The Impacts of Simultaneous Disease Intervention Decisions on Epidemic Outcomes

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Abstract

Mathematical models of the interplay between disease dynamics and human behavioural dynamics can improve our understanding of how diseases spread when individuals adapt their behaviour in response to an epidemic. Accounting for behavioural mechanisms that determine uptake of infectious disease interventions such as vaccination and non-pharmaceutical interventions (NPIs) can significantly alter predicted health outcomes in a population. However, most previous approaches that model interactions between human behaviour and disease dynamics have modelled behaviour of these two interventions separately. Here, we develop and analyze an agent based network model to gain insights into how behaviour toward both interventions interact adaptively with disease dynamics (and therefore, indirectly, with one another) during the course of a single epidemic where an SIRV infection spreads through a contact network. In the model, individuals decide to become vaccinated and/or practice NPIs based on perceived infection prevalence (locally or globally) and on what other individuals in the network are doing. We find that introducing adaptive NPI behaviour lowers vaccine uptake on account of behavioural feedbacks, and also decreases epidemic final size. When transmission rates are low, NPIs alone are as effective in reducing epidemic final size as NPIs and vaccination combined. Also, NPIs can compensate for delays in vaccine availability by hindering early disease spread, decreasing epidemic size significantly compared to the case where NPI behaviour does not adapt to mitigate early surges in infection prevalence. We also find that including adaptive NPI behaviour strongly mitigates the vaccine behavioural feedbacks that would otherwise result in higher vaccine uptake at lower vaccine efficacy as predicted by most previous models, and the same feedbacks cause epidemic final size to remain approximately constant across a broad range of values for vaccine efficacy. Finally, when individuals use local information about others' behaviour and infection prevalence, instead of population-level information, infection is controlled more efficiently through ring vaccination, and this is reflected in the time evolution of pair correlations on the network. This model shows that accounting for both adaptive NPI behaviour and adaptive vaccinating behaviour regarding social effects and infection prevalence can result in qualitatively different predictions than if only one type of adaptive behaviour is modelled.

Keywords: Epidemic Modelling, Vaccinating Behaviour, Non Pharmaceutical Interventions, Adaptive networks, Econophysics

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1 1. Introduction

Infectious disease outbreaks have the potential to cause unexpected burdens and panic in societies. For example, the outbreak of severe acute respiratory syndrome (SARS) in 2003 caused significant economic impacts across the world, despite lasting only six months [1]. Occurring unexpectedly, outbreaks such as the aforementioned SARS outbreak [1, 2], the Middle East respiratory syndrome outbreak in 2012 [3], Ebola outbreak in 2014 [4], or an influenza pandemic, which has happened as recently as 2009 [5], can be difficult to predict and can spread locally or globally and last anywhere from months to years.

Human behaviour can have a large impact on the spread of infectious diseases [6]. People have g been observed to change their regular social routines in response to an epidemic, in order to reduce 10 their risk of becoming infected [7, 8, 9]. The infection prevalence or incidence of a disease in a com-11 munity serves to drive these behavioural changes, as an individual's perceived susceptibility generally 12 rises along with these disease measures [6, 10, 11, 12]. There are two primary self protective interven-13 tion strategies susceptible members of a population can utilize to reduce their chances of contracting 14 a disease. These are pharmaceutical interventions, such as vaccination, and non-pharmaceutical 15 interventions (NPIs), such as social distancing and increased hand washing [13]. The usage of these 16 intervention strategies are voluntary in many health jurisdictions, and so perceived risks play an 17 important role in how often they are utilized [14]. 18

Coupled disease-behaviour models combine human decision making behaviour with traditional 19 transmission dynamics, helping to capture an additional, and often important, aspect of disease 20 spread [6, 15, 16]. Behaviourally based models that incorporate NPIs and social distancing during 21 an outbreak show that these practices can lower the attack rate of a disease [17, 18, 19, 20, 21, 22 22, 23, 24. Suppressing an outbreak using these means can be very critical, as vaccines may not 23 always be immediately available to the general population [25]. Modelling how NPIs are utilized 24 can be approached in various ways by mathematical models. For example, Funk et al [19] allow 25 an individual's level of awareness to the presence of a disease shape their usage of self-protective 26 measures. Rizzo et al. model a population where susceptible individuals base their activity rates 27 on the infection prevalence of a disease in the population or the infection incidence over a time step 28 [20]. Similarly, Bagnoli et al. [21] and Del Valle et al. [17] have individuals lower their susceptibility 29 according to the proportion of their contacts in a transmission network that are infectious, and 30 to the infection prevalence, respectively. Poletti et al. [23, 24] incorporate imitation dynamics to 31 model the behavioural changes of the population. Finally, Fenichel et al. [22] and Chen et al. [26] 32 study models where individuals derive utility from engaging in social contact, but raise their risk of 33 infection when doing so. In these aforementioned models, each individual's behaviour is shaped by 34 the information they gather about the disease status of those around them. Thus, in these models. 35 transmission dynamics depend heavily on the perceived risks that drive contact patterns. 36

Further approaches to mathematical models that integrate self protective behaviour into disease 37 transmission utilize adaptive and multiplex networks. An adaptive network is a network whose edges 38 between contacts change dynamically over time. Using these, Gross et al. [27], Shaw and Schwartz 39 [28, 29] and Zanette and Risau-Gusman [30] allow susceptible nodes to rewire their existing connec-40 tions away from infectious nodes at a given rate. The approach of multiplex networks helps 41 to model the many types of social networks individuals may use to acquire information, 42 and Granell et al. [31] and Cozzo et al. [32] use these to study the impact of different 43 information flows on the spread of epidemics. On the other hand, Glass et al. [33] and Kelso 44 et al. [34] use contact networks which include families, schools, and workplaces to study the effects 45 of various NPIs such as school closures and staying at home while infectious. 46

Additionally, vaccines (if available) play a major role in reducing infection rates during an epi-47 demic. Some mathematical models have shown that under voluntary vaccination, populations may 48 not reach sufficient uptake levels to stop an epidemic [35, 36]. However, under voluntary policies 49 in a network, Zhang et al. demonstrate that nodes with high degree can help to suppress disease 50 spread through their increased desires to vaccinate [37]. During an outbreak, complications may 51 arise when there are delays in vaccination. As a result of a delay, epidemic final size can increase 52 significantly [38], especially as the delay lengthens [39]. When considering the efficacy of a vaccine, 53 Wu et al. suggest through their model that a less effective vaccine causes vaccine uptake to increase 54 (to an effectiveness of about 50%), especially for more serious diseases [40]. Insights from the models 55 discussed above, as well as more empirically based research [41], have shown that perceived risks play 56 an important factor in an individual's decision to protect themselves through vaccination. These 57 risks include perceived susceptibility to the illness and perceived risks associated with vaccinating 58 (due to potential side effects) [42, 43]. Much like NPIs, members of a population will base their 59 vaccination decisions on information they are able to gather about the disease during an outbreak. 60 Perceived risks surrounding a disease play a crucial role in vaccination and NPI decisions. Infor-61 mation that shapes these perceptions is gathered by individuals in a population and may be derived 62 from local information [9, 8, 44] (such as social contact networks), or through global information 63 such as media reports about the population as a whole [44, 45]. We note that disease-behaviour 64 models like those discussed above do not typically consider the intervention strategies of vaccination 65 and NPIs simultaneously. However, it is clear that both are important factors in the spread of a 66 disease. Andrews and Bauch [46] have studied the interactions of these two disease interventions 67 with a utility based decision framework model in the context of seasonal influenza. In contrast to our 68 previous work that considers long-term, year-to-year dynamics, here we develop a disease-behaviour 69 70 individual based network simulation model to study interactions between vaccinating behaviour and NPI behaviour and their impact on health outcomes during the course of a single, and sudden, 71 epidemic outbreak of a novel, self limiting infection, where perceived risks and social influence serve 72 as the primary drivers of individual behaviour. Moreover, we include parameters that allow con-73 trolling the relative influence of local versus global information on behaviour. Our main objective is 74 to compare how our model predictions differ from predictions of models that capture behaviour for 75 only one of the two interventions, under various assumptions for (1) transmission probabilities, (2)76 timing of vaccine introduction, and (3) vaccine efficacy, and how efficacy influences vaccine uptake. 77 Furthermore, we explore how the utilization of local versus global information regarding disease 78 spread and vaccine uptake can alter network wide outcomes. 79

80 2. Methods

⁸¹ 2.1. Disease Dynamics

We consider a disease with a susceptible - infectious - recovered - vaccinated (SIRV) natural 82 history. Susceptible individuals may become infected by their infectious neighbours with probability 83 $P(N_{Inf}) = 1 - (1 - \beta)^{N_{Inf}}$ per day, where N_{Inf} is the number of infectious network neighbours, and 84 β is the transmission rate. Infectious individuals move to a recovered (and immune) state for the 85 remainder of the epidemic in a number of days sampled from a Poisson distribution with a mean of 86 7 days. Finally, susceptible individuals may choose to vaccinate and thus become immune for the 87 duration of the epidemic. Baseline parameter values were calibrated to obtain epidemic final size and 88 vaccine uptake trends within the plausible ranges of the corresponding measures in the United States 89 for the 2009 H1N1 influenza pandemic [47, 48], although we emphasize that we are not modelling influenza in particular, but rather we intend our disease represent a hypothetical self-limiting, acute
 infection where individuals only lose natural immunity on a time scale of years. We also assume
 that this is a novel strain of a disease, and individuals have no prior immunity - either
 natural or vaccine conferred. Full details regarding network structure, transmission dynamics,
 and decision modelling appear in the following subsections.

96 2.2. Contact Network

The disease is transmitted on a network consisting of 10,000 nodes which was constructed by 97 sampling from a large contact network derived from empirical contact data in Portland, Oregon 98 [49]. Previous research has shown that the subnetwork is a good approximation to the full network 99 [50]. This network's structure (see Supplementary Information (SI) Figure 1) remains static 100 throughout an epidemic, and we assume that the edges in the network provide sufficient contact 101 between individuals to allow potential disease transmission. We also run simulations testing 102 our primary results on two other types of networks: random networks and power law 103 networks. For details regarding these results, we direct the reader to the SI. 104

¹⁰⁵ 2.3. Non-Pharmaceutical Interventions and Vaccination

Susceptible individuals in the population may engage in self-protective behaviour in response to a growing epidemic. Their self-protecting activity is governed by both the presence of the disease itself [51] and by the social influence of their contacts and the population as a whole [23, 52]. To model this intervention use, we begin by allowing an individual to reduce their susceptibility to $\beta_{NPI} = (e^{-(\Phi + \Gamma_{NPI})})\beta$. Firstly, Φ is an individual's risk perception of the disease, given by

$$\Phi = \sigma f\left(\lambda, \frac{I_{Net}}{k}\right) + (1 - \sigma) f\left(\lambda, \frac{I_{Pop}}{N_{Pop}}\right),\tag{1}$$

where I_{Net} is the number of a given individual's contacts that have been infected, k is the node degree of the individual on the network, I_{Pop} is the number of individuals in the population that have been infected, N_{Pop} is the population size, and σ dictates how members of the population weigh information gathered from their contacts and the population as a whole. Finally, f is a function that determines an individual's response level to increasing infection incidence, given by

$$f(x,y) = 1 - \exp(-xy),$$
 (2)

where x is a proportionality constant that governs the response dynamic (λ in (1)). Since perceived risks only increase in our model (due to the relatively small timespan of one epidemic), we use this functional form. Also, it is an increasing function bounded between 0 and 1 whose shape (or response of increasing perceived risk to incidence) can be governed by a single parameter. Similar functions have been used in the literature surrounding disease spread and self-protective behaviour, for example, see [19]. Secondly, Γ_{NPI_j} measures an individual j's imitation of others who are utilizing self-protective NPI practices, given by

$$\Gamma_{NPI_j} = \sigma f\left(\gamma, \frac{\sum_{i=1}^{k_j^{Vuln}} 1 - \exp(-(\Phi_i + \Gamma_{NPI_i}))}{k_j^{Vuln}}\right) + (1 - \sigma) f\left(\gamma, \frac{\sum_{i\neq j=1}^{N_{Pop}^{Vuln}} 1 - \exp(-(\Phi_i + \Gamma_{NPI_i}))}{N_{Pop}^{Vuln} - 1}\right),$$
(3)

where N_{Pop}^{Vuln} is the number of susceptible or vaccinated (potentially vulnerable) individuals in the population, k_j^{Vuln} is the number of susceptible or vaccinated neighbours individual j has, 124

$$\sum_{i=1}^{k_j^V uin} 1 - \exp(-(\Phi_i + \Gamma_{NPI_i}))$$

$$\frac{1}{k_{Vuln}}$$
is the average amount of transmission rate reduction caused by self-protective

behaviour amongst an individual's susceptible neighbours, $\frac{\sum_{\substack{i \neq j=1}}^{N_{Pop}^{Vuln}} 1 - \exp(-(\Phi_i + \Gamma_{NPI_i}))}{N_{Pop}^{Vuln} - 1}$ is the similar av-126 erage reduction induced by the susceptible population as a whole, and γ is a parameter that governs 127 the response strength of imitation behaviour. Equation (3) captures how individuals reduce their 128 probabilities of becoming infected through observations of others doing the same. This includes imi-129 tation of both network neighbours (σ), and imitation of how the entire population is behaving $(1-\sigma)$. 130 Thus, $0 < \exp(-(\Phi + \Gamma_{NPI})) \leq 1$ dictates how individuals lower their probabilities of becoming 131 infected as they gain awareness of the epidemic by witnessing the disease spread throughout the pop-132 ulation. In our simulations, NPI use for each individual is updated non-synchronously 133 in a new random order at the beginning of each day. Also, transmission rate reduction 134 through NPI use will typically be $\leq 50\%$ for any given individual, which is consistent

135 with the available literature regarding the efficacy of NPIs [53, 54]. 136

If vaccines are available, members of the population may also choose to protect themselves from 137 infection by receiving a vaccine. The decision to vaccinate becomes a more attractive option as 138 vaccine uptake increases [55], and thus an individual's vaccination decision will depend both on 139 their perceived risk of the disease as well as the decisions of others to vaccinate. We represent this 140 as 141

$$\sigma\left(f\left(\lambda, \frac{I_{Net}}{k}\right) + f\left(\gamma, \frac{V_{Net}}{k}\right)\right) + (1 - \sigma)\left(f\left(\lambda, \frac{I_{Pop}}{N_{Pop}}\right) + f\left(\gamma, \frac{V_{Pop}}{N_{Pop}}\right)\right),\tag{4}$$

where V_{Net} and V_{Pop} are the numbers of a given individual's contacts and total number of indi-142 viduals that have been vaccinated, respectively. If we define $\Gamma_V = \sigma f\left(\gamma, \frac{V_{Net}}{k}\right) + (1-\sigma) f\left(\gamma, \frac{V_{Pop}}{N_{Pop}}\right)$, 143 then (4) can simply be written as 144

$$\Phi + \Gamma_V. \tag{5}$$

Equation (5) combines an individual's risk perception of becoming infected, which is based on 145 local and global information of disease incidence, with an individual's imitation of self protective 146 behaviour, which also based on local and global information. 147

If on any day a susceptible individual's preference towards vaccinating, which we set as $\exp(-(\Phi +$ 148 Γ_V), exceeds a given threshold, θ , then that individual will be transferred to the vaccinated compart-149 ment. Otherwise, this is interpreted as an individual being undecided, and they therefore remain 150 susceptible. This process is similar to methods from decision field theory [56], where individuals 151 update their preferences towards making certain decisions based on available information. If their 152 preference toward making an action reaches a pre-defined level, a decision is then subsequently made. 153

3. Results 154

3.1. Baseline Dynamics 155

The baseline scenario of our model (Table 1) simulates an outbreak in a population whose 156 individuals may protect themselves from infection using NPIs or vaccination. We call this the 157

baseline scenario as it was calibrated to achieve plausible epidemic outcomes under the realistic 158 assumption that both vaccination and NPIs are available simultaneously. Henceforth, we will refer 159 to this scenario as the "combined scenario", as both interventions may be used. If both intervention 160 options are available, the final size of the epidemic is lowest compared to when only one of the 161 two interventions are used, as expected (Fig 1a). We also compare the epidemic time series of the 162 combined scenario to hypothetical scenarios where there is no vaccine available over the course of 163 the outbreak ("NPI-only scenario"), or self-protective behaviour is completely ineffective ("vaccine-164 only scenario"). We note that the NPI-only scenario gives similar infection rates as the combined 165 scenario for the first 3 weeks of the epidemic. This occurs because vaccine uptake in the combined 166 scenario is close to zero in the first few weeks, due to low perceived risks of becoming infected 167 while infection prevalence is still minimal, and therefore the differences between scenarios with and 168 without vaccination are small during this period of time. The implication of this is that delays in 169 vaccine availability in the first few weeks of an epidemic may not hinder vaccine uptake under a 170 voluntary vaccination policy. After this initial period, we observe consistently higher cumulative 171 infected for the NPI-only scenario over the combined scenario for the remainder of the epidemic. 172 The NPI-only simulations yield the greatest average cumulative infection incidence, as the response 173 from solely NPIs amongst susceptible individuals cannot match the disease mitigation of a perfectly 174 efficacious vaccine, in the long term (however, we note that the difference in cumulative incidence 175 is relatively small). In the vaccine-only scenario, infection incidence spikes rapidly but the epidemic 176 lasts a shorter amount of time than in the NPI-only scenario. The relatively rapid early spike in 177 total infected individuals is due to the lack of vaccine uptake in the first weeks of the epidemic, as 178 vaccination decisions are not activated until the perceived threat of becoming infected is sufficiently 179 high. In all these cases, self-protective behaviour serves to slow the spread of an epidemic, but 180 does not successfully reduce the final attack rate as significantly compared to when it is aided by 181 vaccination. 182

In the NPI-only scenario, NPI uptake amongst susceptible individuals is much more pronounced 183 than when vaccination is also an option (Fig 1b). This occurs for two reasons. Firstly, if vaccination 184 can occur, those that practice the strongest self protective behaviour due to having high levels 185 of perceived risk will be amongst the first to vaccinate. In turn, this will lower the average NPI 186 uptake amongst the remaining susceptible population. Secondly, if members of the population 187 are vaccinating, the spread of the disease will be suppressed causing perceived risks of becoming 188 infected to be lower. Thus, resulting NPI use will be less prominent. In the absence of vaccination, 189 transmission reduction through NPI use simply continues to rise along with the infection incidence 190 seen in Figure 1a. In the final vaccine-only scenario, total vaccine uptake is increased on average 191 192 (Fig 1c). Moreover, vaccine coverage begins to rise earlier in response to the rapid spike in infection incidence that is observed when no transmission reduction is present through NPIs. Thus, when 193 NPI effects are not considered, predicted vaccine uptake is significantly higher. 194

195 3.2. Transmission Rate

Time series of infection prevalence corresponding to different transmission rates can help us understand epidemic spread in our 3 scenarios (Fig 2). When the transmission rate is low, NPIs alone are relatively effective at hindering the growth of the epidemic, lowering the peak infection prevalence compared to the vaccine-only case (Fig 2a). For a transmission rate of $\beta = 0.00493$ per infectious contact per day, simulations that utilize NPIs only or vaccination only result in the same epidemic final size (Fig 2b). Although the peak infection prevalence in this scenario is larger for vaccine-only simulations, the epidemic dies out more quickly compared to the NPI-only scenario, resulting in

the same cumulative infection incidence over the epidemic duration. For higher transmission rates. 203 the vaccine only scenario outperforms the NPI only scenario (Fig 2c). Although NPIs delay the 204 peak of the epidemic, infection prevalence dies out more slowly than when the population uses solely 205 vaccination instead. However, this highlights the importance of NPIs in epidemics where vaccination 206 may not be immediately available. Considering the combined scenario data in Fig 2c, which indicates 207 infection prevalence in simulations utilizing both NPIs and vaccination, the initial disease spread is 208 very similar to that of the NPI-only scenario. Only when individuals begin to vaccinate does the 209 infection prevalence in the combined scenario show quantitative difference to the infection prevalence 210 in the NPI-only scenario. Thus, as long as a vaccine is made available within a given time frame, 211 the final size can be expected to be the same due to the early activation of NPIs. 212

Vaccine timing plays a critical role in the health outcomes of the population during an epidemic, 213 across a range transmission rates (Fig 3). In the combined scenario (Fig 3a), a vaccine can be 214 introduced up to 20 days after the start of an epidemic for the epidemic final size to be roughly the 215 same as the scenario when a vaccine is available from day one, for baseline transmission rates. If 216 we disregard the use of NPIs (Fig 3b), the vaccine must be made available within 15 days before 217 we begin to observe larger epidemic final sizes. This effect is similar for $\beta = 0.006$ per infectious 218 contact per day. In the combined scenario, a vaccine must be made available within 15 days before 219 epidemic sizes increase. However, in the vaccine-only case, vaccine availability must occur within 220 just 10 days. Finally, for lower disease transmission, vaccine introduction timing has little impact 221 on infection incidence in the combined scenario. However, in the vaccine-only scenario, we see final 222 sizes begin to increase when availability occurs after 20 days. From these results, we also notice that 223 the rate of increase of epidemic final sizes corresponding to the timing of vaccine introduction are 224 much greater. For example, given the baseline transmission rate, the difference in infection incidence 225 between immediate vaccine availability and availability beginning on day 60 is $\approx 20\%$ in the vaccine-226 only case. However, the same measure in the combined case is only $\approx 4\%$. Thus, the prediction 227 from these two modelling approaches of epidemic size induced by vaccine timing introduction differs 228 by about 16% of the entire population size. 229

Finally, we also consider measures for epidemic final size (Fig 4). When the transmission rate 230 is low, the final size is the same for the combined scenario as for the NPI-only scenario, but much 231 higher for the vaccine-only scenario (Fig 4a). Hence, for low transmission rates, NPIs on their own 232 can reduce final size as much as combined use of NPIs and vaccines, although the same is not true 233 for vaccines on their own. This is due to individuals promptly adopting NPIs, which are targeted at 234 the leading edge of the epidemic and quick to implement, curbing disease spread immediately. Also, 235 vaccine uptake is much larger in the vaccine-only scenario than in the combined scenario (Fig 4b). 236 In contrast, when the transmission rate is high, the final size is almost (but not quite) the same for 237 the combined scenario as for the vaccine-only scenario, but much higher for the NPI-only scenario. 238 Moreover, vaccine uptake is also nearly the same for both of the scenarios that include vaccination. 239 Hence, for high transmission rates, vaccines on their own can reduce final size almost as much as 240 combined use of NPIs and vaccines, although the same is not true for NPIs on their own. In the 241 case of NPIs, the NPI uptake amongst susceptible individuals does not change for $\beta > 0.0045$, due 242 to the adoption of vaccination (Fig 4c). However, in the NPI-only scenario, susceptibility reduction 243 through NPIs continues to rise along with the transmission rate. 244

In summary, when transmission rates are sufficiently low, NPIs alone can be almost as effective as having both vaccines and NPIs (whereas vaccination alone is relatively less effective), but when transmission rates are sufficiently high, vaccines alone can be almost as effective as having both interventions (whereas NPIs alone are relatively less effective).

249 3.3. Vaccine Efficacy

Vaccines are never 100% efficacious. For less effective vaccines, we can expect infection incidence and vaccination coverage to change as individuals in the population adapt to the quality of interventions available to them. Thus, we explore the dynamics under various vaccine efficacies (denoted ϵ), and how they relate to vaccine coverage and epidemic final size with and without the additional impacts of NPIs (Fig 5). We note that in our simulations, vaccines give full protection with probability ϵ , and no additional protection with probability $1 - \epsilon$.

As vaccine efficacy decreases, the vaccine-only scenario overestimates the amount of vaccine 256 uptake demanded by up to 16.5% of the population size relative to the combined scenario. We 257 also observe that as vaccine efficacy decreases, the subsequent increase in vaccine coverage of the 258 population is larger when NPI effects are not incorporated. For example, between efficacies of 100% 259 to 50%, the combined scenario of the model predicts $\sim 3.5\%$ more of the population vaccinating, 260 whereas with the vaccine-only scenario, simulations predict an $\sim 8\%$ increase. This effect is also 261 seen with epidemic final size (Fig 5a). Across all efficacies, final size increase is only $\sim 1.5\%$ of the 262 entire population size with combined NPI and vaccine utilization, and $\sim 5.5\%$ with only vaccination. 263 Thus, we see that disregarding the impact of NPIs may lead to an overestimation of the population's 264 vaccine demand and final epidemic size. Moreover, the increases in vaccine uptake and final size with 265 decreasing vaccine efficacy may be less significant than what previous predictions which disregard 266 NPI effects show [40]. Finally, when incorporating vaccination decisions and self protective behaviour 267 simultaneously into the model, we observe that predicted levels of vaccine uptake are much smaller 268 than when no NPIs are implemented (Fig 5b). 269

270 3.4. Pairwise Correlations

As an epidemic unfolds across a network, the status of the nodes will develop while the disease spreads and intervention decisions are made. As a result, the spatial structure of infected and susceptible individuals on the networks will evolve over time as well. The correlation between these pairs can offer insight on the vulnerability of the network to disease spread and how individuals react to infection prevalence according to the information available to them, which we control with the parameter σ . To measure the correlation between node pairs, we follow Keeling [57]:

$$C_{AB} = \frac{N_{Pop}}{k_{avg}} \frac{[AB]}{[A][B]},\tag{6}$$

where k_{avg} is the average node degree on the network. With this formulation, an increase in C_{AB} indicates an increase in correlation as the number of [AB] pairs in the network relative to the number of type [A] nodes and type [B] nodes also increases. A value of $C_{AB} = 1$ indicates no correlation [57].

Considering the correlation between susceptible-infected ([SI]) pairs (Fig 6c), we observe a rapid 281 initial spike in the network. This early increase is due to the first infected individuals spreading 282 the disease to their network contacts, enabling more opportunities for transmission. As infection 283 prevalence begins to peak (Fig 6a), infected individuals have a higher probability of being connected 284 to a non-susceptible node, which results in the decline of C_{SI} in the network, as distinct clusters of 285 infected and other infected, recovered or vaccinated individuals develop. However, the correlation 286 rises again as infectious nodes recover and only a final few clusters of infected and susceptible nodes 287 remain, more so for lower σ as those who vaccinated are less likely to be connected to an infectious 288 node. Correlations of vaccinated nodes with nodes that are or have been infected, [VI] and [VR], 289

respectively, also show how network dynamics respond to different levels of σ (Fig 6d,e). When 290 individuals base their decisions on local information, that is, on the basis of the number of infectious 291 neighbours ($\sigma = 1.0$), C_{VI} and C_{VR} are higher. This indicates successful ring vaccination occurring 292 in proximity to the infectious individuals. Under strong influence of local information, neighbours 293 of infectious individuals develop a high perceived risk and decide to vaccinate earlier. Then, so-294 cial influences reinforce this vaccinating behaviour, resulting in clusters of vaccinated individuals 295 around infectious individuals. However, when decisions are made more strongly based on popula-296 tion level infection prevalence ($\sigma = 0.5$), [VI] and [VR] pairs become less common in proportion to 297 all vaccinated and infected/recovered nodes since vaccinations do not always occur on the epidemic 298 front. The cumulative vaccine uptake under global information is higher than under local informa-299 tion, however, vaccine uptake increases more rapidly in the early stages of the epidemic under local 300 information (Fig 6b). This reflects the efficiency of targeted vaccination under local information, 301 where vaccines are administered to the contacts of infectious individuals so that infection spread is 302 efficiently prevented. Finally, as the epidemic dies out and infectious nodes become rare, [VI] and 303 [VR] correlations across varying levels of σ converge to similar values. 304

As σ decreases in our model, [SS] pairs become more common relative to the total number of susceptible nodes towards the end of an epidemic, increasing C_{SS} (Fig 6f). On the other hand, with higher σ , final values of C_{SS} continue to decrease. However, we note that during an epidemic, the opposite is true, albeit to a lesser extent. When $\sigma = 0.5$, vaccination occurs in locations other than the epidemic front, in turn decreasing C_{SS} compared to higher values of σ . Nonetheless, the ring vaccination observed with increased σ is more efficient than the more random vaccine allocation seen when $\sigma = 0.5$, for example, due to the disease only being able to spread along the network edges.

312 4. Discussion

We have developed and analyzed a model that simulates a population's adaptive self protective behaviour (use of NPIs and vaccination) in the face of a disease outbreak, in contrast to most previous approaches that model only vaccinating behaviour or only NPI behaviour. We allow an individual's actions to depend both on their perceived risk of infection developed from their experiences with the disease on the network (both from their network neighbours and from the population as a whole), as well as imitation of the behaviour of others in the population.

Surprisingly, when transmission rates are low, the NPI-only scenario offers comparable disease 319 mitigation effectiveness to the combined scenario, while the vaccine-only scenario results in relatively 320 larger epidemic sizes than either the NPI-only scenario or the combined scenario. For higher trans-321 mission rates, the opposite becomes true. That is, the vaccine-only scenario is almost as effective as 322 the combined scenario for reducing infection incidence, but the NPI-only scenario fares worse. If a 323 vaccine is not available immediately to the population at the start of an epidemic, epidemic mitiga-324 tion through adaptive NPI behaviour can curb the growth of an epidemic. Thus, if vaccination is 325 made available to the population within a given time frame, health outcomes will be very similar to 326 situations where a vaccine was always available. If, however, the effects of NPIs are not incorporated, 327 then these time frames are comparatively shorter. Moreover, the increases in infection incidence for 328 the vaccine-only scenarios are significantly higher the later the vaccine is introduced, resulting in in-329 creasingly higher predictions of epidemic final size. Finally, the impact of varying vaccine efficacy on 330 both vaccine uptake and epidemic final size varies significantly between scenarios with and without 331 adaptive NPI behaviour. The increases in both final size and vaccine uptake when vaccine efficacy 332 is decreased are much higher for the vaccine-only scenario than the combined scenario. Hence, a 333

model of adaptive vaccinating behaviour that does not also account for adaptive NPI behaviour
 will make very different predictions than a model that accounts for adaptive behaviour toward both
 interventions. This again highlights the positive benefits of epidemic mitigation through adaptive
 NPI behaviour.

From a network perspective, individuals basing their decisions to practice NPIs or become vac-338 cinated based on the infection prevalence and behaviour in their infection contact network leads to 339 the most effective disease control. Pairwise correlations between vaccinated and infected nodes are 340 highest when this information gathering is possible, as those that vaccinate are typically connected 341 to infected nodes. We also tested the main results with two additional types of networks: 342 random networks and power law networks (see SI). While the dynamics are qualita-343 tively the same, the amount of change in epidemic final size or vaccine uptake with 344 differing vaccine delays or vaccine efficacies can depend on the specific network type. 345 Assumptions about network structure and transmission are an important consideration 346 - particularly when modelling a specific disease. For example, a transmission network 347 for influenza likely has a different structure than one that would be used to model HIV 348 transmission. 349

In the combined scenario, epidemic final size is suppressed most effectively compared to when only single interventions are possible. Also, when the effects of NPIs are not considered in our vaccineonly scenario (an assumption which is common in previous behaviour-disease models focusing on vaccinating behaviour), vaccine uptake predictions are higher compared to when these effects are considered by our model, on account of counteractive feedbacks from NPI behaviour.

Our model includes some simplifying assumptions about behaviour-disease dynamics. For exam-355 ple, NPI efficacy is poorly quantified in the epidemiological literature, and it is not always known in 356 357 what situations individuals may practice them most often. Thus, we assume that NPIs for the spreading disease are not used initially, but in reality there may be some baseline level 358 of NPIs used in the population due to other circulating diseases. Moreover, we do not 359 model the effects of NPI practices that infectious individuals may utilize, such as self 360 isolation. Instead, we make the assumption that infectious NPI use is absorbed into the 361 transmission rate. Also, the network we used in our simulations could be extended to distinguish 362 family, friend, and work structures, where transmission rates to an individual can vary depending 363 on what category certain network contacts fall in. Similarly, age structure can be introduced into 364 the model. As children will be much less likely to effectively practice NPIs, disease transmission 365 in these groups may be more rapid than our model predicts. Finally, we did not include the 366 impact of asymptomatic infections, and assumed all cases were identifiable in our main 367 results. However, we also considered a scenario where 50% of cases were asymptomatic 368 (see SI), and the primary results regarding vaccine efficacy and vaccine availability de-369 lays across various transmission rates are qualitatively the same. Although the main 370 results are similar, in future work that aims to model a specific disease, accounting for 371 asymptomatic infections is an important factor. 372

Through these experiments, we see that predictions of health outcomes and vaccine uptake in a population can vary significantly when NPI use is, or is not, considered. It is important for behaviourally based epidemiological models to incorporate the effects of transmission reduction through this adaptive behaviour, as perceived risks of a disease will in turn be shaped by them subsequently altering the outcomes of an epidemic. The same is also true of models that focus on modelling NPI behaviour, in populations where adaptive vaccinating behaviour could significantly alter model predictions of NPI practices.

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Parameter	Table 1: Baseline Parameter Values. Description	Value
λ	Constant governing awareness/risk perception of disease	1.5
γ	Constant governing behaviour imitation	0.5
heta	Vaccinating threshold	0.35
σ	Weighting for global versus local information	0.8
β	Transmission rate	0.005
N_{Pop}	Population size	10,000
I_0	Initial number of infectious persons	20
η	Mean infectious period, in days	7

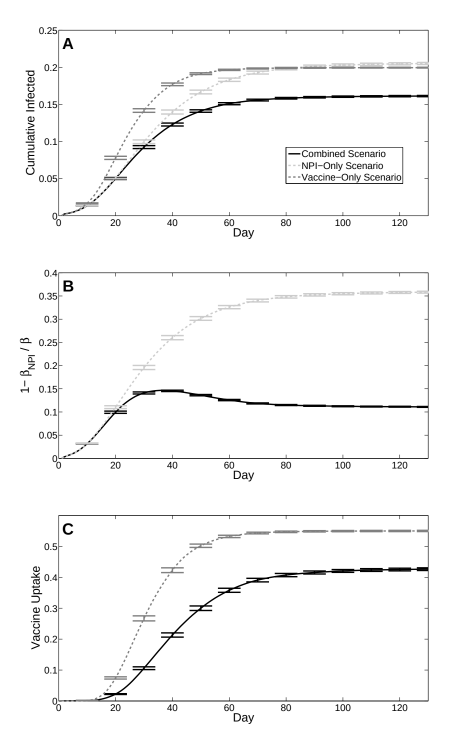


Figure 1: Time series of an epidemic, 95% confidence intervals shown every 10 days around the mean of 500 realizations. a) Cumulative infection incidence b) Transmission rate reduction due to self-protective behaviour (NPIs) amongst the susceptible population, c) Cumulative vaccine coverage.

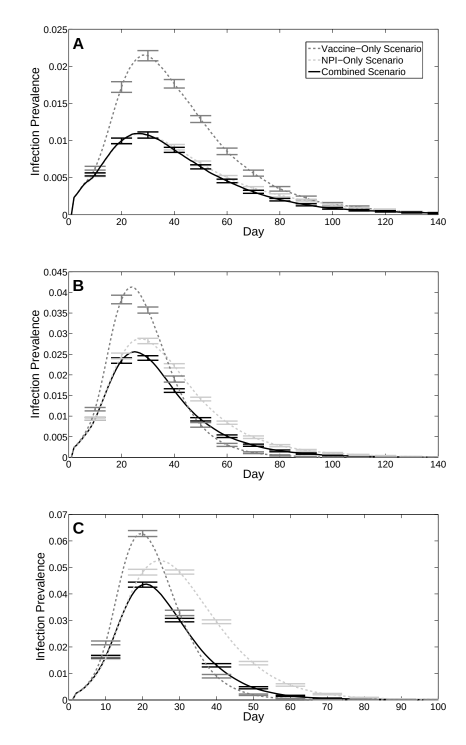


Figure 2: Time series of infection prevalence with the vaccine-only scenario, the NPI-only scenario, and the combined scenario. 95% confidence intervals shown every 10 days around the mean of 500 realizations. (a) $\beta = 0.004$ (b) $\beta = 0.0043$ (c) $\beta = 0.006$.

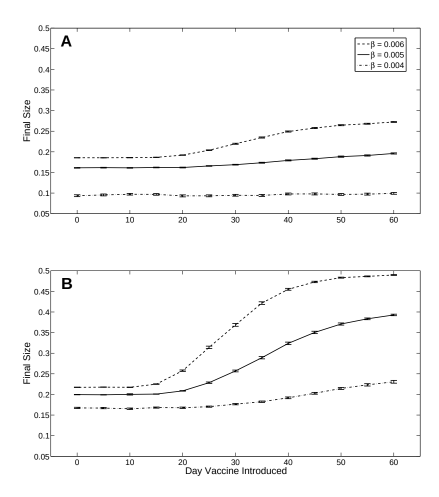


Figure 3: Epidemic final sizes with respect to when vaccination is made available. (a) With NPIs. (b) Without NPIs

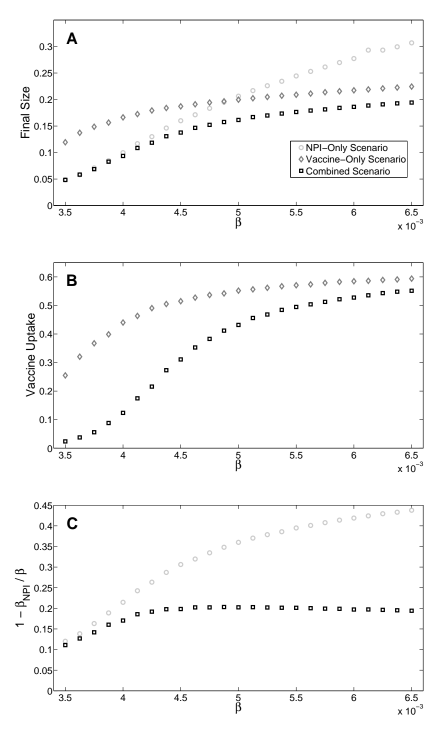


Figure 4: Epidemic measures with respect to transmission rate. (a) Epidemic final size. (b) Vaccine uptake. (c) Transmission rate reduction amongst susceptible individuals.

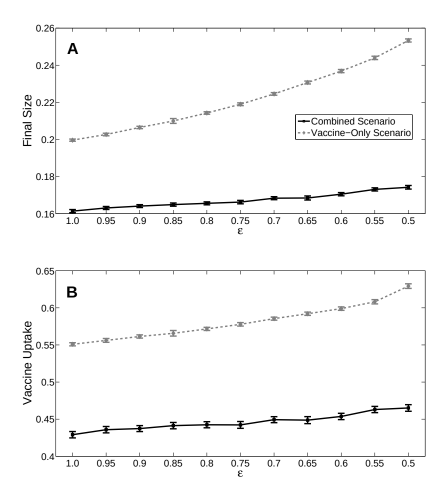


Figure 5: Effects of vaccine efficacy between scenarios with and without NPIs on (a) Vaccine uptake, and (b) Final epidemic size.

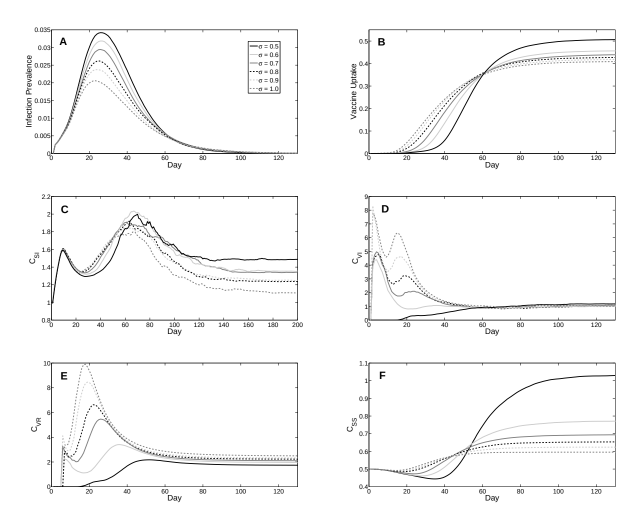


Figure 6: Time series of epidemics over different values of σ , the weighting for global versus local information . (a) Infection prevalence, (b) Vaccine uptake, (c) Correlation between SS pairs, (d) SI pairs, (e) VI pairs, and (f) VR pairs. Lines show the average values over 500 realizations.

383 References

- P. Y. Lee, D. B. Matchar, D. A. Clements, J. Huber, J. D. Hamilton, E. D. Peterson, Economic
 analysis of influenza vaccination and antiviral treatment for healthy working adults, Annals of
 Internal Medicine 137 (4) (2002) 225–231.
- [2] H. Pearson, T. Clarke, A. Abbott, J. Knight, D. Cyranoski, SARS: What have we learned?,
 Nature 424 (2003) 121–126.
- [3] A. Balkhair, K. A. Maamari, F. B. Alawi, The struggle against MERS-CoV (the novel coronavirus), Oman Medical Journal 28 (4) (2013) 226–227.
- [4] WHO Ebola Response Team, Ebola virus disease in west africa the first 9 months of the
 epidemic and forward projections, The New England Journal of Medicine 371 (2014) 1481–
 1495.
- [5] M. P. Girard, J. S. Tam, O. M. Assossou, M. P. Kieny, The 2009 A(H1N1) influenza virus pandemic: A review, Vaccine 28 (2010) 4895–4902.
- [6] S. Funk, M. Salathé, V. A. A. Jansen, Modelling the influence of human behaviour on the
 spread of infectious diseases: A review, Journal of the Royal Society Interface 7 (50) (2010)
 1247–1256.
- J. T. Lau, X. Yang, E. Pang, et al., SARS-related perceptions in hong kong, Emerging Infectious
 Diseases 11 (3).
- [8] T. Philipson, Private vaccination and public health: An empirical examination for U.S. measles,
 The Journal of Human Resources 31 (3).
- [9] A. Ahituv, V. J. Hotz, T. Philipson, The responsiveness of the demand for condoms to the local
 prevalence of AIDS, The Journal of Human Resources 31 (4).
- [10] D. P. Durham, E. A. Casman, Incorporating individual health-protectice decisions into disease
 transmission models: A mathematical framework, Journal of the Royal Society Interface 9 (68)
 (2012) 562–570.
- [11] O. de Zwart, I. Veldhuijzen, G. Elam, Perceived threat, risk perception, and efficacy beliefs
 related to SARS and other (emerging) infectious diseases: results of an international survey,
 International Journal of Behavioral Medicine 16 (2009) 30–40.
- [12] D. Koh, K. Takahashi, M.-K. Lim, et al., SARS risk perception and preventive measures,
 Singapore and Japan, Emerging Infectious Diseases 11 (4) (2005) 641–642.
- [13] Centers for Disease Control and Prevention, Nonpharmaceutical Interventions (NPIs), http:
 //www.cdc.gov/nonpharmaceutical-interventions/ (August 2012).
- [14] G. B. Chapman, E. J. Coups, Predictors of influenza vaccine acceptance among healthy adults,
 Preventive Medicine 29 (1999) 249–262.
- ⁴¹⁷ [15] C. Bauch, P. Manfredi, A. d'Onofrio, Behavioral epidemiology of infectious diseases: An
 ⁴¹⁸ overview, in: P. Manfredi, A. d'Onofrio (Eds.), Modeling the Interplay Between Human Behav⁴¹⁹ ior and the Spread of Infectious Diseases, Springer, 2013, Ch. 1, pp. 1–19.

- [16] Z. Wang, M. A. Andrews, Z.-X. Wu, L. Wang, C. T. Bauch, Coupled disease-behavior dynamics
 on complex networks: A review, Physics of Life Reviews.
- [17] S. Del Valle, H. Hethcote, J. Hyman, C. Castillo-Chavez, Effects of behavioral changes in a
 smallpox attack model, Mathematical Biosciences 195 (2005) 228–251.
- [18] T. C. Reluga, Game theory of social distancing in response to an epidemic, PLOS Computational
 Biology 6 (5).
- [19] S. Funk, E. Gilad, C. Watkins, V. A. A. Jansen, The spread of awareness and its impact on
 epidemic outbreaks, Proceedings of the National Academy of Sciences of the United States of
 America 106 (16) (2008) 6872–6877.
- [20] A. Rizzo, M. Frasca, M. Porfiri, Effect of individual behavior on epidemic spreading in activitydriven networks, Physical Review E 90.
- ⁴³¹ [21] F. Bagnoli, P. Liò, L. Sguanci, Risk perception in epidemic modeling, Physical Review E 76.
- E. P. Fenichel, C. Castillo-Chavez, M. G. Ceddia, G. Chowell, P. A. G. Parra, et al., Adaptive human behavior in epidemiological models, Proceedings of the National Academy of Sciences of the United States of America 108 (15) (2011) 6306–6311.
- [23] P. Poletti, B. Caprile, M. Ajelli, A. Pugliese, S. Merler, Spontaneous behavioural changes in
 response to epidemics, Journal of Theoretical Biology 260 (2009) 31–40.
- ⁴³⁷ [24] P. Poletti, M. Ajelli, S. Merler, Risk perception and effectiveness of uncoordinated behavioral
 responses in an emerging epidemic, Mathematical Biosciences 238 (2012) 80–89.
- ⁴³⁹ [25] E. Check, Avian flu special: Is this our best shot?, Nature 435 (2005) 404–406.
- ⁴⁴⁰ [26] F. Chen, M. Jiang, S. Rabidoux, S. Robinson, Public avoidance and epidemics: Insights from ⁴⁴¹ an economic model, Journal of Theoretical Biology 278 (2011) 107–119.
- [27] T. Gross, C. J. D. D'Lima, B. Blasius, Epidemic dynamics on an adaptive network, Physical Review Letters 96.
- [28] L. B. Shaw, I. B. Schwartz, Fluctuating epidemics on adaptive networks, Physical Review E 77.
- [29] L. B. Shaw, I. B. Schwartz, Enhanced vaccine control of epidemics in adaptive networks, Physical
 Review E 81.
- [30] D. H. Zanette, S. Risau-Gusman, Infection spreading in a population with evolving contacts,
 Journal of Biological Physics 34 (2008) 135–148.
- [31] C. Granell, S. Gómez, A. Arenas, Dynamical interplay between awareness and epidemic spread ing in multiplex networks, Physical Review Letters 111.
- [32] E. Cozzo, R. A. B. nos, S. Meloni, Y. Moreno, Contact-based social contagion in multiplex
 networks, Physical Review E 88.
- [33] R. J. Glass, L. M. Glass, W. E. Beyeler, H. Min, Targeted social distancing design for pandemic
 influenza, Emerging Infectious Diseases 12 (11) (2006) 1671–1681.

- ⁴⁵⁵ [34] J. K. Kelso, G. J. Milne, H. Kelly, Simulation suggests that rapid activation of social distancing ⁴⁵⁶ can arrest epidemic development due to a novel strain of influenza, BMC Public Health 9 (117).
- [35] R. Vardavas, R. Breban, S. Blower, Can influenza epidemics be prevented by voluntary vacci nation?, PLOS Computational Biology 3 (5).
- [36] C. T. Bauch, A. P. Galvani, D. J. Earn, Group interest versus self-interest in smallpox vaccina tion policy, Proceedings of the National Academy of Sciences of the United States of America
 100 (18) (2003) 10564–10567.
- [37] H. Zhang, J. Zhang, C. Zhou, M. Small, B. Wang, Hub nodes inhibit the outbreak of epidemic
 under voluntary vaccination, New Journal of Physics 12.
- ⁴⁶⁴ [38] Y. Yang, J. D. Sugimoto, M. E. Halloran, N. E. Basta, D. L. Chao, et al., The transmissibility ⁴⁶⁵ and control of pandemic influenza A (H1N1) virus, Science 326 (2009) 729–733.
- [39] M. Z. Gojovic, B. Sander, D. Fisman, M. D. Krahn, C. T. Bauch, Modelling mitigation strategies
 for pandemic (H1N1) 2009, PMC Canadian Medical Association Journal 181 (2009) 673–680.
- ⁴⁶⁸ [40] B. Wu, F. Fu, L. Wang, Imperfect vaccine aggravates the long-standing dilemma of voluntary ⁴⁶⁹ vaccination, PLOS One 6.
- [41] N. T. Brewer, G. B. Chapman, F. X. Gibbons, et al., Meta-analysis of the relationship between
 risk perception and health behaviour: The example of vaccination, Health Psychology 26 (2)
 (2007) 136–145.
- [42] R. Roberts, Q. Sandifer, M. Evans, M. Noland-Farrell, P. Davis, Reasons for non-uptake of
 measles, mumps, and rubella catch up immunisation in a measles epidemic and side effects of
 the vaccine, BMJ 310 (1995) 1629–1639.
- [43] P. H. Streefland, Public doubts about vaccination safety and resistance against vaccination,
 Health Policy 5 (2001) 159–172.
- [44] E. Klein, D. L. Smith, C. A. Gilligan, Economic incentives and mathematical models of disease,
 Environment and Development Economics 12 (2007) 707–732.
- ⁴⁸⁰ [45] T. R. Berry, J. Wharf-Higgins, P. Naylor, SARS wars: An examination of the quantity and ⁴⁸¹ construction of health information in the news media, Health Communication 21.
- [46] M. A. Andrews, C. T. Bauch, Disease interventions can interfere with one another through
 disease-behaviour interactions, PLOS Computational Biology 11.
- [47] S. Shrestha, D. Swerdlow, R. Borse, V. Prabhu, L. Finelli, et al., Estimating the burden of 2009
 pandemic influenza A (H1N1) in the United States (April 2009-April 2010), Clinical Infectious
 Diseases 52 (2011) S75–S82.
- [48] Centers for Disease Control and Prevention, Flu Vaccination Coverage, United States, http:
 //www.cdc.gov/flu/fluvaxview/coverage-1213estimates.htm (April 2015).
- [49] Network Dynamics and Simulation Science Laboratory, Synthetic data products for societal
 infrastructures and proto-populations: Data set 1.0, Tech. rep., Virginia Polytechnic Institute
 and State University (2008).

- ⁴⁹² [50] C. R. Wells, C. T. Bauch, The impact of personal experiences with infection and vaccination
 ⁴⁹³ on behaviour-incidence dynamics of seasonal influenza, Epidemics 4 (2012) 139–151.
- ⁴⁹⁴ [51] M. Z. Sadique, W. J. Edmunds, R. D. Smith, W. J. Meerding, O. de Zwart, et al., Precautionary
 ⁴⁹⁵ behavior in response to perceived threat of pandemic influenza, Emerging Infectious Diseases
 ⁴⁹⁶ 13 (9) (2007) 1307–1313.
- ⁴⁹⁷ [52] C. T. Bauch, Imitation dynamics predict vaccinating behaviour, Proceedings of the Royal So-⁴⁹⁸ ciety B Biological Sciences 272 (1573) (2005) 1669–1675.
- E. L. Larson, Y. hui Ferng, J. Wong-McLoughlin, S. Wang, M. Haber, S. S. Morse, Impact of
 non-pharmaceutical interventions on uris and influenza in crowded, urban households, Public
 Health Reports 125 (2) (2010) 178–191.
- J. J. Sheehan, P. J. Mott, B. W. Sisk, J. W. Arbogast, C. Ferrazzano-Yaussy, C. A. Bondi,
 Alcohol-based instant hand sanitizer use in military settings a prospective cohort study of
 army basic trainees, Military Medicine 172 (11) (2007) 1170–1176.
- ⁵⁰⁵ [55] S. Bhattacharyya, C. T. Bauch, "wait and see" vaccinating behaviour during a pandemic: A ⁵⁰⁶ game theoretic analysis, Vaccine 29 (2011) 5519–5525.
- ⁵⁰⁷ [56] J. R. Busemeyer, J. T. Townsend, Decision field theory: A dynamic-cognitive approach to decision making in an uncertain environment, Psychological Review 100 (3) (1993) 432–459.
- [57] M. J. Keeling, K. T. Eames, Networks and epidemic models, Journal of the Royal Society: Interface 2 (2005) 295–307.