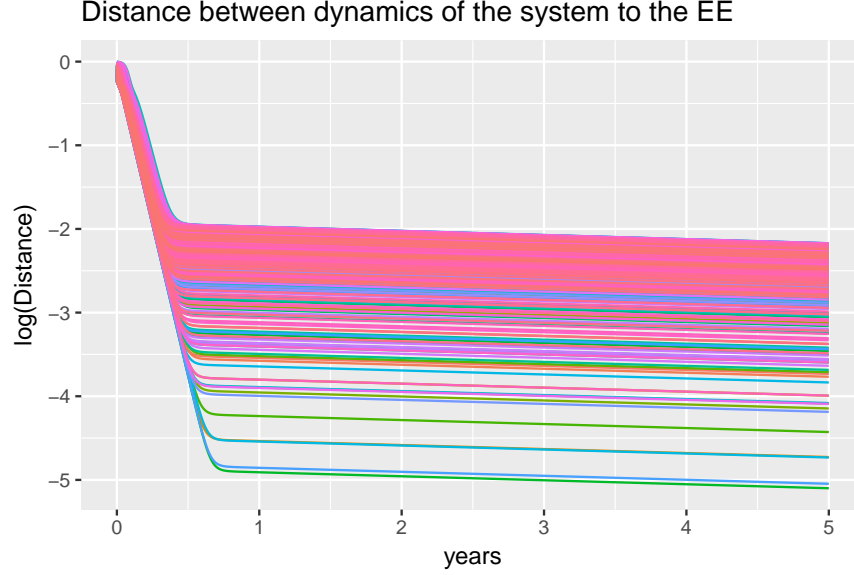


Figure 3: Distance between the dynamics of the system to EE, as a function of time. If EE is GAS, distance function should approach 0, indifferent of which initial condition was chosen. This figure shows the distance function of 1000 random initial conditions. In the long run, the distance between dynamics of the system to EE decays exponentially. We have enough confidence to conclude that EE is GAS.

2.2.3 Effect of intentional infection on total mortality

In epidemic analysis, one way of measuring whether a certain intervention is more advantageous than another is to compare the total disease induced mortality.

We first take a look at the mortality rate at EE, and insert \hat{V} [Equation 16b] and $\hat{I} (= 0)$ in Equation 13d to obtain

$$\left. \frac{dM}{d\tau} \right|_{EE} = p_V(1 - \epsilon)V = \frac{p_V(1 - \epsilon)\epsilon(\mathcal{R}_0 - 1) + p_V(1 - \epsilon)\epsilon\sqrt{(\mathcal{R}_0 - 1)^2 + 4\mathcal{R}_0 p}}{2\mathcal{R}_0}, \quad (26)$$

An important observation is that, at EE, mortality rate increases as p increases. This shows in the long run, having a larger proportion of intentional infection is unwise as it leads to additional deaths.

253 In history, smallpox was present in the human population long before variolation was in-
254 vented. When variolation was introduced, the population was already near equilibrium with
255 no intentional infection ($p = 0$). We are interested in the scenario where intentional infection
256 is introduced after the population is in equilibrium without intentional infection.

257 One of the questions we want to answer is how long does our system take to reach the new
258 EE. In fact, the system will reach the equilibrium only after infinite time. Therefore, we need
259 to define a threshold distance from the equilibrium that we consider to be “at equilibrium”.
260 Since the new equilibrium has $\hat{I} = 0$, we define reaching equilibrium by $I \leq 1 \times 10^{-6}$ (one
261 in a million).

262 If the case fatality proportion for intentionally infected cases is 1%, we can hardly observe
263 any difference when changing proportion of intentional infection. To help us observe the
264 dynamics, we assume intentional infection has a 20% case fatality proportion.

265 Figure 4 shows that it takes shorter time to reach the new EE for a larger proportion of
266 intentional infection.

267 We are also interested in the time it takes for intentional infection to be more advantageous
268 than non-intentional infection, by comparing total mortality.

269 From previous analysis, we have learned that $\frac{dM}{d\tau}$ decreases over time, and stabilizes at a
270 constant rate when reaching the new EE. However, at the new EE, the magnitude of $\frac{dM}{d\tau}$ is
271 larger when p is larger.

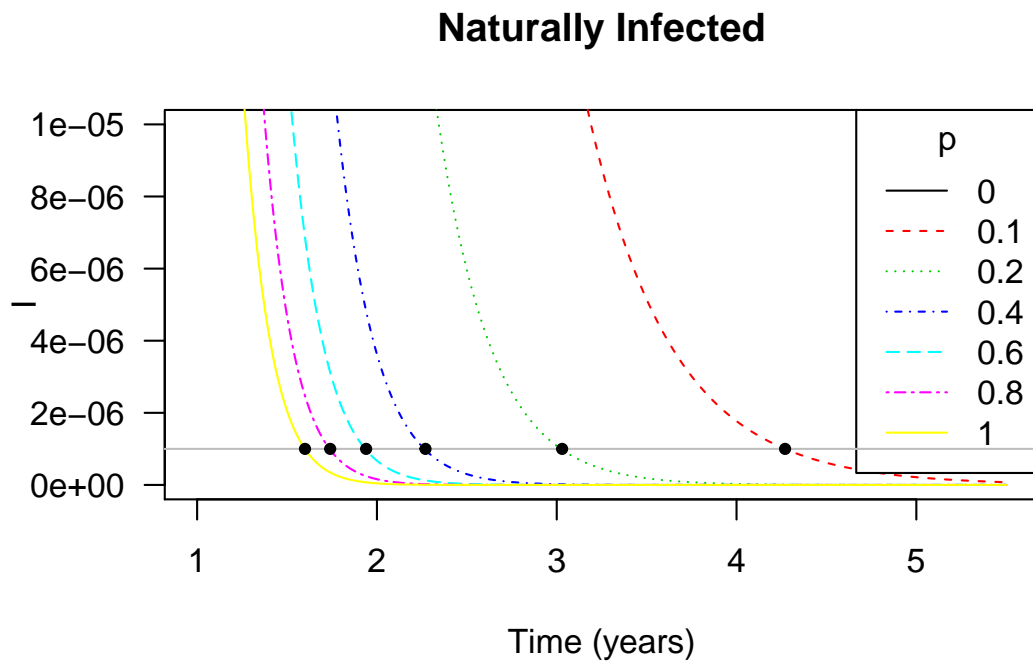


Figure 4: Determining time taken to reach equilibrium. The horizontal line at $I = 10^{-6}$ is the threshold we define to be “at equilibrium”. This figure shows, with a larger proportion of intentional infection on newborn, the system reaches the equilibrium more quickly.

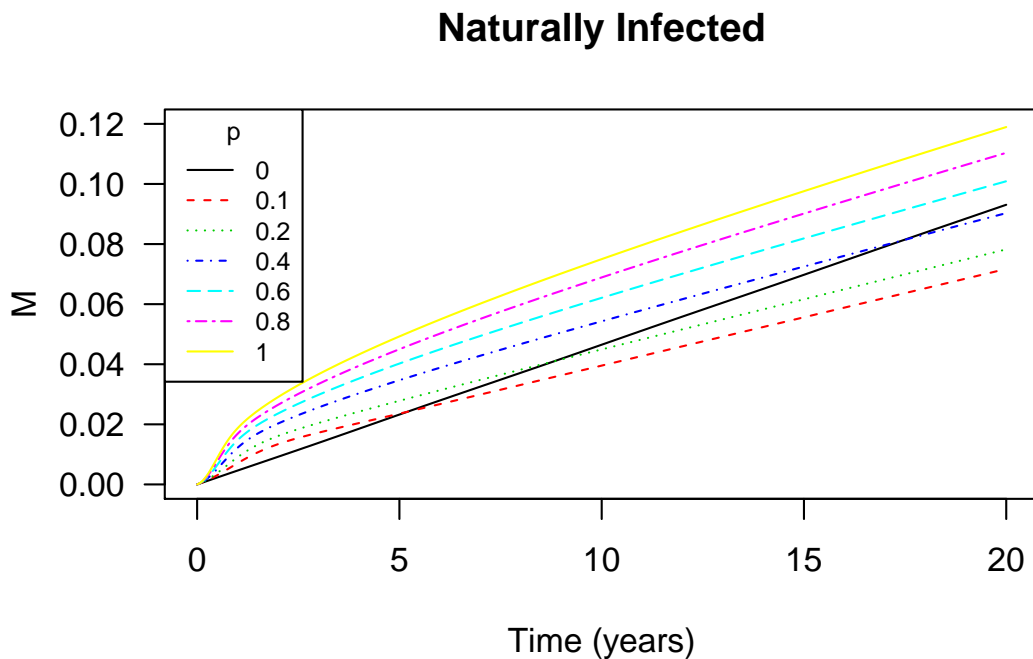


Figure 5: An illustration of intentional infection having advantages over non-intentional infection, with a case fatality proportion from intentional infection of $p_V = 0.2$. Cumulative

Figure 5 shows, at earlier times, intentional infections have steeper slopes than the black line, which is non-intentional infection. The slopes for intentional infection decreases over time, and eventually intentional infection with any proportion will result in fewer total death than non-intentional infection.

Table 2 summarize the time required to reach the new EE and the time required to have advantages over non-intentional infection.

Table 2: Time required to get close to the equilibrium ($\hat{I} < 10^{-6}$) and time to gain advantages over non-intentional infection (i.e., lower cumulative mortality M).

p	Time to EE	Time to have advantages
0.1	4.27 yrs	5.20 yrs
0.2	3.03 yrs	8.81 yrs
0.4	2.27 yrs	17.45 yrs
0.6	1.94 yrs	28.37 yrs
0.8	1.74 yrs	42.43 yrs
1.0	1.60 yrs	61.46 yrs

Above table shows that as p increases, the system reaches the new EE faster (as shown in Figure 6), but time required to gain advantages over the non-intentional infection case increases.

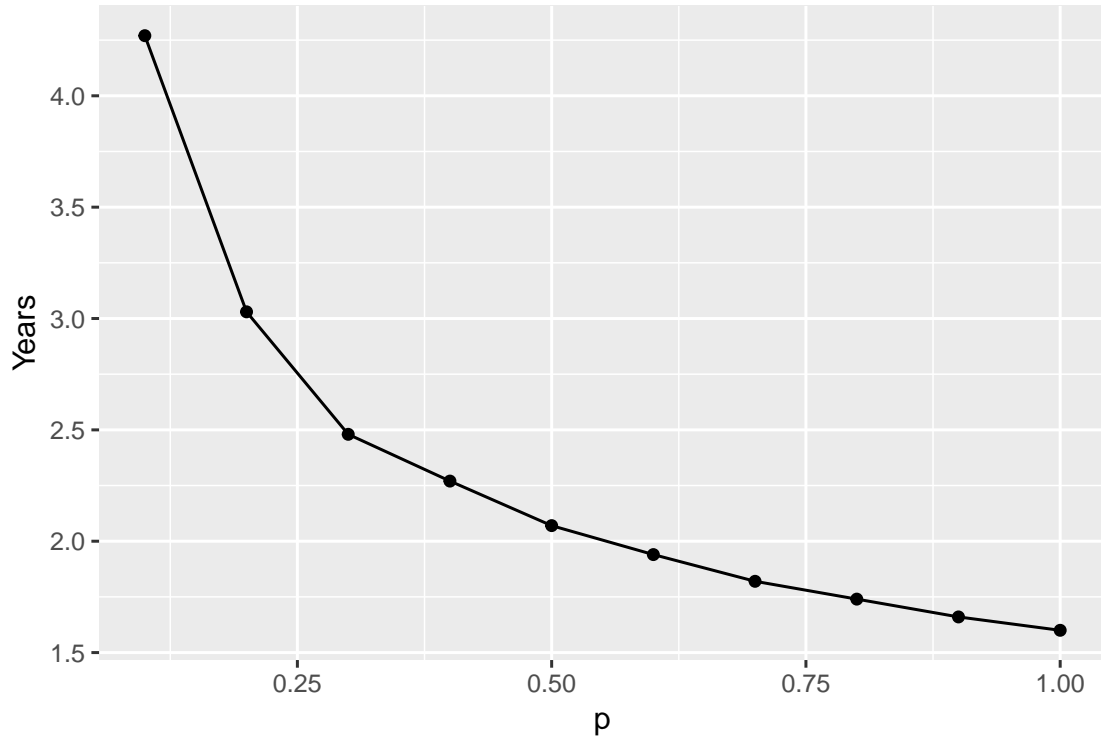


Figure 6: Time to EE as a function of p .

281 If we were to make the equivalent of Figure 5 with the CFP for intentional infection $p_V = 0.01$,
 282 then the time required to have advantages over non-intentional infection is minimal for any p
 283 (the difference in time to advantage is less than a year regardless of the value of p). Figure 7
 284 shows the general relationship between time to advantage and p .

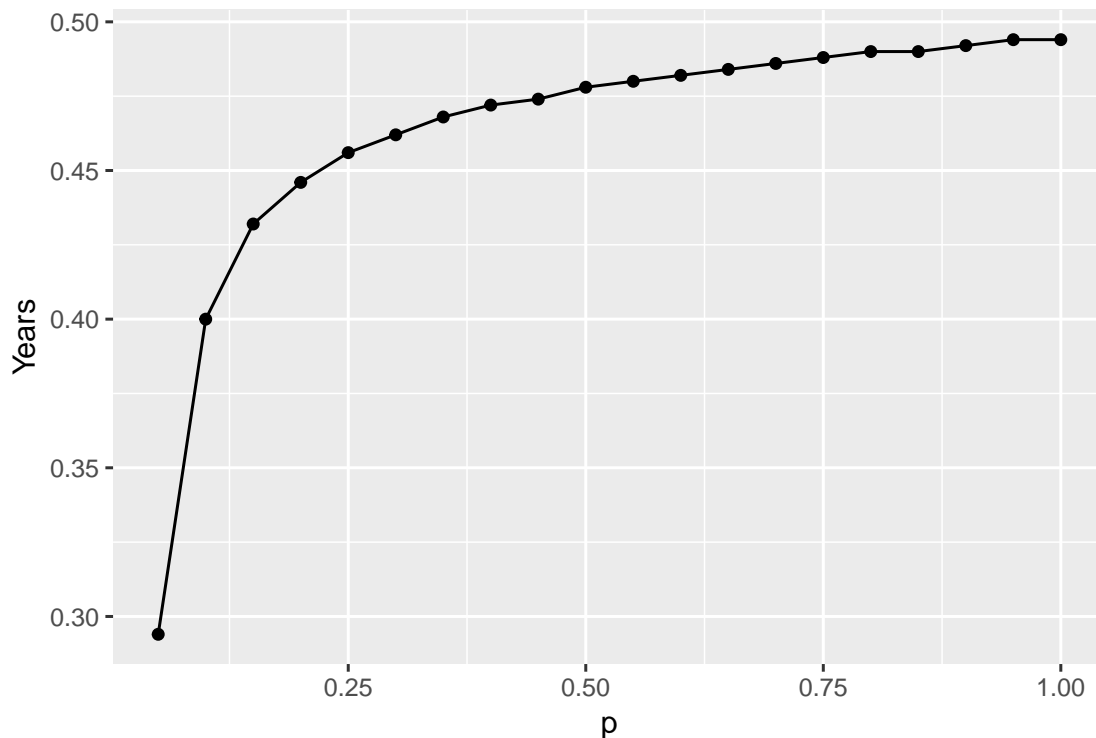


Figure 7: Time to advantage as a function of p . The time required for intentional infection to be more advantageous than non-intentional infection increases as the proportion of intentional infection on newborn increases.

Figure 7 shows that with a lower proportion of intentional infection p , we gain advantage faster. This conclusion is misleading, because it is suggesting that, with a minimal proportion of intentional infection, we can minimize the time it takes to gain advantage. We found that, if p is small, though it gains advantage faster, the number of deaths actually stays very close to non-intentional infection. In other words, although the advantage is obtained quickly, its magnitude is too small to be significant. With this in mind, we will consider one intentional infection strategy (p value) to be “more advantageous” than another if mortality with the strategic value of p is at least 10% lower than mortality with no intentional infection ($p = 0$).

Figure 8: Time required for cumulative mortality of intentional infection becomes at least 10% lower than of non-intentional infection. Under our new definition, with a larger proportion of intentional infection on newborn, intentional infections has fewer death than non-intentional infection in shorter time.

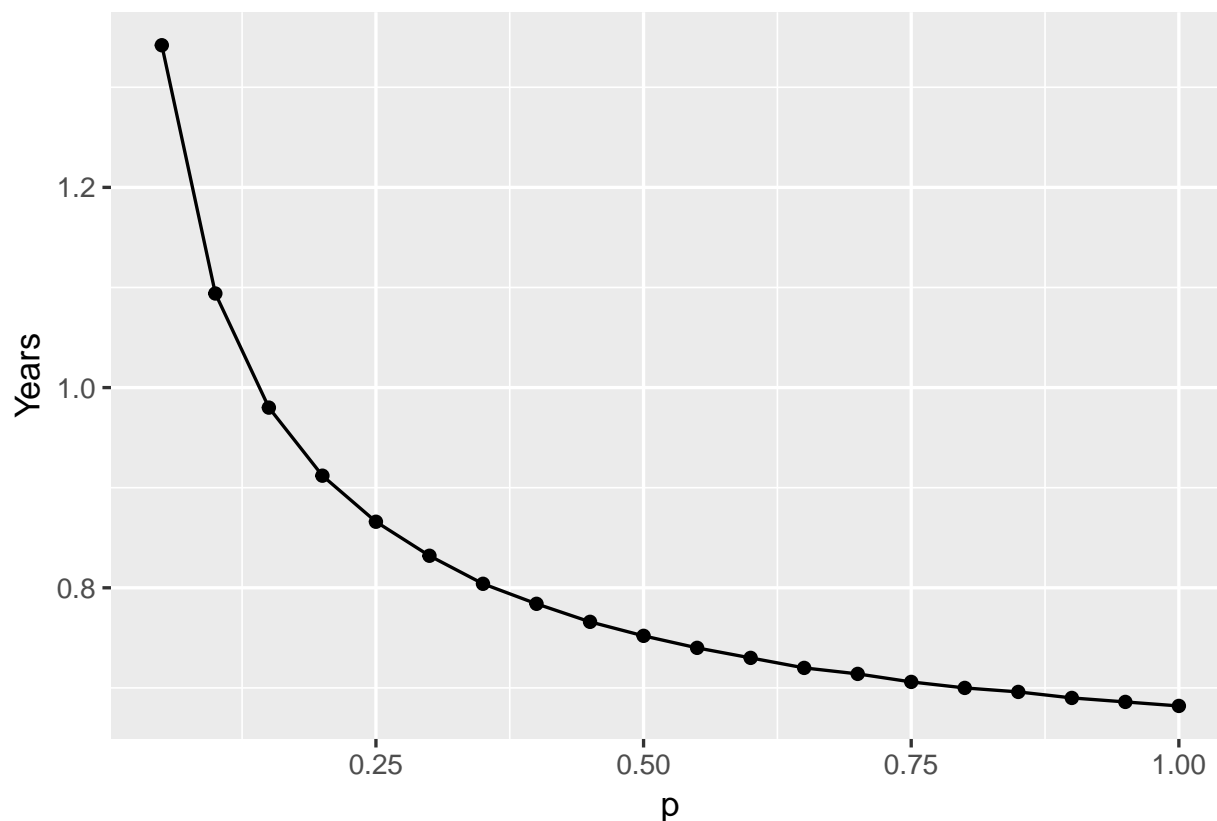


Figure 8 indicates that under our new definition of “more advantageous”, the situation is reversed: a larger proportion of intentional infection yields an advantage more quickly.

2.2.4 Comments and discussion on this model

In this model, we have shown that intentional infection has an advantage over non-intentional infection in terms of total mortality. This suggests, in history, the introduction of intentional infection does have positive effects on disease control, intuitively by reducing the number of deaths. However, as we have discussed in the beginning of [subsubsection 2.2.3](#), a larger

proportion of intentional infection on newborn will have a larger mortality rate once the equilibrium is reached. Though it can reduce total number deaths and become more advantageous than non-intentional infection faster, it is unwise to keep infecting such a large proportion of newborns simply because it will lead to more deaths.

To summarize, if our strategy of intentional infection remains the same over time, in the long run, total mortality could be minimized with a minimal proportion of intentional infection. However, in a real life scenario, strategy does not need to be constant. It may be possible to intentionally infect at a relatively higher proportion initially, and decrease the proportion, or even stop intentional infection. With a combination of strategies like that, we could possibly minimize the total mortality, or even eradicate the disease.

Other than the possibilities of improving on the strategies of intentional infection, there could still be some development to the model itself to make it more suitable to a realistic scenario. For example, since Intentional infection has a lower death rate, we could assume a milder symptom for intentionally infected cases. As a result, because aggressive symptoms are typical routes of disease transmission, lack of such pathways will lead to a decrease in transmission rate. On the other hand, intentionally and naturally infected cases could have different recovery rates. Therefore, our next model could consider various possibilities of these parameters and draw conclusions to its behavior.

2.2.5 Possibility of eradication

If we define the disease as have been eradicated if the proportion of infected is less than one in a million, our model is able to predict possible eradication of the disease.

At EE, $I = 0$. Although I is going to approach 0 asymptotically, it will never completely cease to exist. However, because we know that there is a point in forward time where $I < 1 \times 10^{-6}$, we can claim that naturally infected cases could be eradicated, this will be demonstrated in the next section.

The next question is whether it is possible for intentionally infected cases to burn out. After

326 I is wiped out, we can stop intentionally infecting any other newborns. From then on, if
 327 $S < \frac{1}{\mathcal{R}_0}$ for a long enough period of time, which means the effective \mathcal{R}_0 is less than 1, we
 328 could possibly observe $V < 1 \times 10^{-6}$, which also represent the eradication of intentionally
 329 infected cases.

330 We use one example to illustrate the occurrence of such scenario. Assume we have $p = 1$
 331 until we reach EE, we could calculate \hat{S} and \hat{I} . Now if we set the initial condition as \hat{S} and
 332 \hat{I} we just calculated and let $p = 0$, we will obtain the following after simulation.

S as a function of time, after we stop intentional infection

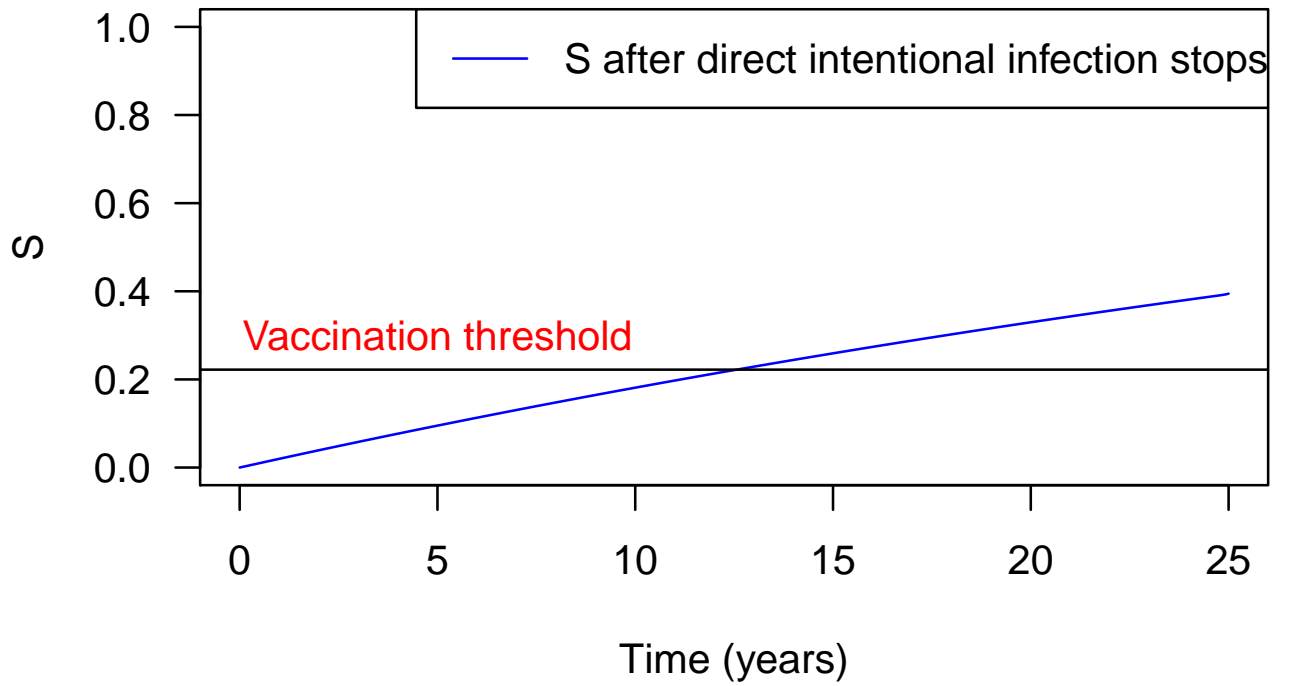


Figure 9: The black line is the vaccination threshold for a disease with $\mathcal{R}_0 = 4.5$. The blue curve represent the dynamics of susceptible, after we stop intentional infection. This figure shows, for more than 10 years after we stop intentional infection, the amount of susceptible stays below the vaccination threshold.

V as a function of time, after we stop intentional infection

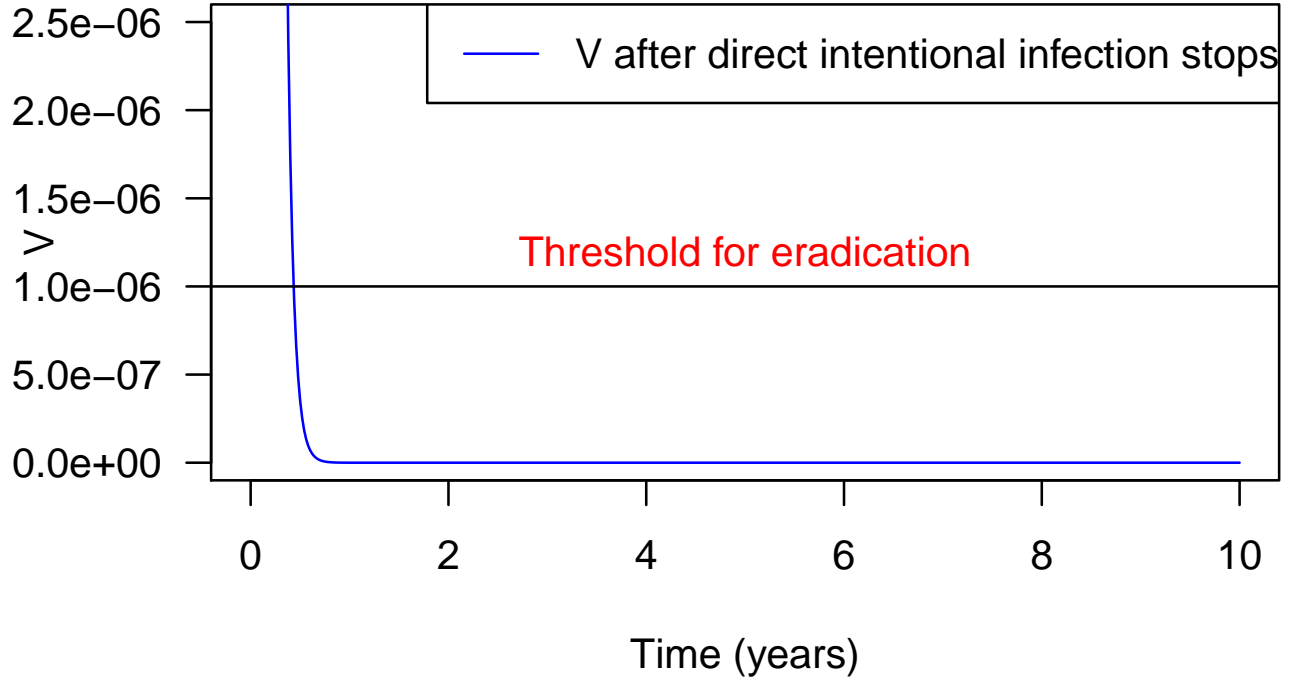


Figure 10: The black line is the pre-defined eradication threshold, $V = 10^{-6}$. The blue curve represents the dynamics of V , after we stop intentional infection. This figure shows that it takes less than 1 year for V to fall below the eradication threshold.

Figure 9 and Figure 10 have shown an example of pre-defined disease eradication. In our example, we assumed $p = 1$, but other values of p will still need to be investigated.

2.3 Model: Different transmission rate and recovery rate

2.3.1 System of differential equations

As mentioned above, we need to involve new parameters for different transmission and recovery rates. Here, let β_V and β_I represent the transmission rate of variolated cases and

339 pathogen infected cases, respectively. γ_V and γ_I are the recovery rate of variolated cases and
 340 pathogen infected cases, respectively.

341 With the new parameters in hand, we now setup our system of differential equation,

$$\begin{aligned}
 \frac{dS}{dt} &= \mu(1-p) - \beta_V SV - \beta_I SI - \mu S, \\
 \frac{dV}{dt} &= \beta_V SV + \mu p - \gamma_V V - \mu V, \\
 \frac{dI}{dt} &= \beta_I SI - \gamma_I I - \mu I, \\
 \frac{dM}{dt} &= p_V \gamma_V V + p_I \gamma_I I, \\
 \frac{dR}{dt} &= (1-p_V) \gamma_V V + (1-p_I) \gamma_I I - \mu R.
 \end{aligned}
 \tag{27}$$

343 We non-dimensionalize [Equation 27](#) by scaling time, by

$$\tau = (\gamma_I + \mu)t.
 \tag{28}$$

345 As the result, we obtain,

$$\frac{dS}{d\tau} = \epsilon(1-p) - \mathcal{R}_{0,V} SV - \mathcal{R}_{0,I} SI - \epsilon S,
 \tag{29a}$$

$$\frac{dV}{d\tau} = \mathcal{R}_{0,V} SV + \epsilon p - \tilde{\gamma} V,
 \tag{29b}$$

$$\frac{dI}{d\tau} = \mathcal{R}_{0,I} SI - I,
 \tag{29c}$$

$$\frac{dM}{d\tau} = p_V(\tilde{\gamma} - \epsilon)V + p_I(1 - \epsilon)I,
 \tag{29d}$$

$$\frac{dR}{d\tau} = (1-p_V)(\tilde{\gamma} - \epsilon)V + (1-p_I)(1 - \epsilon)I - \epsilon R,
 \tag{29e}$$

352 where $\mathcal{R}_{0,V} = \frac{\beta_V}{\gamma_I + \mu}$, $\mathcal{R}_{0,I} = \frac{\beta_I}{\gamma_I + \mu}$, $\tilde{\gamma} = \frac{\gamma_V + \mu}{\gamma_I + \mu}$, $\epsilon = \frac{\mu}{\gamma_I + \mu}$.

353 2.3.2 Equilibria

354 To solve for all equilibria, we set equations [Equation 29a](#), [Equation 29b](#) and [Equation 29c](#)
 355 equal to 0.

We acquired three solutions. However, given conditions that all \hat{S} , \hat{V} and \hat{I} have to be a non-negative number between 0 and 1. The only solution that is biologically meaningful regardless of parameter value is,

$$\hat{S} = \frac{\mathcal{R}_{0,V} + \tilde{\gamma} - \sqrt{\mathcal{R}_{0,V}^2 - 2\mathcal{R}_{0,V}\tilde{\gamma} + \tilde{\gamma}^2 + 4p\mathcal{R}_{0,V}\tilde{\gamma}}}{2\mathcal{R}_{0,V}}, \quad (30a)$$

$$\hat{V} = \frac{\epsilon - \frac{\tilde{\gamma}\epsilon}{\mathcal{R}_{0,V}} + \frac{\epsilon\sqrt{\mathcal{R}_{0,V}^2 - 2\mathcal{R}_{0,V}\tilde{\gamma} + \tilde{\gamma}^2 + 4p\mathcal{R}_{0,V}\tilde{\gamma}}}{\mathcal{R}_{0,V}}}{2\tilde{\gamma}}, \quad (30b)$$

$$\hat{I} = 0. \quad (30c)$$

Similar to our previous models, since infected population at equilibrium is non-zero, it is not a disease free equilibrium but rather an endemic equilibrium.

There exist another solution, but only valid with restrictions on parameter values,

$$\hat{S} = \frac{1}{\mathcal{R}_{0,V}}, \quad (31a)$$

$$\hat{V} = \frac{p\mathcal{R}_{0,V}\epsilon}{-\mathcal{R}_{0,I} + \mathcal{R}_{0,V}\tilde{\gamma}}, \quad (31b)$$

$$\hat{I} = \frac{\mathcal{R}_{0,I}\epsilon - \mathcal{R}_{0,V}\mathcal{R}_{0,I}\epsilon - \mathcal{R}_{0,V}\tilde{\gamma}\epsilon + \mathcal{R}_{0,V}^2\tilde{\gamma}\epsilon - p\mathcal{R}_{0,V}^2\tilde{\gamma}\epsilon}{\mathcal{R}_{0,V}(-\mathcal{R}_{0,I} + \mathcal{R}_{0,V}\tilde{\gamma})}. \quad (31c)$$

This equilibrium will be discussed in later subsection.

Stability analysis rely on Jacobian Matrix,

$$\mathcal{J} = \begin{bmatrix} -\mathcal{R}_{0,V}V - \mathcal{R}_{0,I}I - \epsilon & -\mathcal{R}_{0,V}S & -\mathcal{R}_{0,I}S \\ \mathcal{R}_{0,V}V & \mathcal{R}_{0,V}S - \gamma & 0 \\ \mathcal{R}_{0,I}I & 0 & \mathcal{R}_{0,I}S - 1 \end{bmatrix}. \quad (32)$$

373 Eigenvalues of Jacobian are given as follow,

$$374 \quad \lambda_1 = -1 + \mathcal{R}_{0,I}S \quad (33a)$$

$$375 \quad \lambda_2 = \frac{-\tilde{\gamma} + \mathcal{R}_{0,V}S - \epsilon - \mathcal{R}_{0,V}V - \sqrt{(-\tilde{\gamma} + \mathcal{R}_{0,V}S - \epsilon - \mathcal{R}_{0,V}V)^2 - 4(\mathcal{R}_{0,V}\tilde{\gamma} + \epsilon\tilde{\gamma} - \mathcal{R}_{0,V}S\epsilon)}}{2} \quad (33b)$$

$$376 \quad \lambda_3 = \frac{-\tilde{\gamma} + \mathcal{R}_{0,V}S - \epsilon - \mathcal{R}_{0,V}V + \sqrt{(-\tilde{\gamma} + \mathcal{R}_{0,V}S - \epsilon - \mathcal{R}_{0,V}V)^2 - 4(\mathcal{R}_{0,V}\tilde{\gamma} + \epsilon\tilde{\gamma} - \mathcal{R}_{0,V}S\epsilon)}}{2} \quad (33c)$$

377
378 Looking at the real part of λ_2 and λ_3 , first we look at the terms before square root.

379 By using [Equation 30a](#), we have

$$380 \quad -\tilde{\gamma} + \mathcal{R}_{0,V}S - \epsilon - \mathcal{R}_{0,V}V < 0. \quad (34)$$

381 Therefore,

$$382 \quad \Re(\lambda_2) < -\tilde{\gamma} + \mathcal{R}_{0,V}S - \epsilon - \mathcal{R}_{0,V}V < 0. \quad (35)$$

383 Next, notice

$$384 \quad \mathcal{R}_{0,V}\tilde{\gamma} + \epsilon\tilde{\gamma} - \mathcal{R}_{0,V}S\epsilon > 0. \quad (36)$$

385 It follows that

$$386 \quad \sqrt{(-\tilde{\gamma} + \mathcal{R}_{0,V}S - \epsilon - \mathcal{R}_{0,V}V)^2 - 4(\mathcal{R}_{0,V}\tilde{\gamma} + \epsilon\tilde{\gamma} - \mathcal{R}_{0,V}S\epsilon)} < |-\tilde{\gamma} + \mathcal{R}_{0,V}S - \epsilon - \mathcal{R}_{0,V}V| \quad (37)$$

387 As the result, $\Re(\lambda_3) < 0$.

388 Lastly, we look at λ_1 . Here we noticed that $\Re(\lambda_1)$ is not necessarily negative as it depends
389 on all other parameters. For the purpose of our interest, we would like to use specific value
390 of smallpox to simulate the result.

391 We set $\mathcal{R}_{0,I} = 4.5$. As $\mathcal{R}_{0,V}$ should be logically less than $\mathcal{R}_{0,I}$, we set $\mathcal{R}_{0,V} = 2.5$, and plotted
392 the (γ, p) plane.

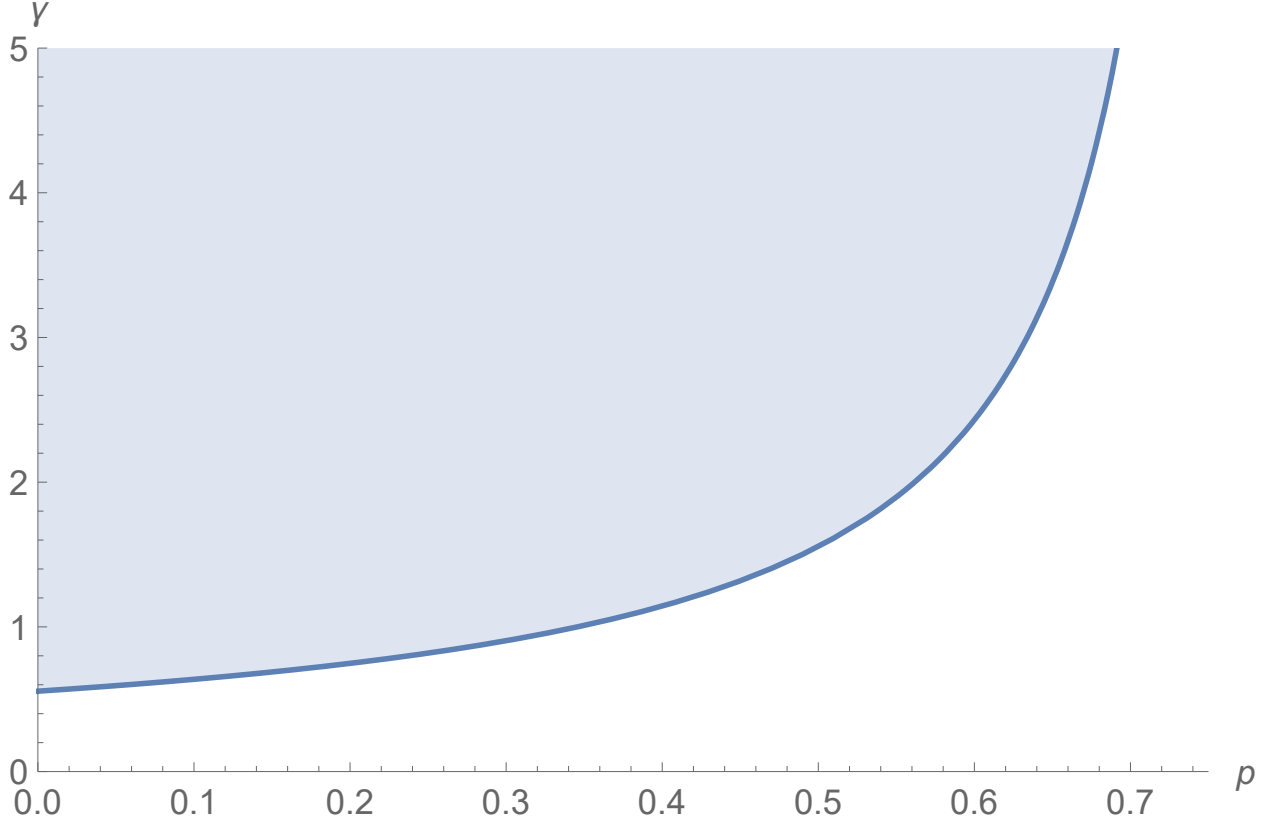


Figure 11: (γ, p) plane when $\mathcal{R}_{0,V} = 2.5, \mathcal{R}_{0,I} = 4.5$. Shaded area is the region where $\Re(\lambda_1) < 0$, meaning the system is locally asymptotically stable.

It is necessary to investigate the dynamics when EE is unstable. Recall the other equilibrium which is only valid for some parameter values. We discovered that, at the boundary condition in Figure 11, our first equilibrium loses stability. But we also found that the second equilibrium becomes biologically meaningful at the same time. In order to identify whether the second equilibrium is stable, we compute the eigenvalues of the Jacobian again,

$\Re(\lambda_2)$ and $\Re(\lambda_3)$ are still both negative.

$$\Re(\lambda_1) = -1 + \frac{\mathcal{R}_{0,V}}{\mathcal{R}_{0,I}}. \quad (38)$$

With our intuition that $\mathcal{R}_{0,V} < \mathcal{R}_{0,I}$, we can conclude that $\Re(\lambda_1) < 0$. Therefore, when the first equilibrium is unstable, the system approaches another equilibrium. Hence, the stability

of the system switches between two equilibria at the boundary of [Figure 11](#).

2.3.3 Comments and discussion on this model

Parameters of this model such as $\mathcal{R}_{0,V}$ and γ_V can be immensely different depending on the design of the pathogen used for intentional infection.

Previous studies have shown that, for a transmissible vaccine, in a similar model which does not consider natural birth and death, as well as no disease induced mortality. Even if $\mathcal{R}_{0,V}$ is less than 1, such as $\mathcal{R}_{0,V} = 0.5$, a significant decrease in the final size of an epidemics is expected [7]. In our model, birth and death rate are considered. Knowing that in our system, the disease can never be eradicated, there will always be newly infected cases. For that reason, we cannot estimate final size for this model. But we are expecting that, the cumulative disease induced mortality will be reduced.

3 Model: Intentionally infect susceptible with a certain rate

3.1 Model: Modification to SIR model

3.1.1 System of differential equations

Our second strategy is to intentionally infect susceptible individuals with a certain rate. Again, we start our analysis by modifying the standard SIR model.

The following assumptions are made:

- No disease induced mortality.
- Birth and natural death rate are the same, the total population remains constant.

- Latent period is short enough to be ignored.
- All susceptible individuals are equally likely to be infected, and all infected individuals are equally infectious.

$$\begin{aligned}
\frac{dS}{dt} &= \mu - \beta SI - rS - \mu S, \\
\frac{dI}{dt} &= rS + \beta SI - \gamma I - \mu I, \\
\frac{dR}{dt} &= \gamma I - \mu R.
\end{aligned} \tag{39}$$

Here, β is transmission rate, γ is recovery rate, μ is the *per capita* rate of birth and death, and r is the rate of intensional infection on susceptible individuals. We will discuss the reasonable value of r in later sections.

By scaling time, we convert our system into dimensionless form, using:

$$\tau = (\gamma + \mu)t. \tag{40}$$

The dimensionless system we arrive at is:

$$\begin{aligned}
\frac{dS}{d\tau} &= \epsilon - \mathcal{R}_0 SI - \eta S - \epsilon S, \\
\frac{dI}{d\tau} &= \eta S + \mathcal{R}_0 SI - I, \\
\frac{dR}{d\tau} &= (1 - \epsilon)I - \epsilon R,
\end{aligned} \tag{41}$$

where $\epsilon = \frac{\mu}{\gamma + \mu}$, $\mathcal{R}_0 = \frac{\beta}{\gamma + \mu}$, $\eta = \frac{r}{\gamma + \mu}$.

3.1.2 Equilibrium

We obtain only one solution by solving the system:

$$\hat{S} = \frac{1}{\mathcal{R}_0} - \frac{2\eta}{\mathcal{R}_0(-(\eta + \epsilon - \epsilon\mathcal{R}_0) + \sqrt{(\eta + \epsilon - \epsilon\mathcal{R}_0)^2 + 4\mathcal{R}_0\epsilon\eta} + 2\eta)} \quad (42a)$$

$$\hat{I} = \frac{-(\eta + \epsilon - \epsilon\mathcal{R}_0) + \sqrt{(\eta + \epsilon - \epsilon\mathcal{R}_0)^2 + 4\mathcal{R}_0\epsilon\eta}}{2\mathcal{R}_0} \quad (42b)$$

Note for any non-zero η , $\hat{I} \neq 0$, it follows that this equilibrium is also an endemic equilibrium, since there is always a proportion of population that are infected.

Stability analysis rely on Jacobian matrix, which is,

$$\mathcal{J} = \begin{bmatrix} -\mathcal{R}_0 I - \eta - \epsilon & -\mathcal{R}_0 S \\ \eta + \mathcal{R}_0 I & \mathcal{R}_0 S - 1 \end{bmatrix}. \quad (43)$$

For simplicity. Let

$$G = -(\eta + \epsilon - \epsilon\mathcal{R}_0) + \sqrt{(\eta + \epsilon - \epsilon\mathcal{R}_0)^2 + 4\mathcal{R}_0\epsilon\eta}. \quad (44)$$

Notice, if $\epsilon, \eta \neq 0$, $G > 0$.

So Jacobian at EE becomes,

$$\mathcal{J}|_{E.E.} = \begin{bmatrix} \frac{G}{2} - \eta - \epsilon & -1 + \frac{2\eta}{G+2\eta} \\ \eta + \frac{G}{2} & -\frac{2\eta}{G+2\eta} \end{bmatrix}. \quad (45)$$

The eigenvalues of of Jacobian are:

$$\lambda_{1,2} = \frac{-(G^2 + 4\eta G + 2\epsilon G + 4\eta^2 + 4\epsilon\eta + 4\eta)}{4(G + 2\eta)} \quad (46)$$

$$\pm \frac{\sqrt{((G^2 + 4\eta G + 2\epsilon G + 4\eta^2 + 4\epsilon\eta + 4\eta)^2 - 4(2G^3 + 12\eta G^2 + 24\eta^2 G + 8\epsilon\eta G + 16\eta^3 + 16\epsilon\eta^2))}}{4(G + 2\eta)} \quad (47)$$

We know that $G > 0$, and since the rate of intentional infection is non-zero, $\eta > 0$.

As we are only interested in the case where the rate of intentional infection is non-zero, we assume that $\eta > 0$. Together with the fact that $G > 0$, the discriminant(Δ) satisfies the following inequality,

$$\Delta < |(G^2 + 4\eta G + 2\epsilon G + 4\eta^2 + 4\epsilon\eta + 4\eta)| \quad (48)$$

Therefore, we can conclude that $\text{Re}(\lambda) < 0$, which means, EE is locally asymptotically stable.

For global stability analysis, we apply the same method as in [subsubsection 2.2.2](#), which is by using distance function.

The identical distance function was used to run the simulation, and the result is obtained in the following plot:

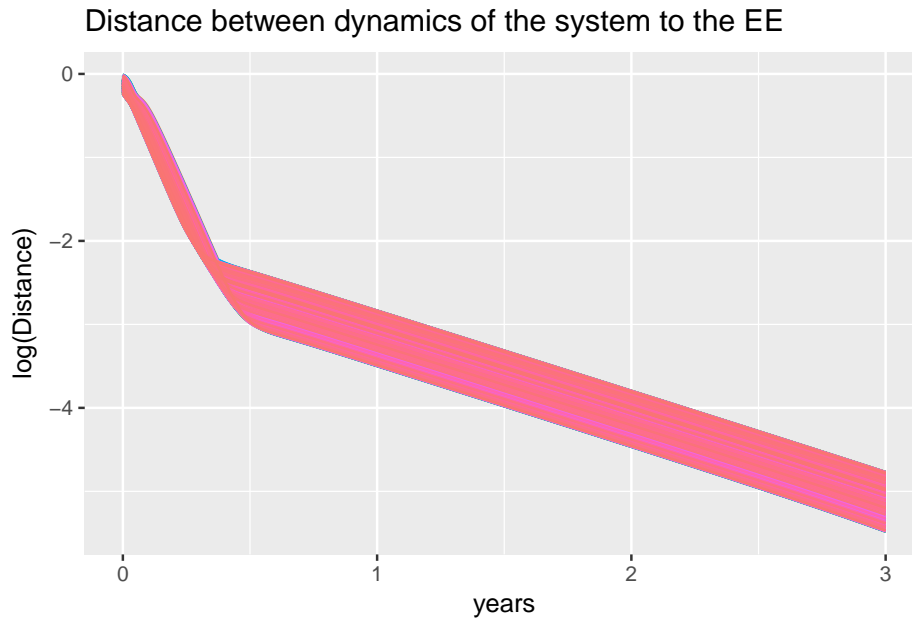


Figure 12: Distance between dynamics of the system to the EE, as a function of time. If EE is GAS, distance function should approach 0, regardless of which initial condition is chosen. This figure shows the distance function of 1000 random initial conditions. Similar to the newborn model, in the long run, the distance decays exponentially. We have enough confidence to conclude that EE is GAS.

To fully understand the dynamics of the system, we need to identify whether or not there is

a damped oscillation. This is done by determining the sign of the discriminant.

Our analysis begins with specific values of each parameter. Again, we will use parameters of smallpox. The values are listed in Table 1.

3.1.3 Region of (\mathcal{R}_0, η) plane where there are damped oscillations (fixed ϵ)

η is the rate of intentional infection, \mathcal{R}_0 is the basic reproduction number.

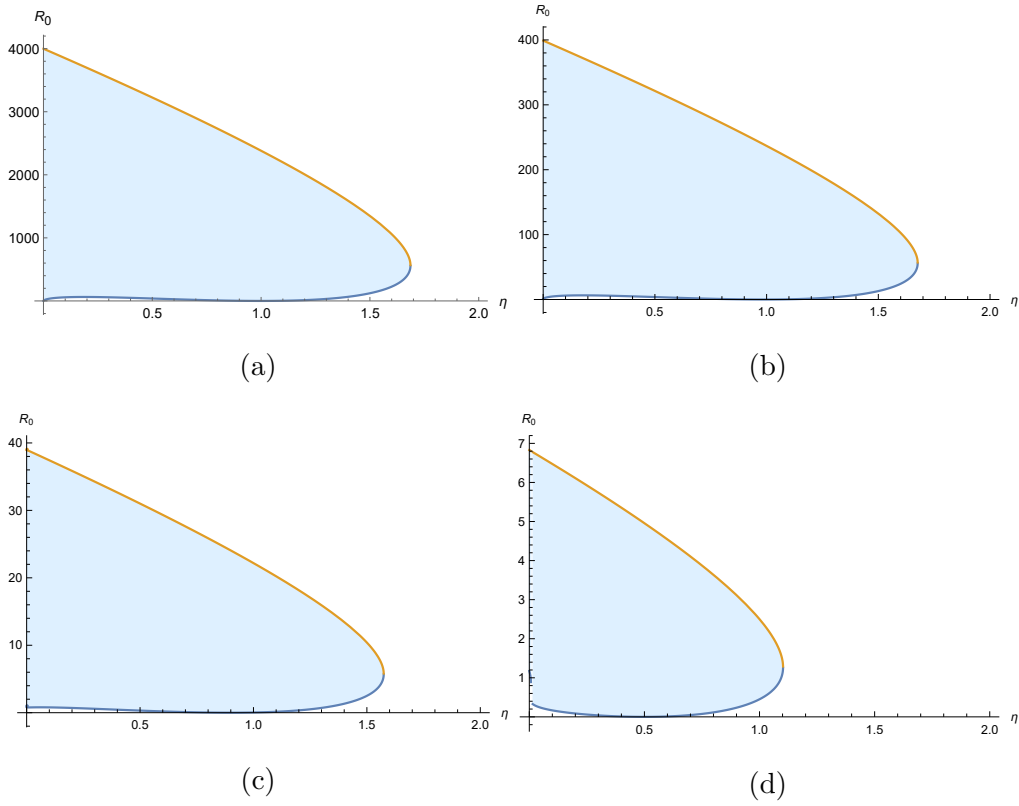


Figure 13: Panel (a)-(d) show (\mathcal{R}_0, η) plane for different ϵ values. The shaded region represents the range of conditions where the system has damped oscillations.

a) $\epsilon = 0.001$, b) $\epsilon = 0.01$, c) $\epsilon = 0.1$, d) $\epsilon = 0.5$

3.1.4 Comments and discussion on this model

Similar to our newborn infection model, the model obtained by modifying the standard SIR model is rather simple. We still need to divide the infected compartment into intentionally and naturally infected classes.

Furthermore, to help us conclude whether or not there is any positive effect by intentionally infecting the population, we need to involve disease induced mortality. This help us to compare two different strategies of intentional infection.

3.2 Model: Addition of disease induced mortality

3.2.1 System of differential equations

The addition of disease induced mortality requires the involvement of case fatality proportion.

Here, we are applying the exact same adjustment as we did in [subsection 2.2](#). Our new system is,

$$\frac{dS}{dt} = \mu - \beta S(V + I) - rS - \mu S, \quad (49a)$$

$$\frac{dV}{dt} = \beta SV + rS - \gamma V - \mu V, \quad (49b)$$

$$\frac{dI}{dt} = \beta SI - \gamma I - \mu I, \quad (49c)$$

$$\frac{dM}{dt} = p_V \gamma V + p_I \gamma I, \quad (49d)$$

$$\frac{dR}{dt} = (1 - p_V) \gamma V + (1 - p_I) \gamma I - \mu R, \quad (49e)$$

where β is transmission rate, γ is recovery rate, μ is the *per capita* rate of birth and death, and r is the rate of intensional infection on susceptible individuals.

We non-dimensionalize [Equation 49](#) by scaling time, by

$$\tau = (\gamma + \mu)t. \quad (50)$$

As the result, we obtain the following:

$$\frac{dS}{d\tau} = \epsilon - \eta S - \mathcal{R}_0 S(V + I) - \epsilon S, \quad (51a)$$

$$\frac{dV}{d\tau} = \mathcal{R}_0 S V + \eta S - V, \quad (51b)$$

$$\frac{dI}{d\tau} = \mathcal{R}_0 S I - I, \quad (51c)$$

$$\frac{dM}{d\tau} = p_V(1 - \epsilon)V + p_I(1 - \epsilon)I, \quad (51d)$$

$$\frac{dR}{d\tau} = (1 - p_V)(1 - \epsilon)V + (1 - p_I)(1 - \epsilon)I - \epsilon R, \quad (51e)$$

where $\epsilon = \frac{\mu}{\gamma + \mu}$, $\mathcal{R}_0 = \frac{\beta}{\gamma + \mu}$, $\eta = \frac{r}{\gamma + \mu}$

3.2.2 Equilibria

If $\eta = 0$, the system will again be simplified to the standard SIR model, the equilibrium of that was previously shown. Here, we are only interested in the case where $\eta \neq 0$.

The only solution we obtain by solving the system is:

$$\hat{S} = \frac{1}{\mathcal{R}_0} - \frac{2\eta}{\mathcal{R}_0(-(\eta + \epsilon - \epsilon\mathcal{R}_0) + \sqrt{(\eta + \epsilon - \epsilon\mathcal{R}_0)^2 + 4\mathcal{R}_0\epsilon\eta} + 2\eta)}, \quad (52a)$$

$$\hat{V} = \frac{-(\eta + \epsilon - \epsilon\mathcal{R}_0) + \sqrt{(\eta + \epsilon - \epsilon\mathcal{R}_0)^2 + 4\mathcal{R}_0\epsilon\eta}}{2\mathcal{R}_0}, \quad (52b)$$

$$\hat{I} = 0, \quad (52c)$$

Notice, \hat{V} is non-zero, therefore, this equilibrium is not a disease free equilibrium. It follows that it is the endemic equilibrium.

3.2.3 Stability of Endemic Equilibrium

The Jacobian matrix of this system is,

$$\mathcal{J} = \begin{bmatrix} -\eta - \mathcal{R}_0(V + I) - \epsilon & -\mathcal{R}_0S & -\mathcal{R}_0S \\ \mathcal{R}_0V + \eta & \mathcal{R}_0S - 1 & 0 \\ \mathcal{R}_0I & 0 & \mathcal{R}_0S - 1 \end{bmatrix}. \quad (53)$$

Eigenvalues of Jacobian are ,

$$\lambda_1 = -1 + \mathcal{R}_0S \quad (54a)$$

$$\lambda_2 = \frac{-1 + \mathcal{R}_0S - \eta - \epsilon - \mathcal{R}_0V + \sqrt{(-1 + \mathcal{R}_0S - \eta - \epsilon - \mathcal{R}_0V)^2 - 4(\eta + \mathcal{R}_0V + \epsilon - \mathcal{R}_0S\epsilon)}}{2} \quad (54b)$$

$$\lambda_3 = \frac{-1 + \mathcal{R}_0S - \eta - \epsilon - \mathcal{R}_0V - \sqrt{(-1 + \mathcal{R}_0S - \eta - \epsilon - \mathcal{R}_0V)^2 - 4(\eta + \mathcal{R}_0V + \epsilon - \mathcal{R}_0S\epsilon)}}{2} \quad (54c)$$

By using Equation 52a and Equation 54a, we get

$$\Re(\lambda_1) = -1 + \mathcal{R}_0S = -\frac{2\eta}{(-(\eta + \epsilon - \epsilon\mathcal{R}_0) + \sqrt{(\eta + \epsilon - \epsilon\mathcal{R}_0)^2 + 4\mathcal{R}_0\epsilon\eta + 2\eta})} < 0 \quad (55)$$

To decide the real parts of λ_2 and λ_3 , we need to determine the sign of the quantity under the square root.

By using Equation 52a again, we have

$$\mathcal{R}_0S\epsilon < \epsilon. \quad (56)$$

Therefore

$$(\eta + \mathcal{R}_0V + \epsilon - \mathcal{R}_0S\epsilon) > 0. \quad (57)$$

It follows that

$$\sqrt{(-1 + \mathcal{R}_0S - \eta - \epsilon - \mathcal{R}_0V)^2 - 4(\eta + \mathcal{R}_0V + \epsilon - \mathcal{R}_0S\epsilon)} < |(-1 + \mathcal{R}_0S - \eta - \epsilon - \mathcal{R}_0V)|. \quad (58)$$

525 We have two cases. If the quantity under square root is negative, then

$$526 \quad \Re(\lambda_2) = \Re(\lambda_3) = -1 + \mathcal{R}_0 S - \eta - \epsilon - \mathcal{R}_0 V < 0. \quad (59)$$

527 Otherwise, we have

$$528 \quad \Re(\lambda_3) < \Re(\lambda_2) < 0. \quad (60)$$

529 Thus, for any $\eta > 0$, we can conclude that the EE is locally asymptotically stable regardless
530 of the values of other parameters.

531 3.2.4 The effect of intentional infection on mortality

532 We will again start by looking at the mortality rate at EE.

533 By inserting \hat{V} [Equation 52b] and $\hat{I} (= 0)$ into Equation 51, we obtain the mortality rate at
534 EE,

$$535 \quad \frac{dM}{d\tau} = p_V(1 - \epsilon)V = \frac{p_V(1 - \epsilon)\epsilon(\mathcal{R}_0 - 1) + p_V(1 - \epsilon)\epsilon\sqrt{(\mathcal{R}_0 - 1)^2 + 4\mathcal{R}_0 p}}{2\mathcal{R}_0}. \quad (61)$$

536 As shown the following graph,

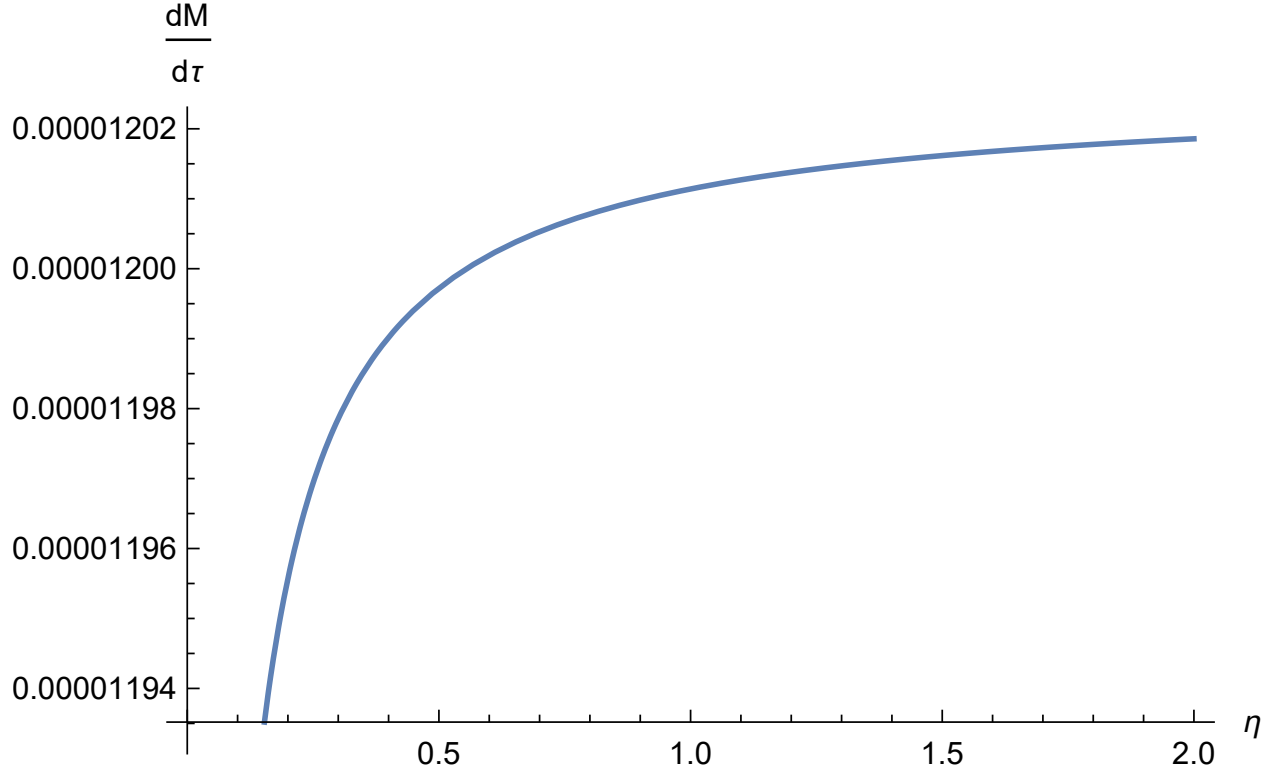


Figure 14: $\frac{dM}{d\tau}$ at EE as a function of η . $\frac{dM}{d\tau}$ increases as the rate of intentional infection on susceptible increases. The magnitude of $\frac{dM}{d\tau}$ eventually reaches an asymptote at $\frac{dM}{d\tau} = 0.00001202$.

An important observation is, as the rate of intentional infection increases, the mortality rate eventually reaches an asymptote at around 1.2×10^{-5} , which is very close to the mortality rate when we intentionally infect newborn individuals with a highest proportion ($p = 1$). This suggests, in the long run, intentionally infect susceptible at a high rate would have similar results as a high proportion of newborn infection. A logical explanation to that is, when $p = 1$ for newborn infection model or when r is very high for susceptible model, at EE, there is almost no individuals that are still susceptible ($\hat{S} = 0$ or $\hat{S} \approx 0$). The only individuals entering the system are the newborns, but newborn model suggests 100% of them will be intentionally infected before becoming susceptible. Similarly, susceptible model will intentionally infect almost all of the newborns immediately after they become susceptible. As a result, the increasing rates of V (intentionally infected) are essentially the same for both approaches. Therefore, due to the fact that $\frac{dM}{d\tau}$ is directly proportional to \hat{V} , the magnitude

549 of $\frac{dM}{d\tau}$ are similar.

550 As we did in the newborn model, we also want to know the time it takes for susceptible
551 intentional infection to reach new equilibrium, as well as the time it takes to be advantageous
552 over non-intentional infection.

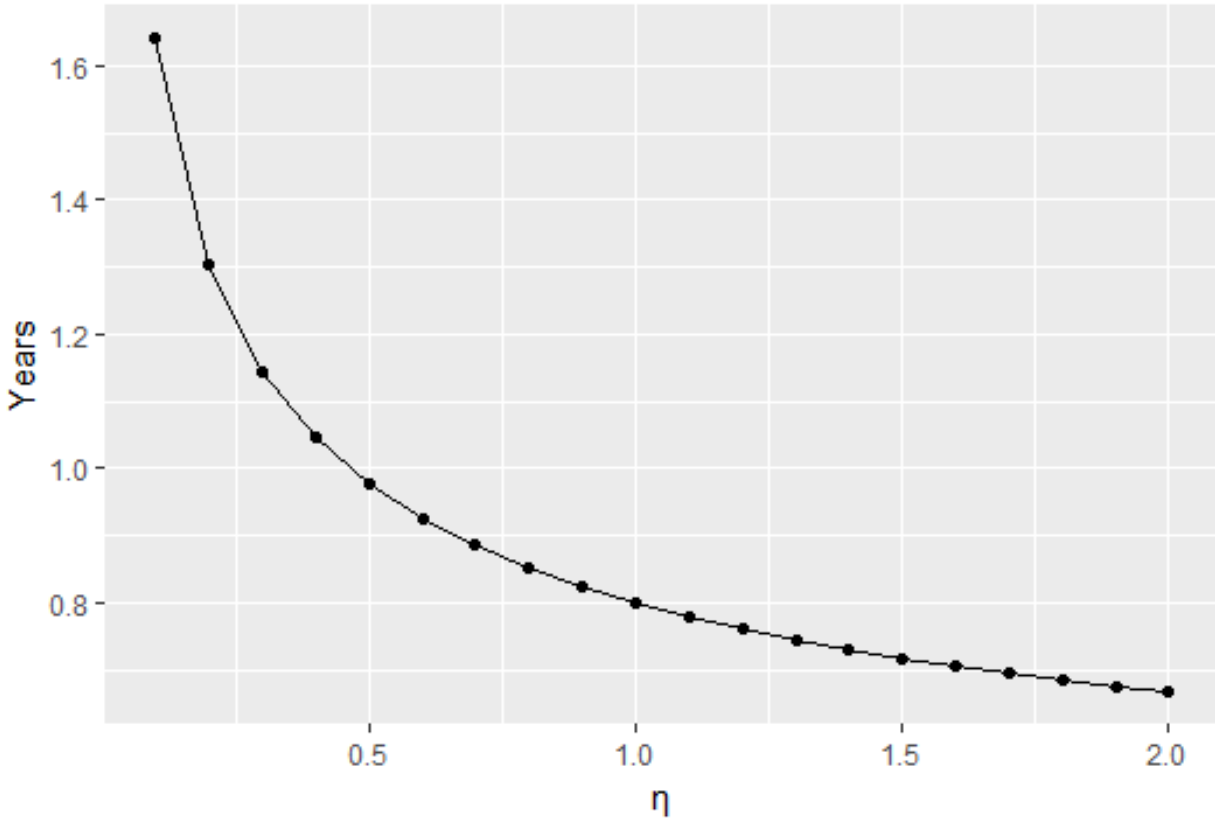


Figure 15: Time taken for susceptible intentional infection to reach the new equilibrium

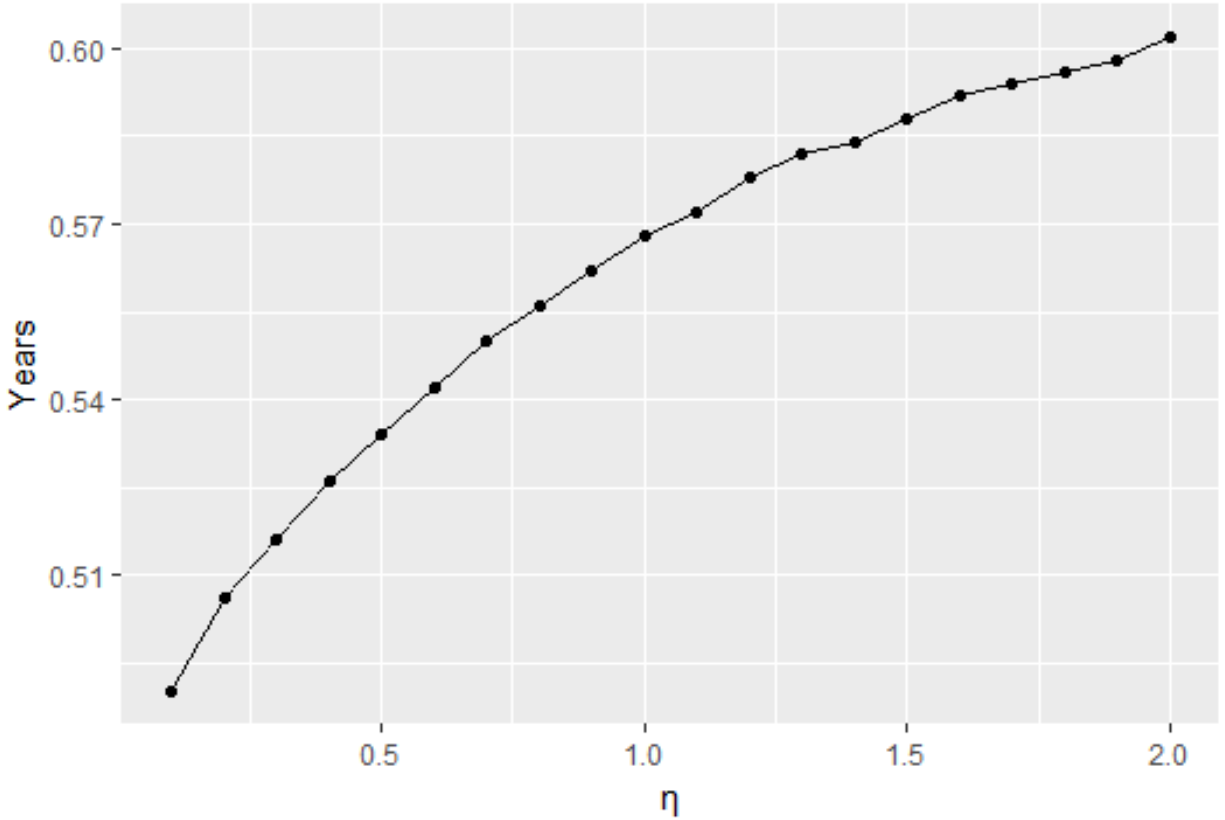


Figure 16: Time taken for susceptible intentional infection to have at least 10% fewer disease induced mortality, as a function of rate of intentional infection on susceptible.

Recall in [subsection 2.2.3](#), we initially found that in the newborn infection model, with a larger proportion of newborn intentional infection, it becomes more advantageous over non-intentional infection more slowly. But after we change the definition of “more advantageous” to be “The cumulative mortality rate of the system with intentional infection is at least 10% less than of the system with no intentional infection”, the situation was reversed. However, unlike the newborn infection model, in susceptible infection, the situation remain the same even after the definition of “more advantageous” was changed. The time required for intentional infection to be more advantageous still increases as the rate of intentional infection increases. This means, with a very low rate of intentional infection on susceptible, not only can we ensure a lower cumulative mortality in the long run, but also can the model be advantageous more quickly. This also means that we cannot conclude an optimal rate of

intentional infection on susceptible.

3.2.5 Comments and discussion to this model

This model has proven the benefit of intentional infection with the approach of infecting susceptible individuals. We have shown that, with any positive rate of intentional infection on susceptible, the cumulative disease induced mortality can always be reduced, compared to not doing so. However, as discussed in the previous section, we cannot conclude an optimal rate of intentional infection. It is because we have shown that the benefit is maximized, in every aspect, when the rate of intentional infection is minimal. Further study is required to answer this question.

4 Discussion

We have used several simple models to illustrate that both approaches of intentional infection could help restrain epidemics by reducing the number of total death caused by life-threatening diseases, in our case, smallpox. This suggests, practices of smallpox variolation in history have indeed saved many lives.

In addition to demonstrating the utility of intentional infection, one of our models have also shown that, under certain circumstances, the disease could be completely eradicated. Future research can perform stochastic analysis using similar systems of differential equations, in order to rigorously prove its possibility.

Even though our results have revealed that intentional infection can be an effective tool against epidemics, our mathematical models rely on several important assumptions. First, we assume that recovered individuals acquires permanent immunity against this infectious disease. Although unrealistic for many diseases such as flu. It is still applicable to many others, specifically diseases such that infection could provide long term immunity up to several decades. For example, measles, whooping cough, etc [1]. Secondly, in addition to

our first assumption, possibility of evolution was ignored in our models, which is extremely common in viruses, as they mutate much faster than most of the organisms on earth [8]. Mutations may lead to different strains of viruses, thus the old immunity may not be effective anymore. Finally, we assumed a single homogeneous population, which does not account for spatial, seasonal and individual heterogeneity.

Moreover, due to the existence of special events or accidents in realistic scenarios, patterns of disease spread could be significantly altered, meaning we have many different factor not yet being considered in our models. For both approaches, the models can still be modified and developed to higher complexity. For instance, in real life, smallpox variolation is done via skin. But since the subject used to variolate is live smallpox virus, there could be other pathways for live smallpox virus to enter respiratory system of the variolated individual, or of the other people who has physically contacted such individual. A specific example of such pathway could be touching the area of variolation, followed by touching nose or mouth.

In such scenario, though the variolated individual might acquire symptoms much less severe than the symptoms of natural infections, when virus happens to enter respiratory system, a natural infection with much more serious symptoms could still occur. Therefore, possibility still exists for a variolated individual to become gravely ill. In our construction of the models, the consequence of that will be, a proportion of variolated individuals (V) entering naturally infected class (I).

It is meaningful to compare the utility of intentional infection to traditional vaccination. Unlike intentional infection, most traditional vaccines are unable to transmit from one individual to another. Intuitively, if intentional infection and traditional vaccination have the same pattern of distribution, the former could spread faster than the latter. Nevertheless, the trade-off for intentional infection is to have a proportion of people still die after being intentionally infected, whereas vaccination is assumed to provide a perfect protection against infectious disease.

We may also investigate the potential of combining intentional infection and traditional vaccination, which is known as transmissible vaccine. It is a vaccine that is capable of

spreading from one individual to another, therefore utilizes the advantage of intentional infection, as well as providing perfect protection.

At last, we could compare the results of our models to historical data of smallpox. This may help us understand the variolation history of smallpox, which can innovate us in developing safer and more effective methods for disease prevention and control.

5 Appendix

Figures used in this document were created by “R” or “Mathematica (Version 11.3)”. The files used to construct figures are under the folder named “Supplementary material”

References

- [1] Ian J Amanna, Nichole E Carlson, and Mark K Slifka. “Duration of humoral immunity to common viral and vaccine antigens”. In: *New England Journal of Medicine* 357.19 (2007), pp. 1903–1915.
- [2] Barry R Bloom et al. “Priorities for global research and development of interventions”. In: (2006).
- [3] Gavin Dabrera et al. “A case-control study to estimate the effectiveness of maternal pertussis vaccination in protecting newborn infants in England and Wales, 2012–2013”. In: *Clinical Infectious Diseases* 60.3 (2014), pp. 333–337.
- [4] Klaus D Elgert. *Immunology: understanding the immune system*. John Wiley & Sons, 2009.
- [5] Donald A Henderson. “The eradication of smallpox”. In: *Scientific American* 235.4 (1976), pp. 25–33.
- [6] Angela Ki Che Leung. ““Variolation” and Vaccination in Late Imperial China, Ca 1570–1911”. In: *History of vaccine development*. Springer, 2011, pp. 5–12.

- 639 [7] Scott L Nuismer et al. “Controlling epidemics with transmissible vaccines”. In: *PloS*
640 *one* 13.5 (2018), e0196978.
- 641 [8] Robert H Purcell and AJ Zuckerman. “Hepatitis B virus mutants and efficacy of vacci-
642 nation”. In: *Lancet* 356.9231 (2000), pp. 769–769.
- 643 [9] Pieter Streefland, AMR Chowdhury, and Pilar Ramos-Jimenez. “Patterns of vaccination
644 acceptance”. In: *Social science & medicine* 49.12 (1999), pp. 1705–1716.