Transmission Heterogeneity and Control Strategies for Infectious Disease Emergence

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Background. The control of emergence and spread of infectious diseases depends critically on the details of the genetic makeup of pathogens and hosts, their immunological, behavioral and ecological traits, and the pattern of temporal and spatial contacts among the age/stage-classes of susceptible and infectious host individuals. Methods and Findings. We show that failing to acknowledge the existence of heterogeneities in the transmission rate among age/stage-classes can make traditional eradication and control strategies ineffective, and in some cases, policies aimed at controlling pathogen emergence can even increase disease incidence in the host. When control strategies target for reduction in numbers those subsets of the population that effectively limit the production of new susceptible individuals, then control can produce a flush of new susceptibles entering the population. The availability of a new cohort of susceptibles may actually increase disease incidence. We illustrate these general points using Classical Swine Fever as a reference disease. Conclusion. Negative effects of culling are robust to alternative formulations of epidemiological processes and underline the importance of better assessing transmission structure in the design of wildlife disease control strategies.

INTRODUCTION

Historically, models for infectious diseases considered populations of host and pathogen to be well-mixed with homogeneous disease transmission among susceptible and infected individuals. In recent years, however, we have recognized that transmission may not be constant but may vary with time, social structure, and/or age/stage-class. In human epidemiology it is well known that transmission may indeed change substantially among social groups and age classes, as observed, for instance, in the early phases of HIV epidemics [1,2]. Recent research has also emphasized the importance of super-spreaders in the dynamics of SARS, sexually transmitted and childhood diseases [3,4]. However, there is only anecdotal information about variation in transmission rate by age/stage structure in wild animals or the implication of such variation on disease dynamics in zoonotic reservoirs [5] and only limited information in the case of domestic animals (see e.g., [6]). This is unfortunate since the majority of emerging infectious diseases are zoonotic and it is well know that many species exhibit a high degree of heterogeneity in their spatial, social and age/stage structure. It is highly likely that age/stage-dependent behavioral differences in reservoir contact rates may significantly affect disease transmission between and within different age/stage classes and, as a consequence, the effectiveness of disease control policies.

While the science of infectious diseases has made tremendous progress in the last several decades thanks, in part, to advances in molecular biology, immunology, medicine and mathematical modeling, the eradication of pathogens and parasites in wildlife relies very often only on two simple strategies, namely vaccination and culling, i.e., the removal of animals to push host population density below the threshold for disease invasion [7]. Quarantine and isolation through the construction of sanitary containments are rare options in the control of wildlife diseases and are applied primarily to domestic animals and farms such in the case of the foot-and-mouth epidemics in UK [8] and of avian flu epidemics in Asia [9] often at the cost of huge economic losses.

Oral vaccination, on the contrary, is a quite widespread technique for disease eradication and control: it has been applied in the USA to control the westward expansion of rabies virus in raccoon hosts and has effectively eliminated raccoons in Texas [10,11]. In Germany, oral vaccination is used to control classical swine fever virus exposure in wild boars [12]. The drawbacks of oral vaccination mainly consist in the difficulty of producing, at a reasonable price, a sufficient amount of a vaccine able to persist long enough in baits so as to be picked up by a suitable fraction of susceptible animals. Moreover, vaccination might be less effective than expected, as older animals–who may have antibodies resulting from prior infection–are often more aggressive in ingesting baits than younger more susceptible individuals.

Culling is usually the most simple and economical measure to control diseases spread in wildlife and its application is strongly supported as an emergency procedure for disease eradication. It has been historically applied to control different domestic and wildlife diseases with the aim of reducing host population or removing infected individuals, such as for bovine tuberculosis in badgers [13] and foot-and-mouth disease in cattle in the UK [14], avian flu in waterfowl birds and poultry in Asia [9] and classical swine fever in wild boars and domestic pigs in Europe [15]. Albeit, sport hunting per se is certainly not aimed at preventing disease spread, it usually exerts effects similar to those of culling in terms of population density reduction. The same is true also for illegal hunting (poaching).

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Despite its simplicity and alleged cost effectiveness, there is some evidence that under a variety of circumstances culling may not generate the benefits anticipated [16–18]. Selective hunting may interfere with establishment of herd immunity inducing faster turnover of the population and decreasing host life expectancy. Furthermore, it may induce long distance host movement, increasing contacts between different groups of animals [19], and it may affect the evolution of some host species traits [20]. Finally, culling often affects age/stage structure by preferentially removing older and less susceptible individuals. This is important, as the basic theory for Susceptible-Infected-Recovery (SIR-like) compartmental models of a self-regulating host suggests that disease prevalence in a homogenous population should monotonically decrease with increasing culling rate [as in 21,22 models]. However, as we will see, the existence of age/stage structure in transmission rate may significantly alter this conclusion.

In the present work we show (using very general assumptions about life history traits of host species) that the presence of age-dependent heterogeneity in the transmission rate may produce the counter-intuitive result that disease prevalence increases over a range of intermediate levels of culling. We recast a simple SIR model for wildlife disease with two age/stage classes, namely, pre-reproductive juveniles and adults and sub-adults (see Material and Methods). Numerical characterization of the problem was examined for the specific case of classical swine fever in wild boar (see Protocol S1 in Supporting Information). We chose CSF as reference disease because it has caused serious economic losses in Europe from spillover infection to pig farms over the last twenty years [23] and it is still endemic in Asia and South America and in some regions of Eastern Europe and Italy (Sardinia). Also, Choisy and Rohani [24] have recently illustrated similar points using CSF but with a different model structure. Comparison across model structures for the same disease will lead to an increased robustness in any general characterization of optimal strategies for control.

While the ideas being investigated in the present work are based on a detailed understanding of wild boar biology, the epidemiological model itself has been simplified to address the following key question: how do age-dependent heterogeneities in transmission interact with culling rate in the control of disease prevalence?

To answer this question, we extend the classical SIR model of infection to allow for an age structured wildlife population. In this simple model, host population is divided into two age classes, juveniles and adults. We assumed that juveniles are highly susceptible to infection with associated high mortality and negligible recovery rates. On the other hand, infected adults and sub-adults exhibit negligible mortality and can recover with lifelong immunity (see Material and Methods for details).

RESULTS

We have computed disease prevalence in the population at equilibrium as a function of culling rate or hunting mortality for different values of age-dependent heterogeneity in transmission, called $\Delta R_0 = R_0 - R_0^*$ (where, $R_0$ is the transmission rate between adults and $R_0^*$ is the transmission rate between juveniles). We held constant the basic reproduction number $R_0$ in model (1) to keep the same level of infection in the population for each value of $\Delta R_0$ when culling is absent (see details in Protocol S2 in Supporting Information).

When the within class transmission rate for adults is larger than that within juveniles, the disease prevalence can actually increase with culling or hunting rate instead of decreasing as expected under the assumption of homogenous mixing (Fig. 1a); disease prevalence eventually peaks for intermediate values of culling and then decreases only for high level of animal removal. Moreover, the absolute number of infected individuals is larger in the case of heterogeneous mixing relative to the number under conditions of homogenous mixing at all levels of culling. The explanation for this effect is that at low and intermediate levels of culling, by removing older resistant individuals, population age structure is skewed in favor of highly susceptible juvenile hosts thus making culling ineffective. The minimum culling rate $C_{min}$ required to bring disease prevalence below the value attained in the absence of culling can be fairly high, as depicted in Fig. 1a. As a consequence, as long as $c < C_{min}$, culling actually performs worse, in terms of disease control, than the do-nothing alternative ($c = 0$). The dynamics of infection also reveal that the population density at
equilibrium decreases with culling more quickly in the presence of transmission heterogeneity (see Fig. 1b) even though, for intermediate values of culling, the actual number of infected hosts is larger in the case of heterogeneous transmission.

We also performed a sensitivity analysis in the transmission parameters by estimating the values of $C_{min}$ for a broad range of $R_0$ and $DB$ values. Fig. 2 shows that negative effects of culling (corresponding to $C_{min} > 0$) can be important when both basic reproduction number and transmission heterogeneity are sufficiently large, otherwise the model predicts a monotonic decrease in prevalence with the culling rate ($C_{min} = 0$).

DISCUSSION

The pattern revealed in Fig. 1a suggests that if the culling rate for disease eradication is computed by assuming homogenous mixing while transmission rate is actually age dependent, then classical culling strategies may prove to be ineffective: in fact, even though the host population is even more depressed then expected under homogenous mixing, not only will the disease not be eradicated from the population but prevalence can be even higher than in the absence of culling. As a consequence, unless it is possible to guarantee a sufficiently high removal of adult individuals, the do-nothing alternative is more effective and less costly than an intermediate culling strategy. For example, in the case of classical swine fever, to move from $c=0$ to $c_{min}$ requires removing at least 22.5% of individuals in the population. Similarly, intermediate levels of hunting pressure, especially when not aimed at disease control, might actually increase disease prevalence as well as the number of infected individuals. We show that similar counter-intuitive results are possible only in the presence of heterogeneity in transmission and sufficiently large $R_0$ values (see Fig. 2).

On the other hand, the maintenance of a high level of culling that guarantees disease eradication is not always feasible in practice. In fact, if the host population is very small, culling might generate conservation concern, as the removal of a large fraction of individuals might drive the host to the brink of extinction, as argued by Dobson & Meagher [25] for the eradication of Brucellosis in Yellowstone National Park bison. On the contrary, if the host population is very large, it might be impossible to cull a large enough fraction of individuals. This may correspond to the case, for instance, for huge colonies of bats in central Africa suspected of being the reservoir of Ebola and Marburg viruses [26] or populations of small rodents in North America, that comprise the main reservoirs for Hanta viruses or the vectors of Lyme disease [27].

The results presented in Fig. 1 contain an important insight concerning the potential for disease to spill-over into domestic animals. If transmission between wildlife and domestic animals is density-dependent, the risk of spill-over decreases for increasing culling rates even though it is higher than in the case of homogenous mixing (dotted lines in Fig. 1a); while if transmission between wildlife and domestic animals is frequency dependent, the risk of spillover can substantially increase for intermediate values of culling rate with respect to the case of homogenous mixing (solid lines in Fig. 1a). Therefore, it is crucial to articulate and understand the exact mechanisms by which infectious wild hosts interact with susceptible domestic animals and humans susceptible in order to predict the effects of wildlife culling on the risks of spillover.

We thus conclude that age/stage structure of transmission rate can be crucial in our understanding of disease pattern and the implementation of control policies. This conclusion likely applies to other wildlife diseases in which age/size structure is an important component of population dynamics.

Recently, Choisy and Rohani [24] presented a model of wildlife disease that focused on the effects of strong density-dependence and seasonality on culling and equilibrium disease prevalence. Their model was different from ours in that it was not age-structured and culling was random over the population of hosts. Nonetheless, they showed a similar response to culling with intermediate levels of culling producing a counter-intuitive increase in disease prevalence. Given the explicit difference in model structures it is instructive to speculate on how two very
different models can generate the same overall effect. As Choisy and Rohani indicate, the result of their model is driven by culling releasing the population from density-dependent reductions in the birth rate thereby producing a flush in new susceptible in the population. In our model we have a similar effect but driven by a completely different mechanism. The age-dependent culling coupled to the intrinsic heterogeneity in transmission similarly produces a flush in the relative abundance of the young susceptible class. We imagine that other mechanisms besides age/stage structure (as in our model) or strong density-dependence and seasonality (as in the Choisy and Rohani model), may interact with culling to produce similar patterns of prevalence.

Given the implications of transmission heterogeneities on dynamics and control, more detailed studies are thus necessary to assess these heterogeneities when structuring control strategies in current and ongoing wildlife epizootics [13,16,22]. What appears certain is that the negative effects of culling are robust to alternative model formulations and highlight the importance of better assessing transmission structure, seasonality, and population regulatory processes in the design of wildlife disease control strategies.

MATERIALS AND METHODS

The model is characterized by five classes: susceptible juveniles and adults, infected juveniles and adults and recovered (immune) adults.

The infection dynamics in this age-structured population are described by:

\[
\begin{align*}
\dot{S}_a &= v(S_a + R_a) - (\mu_a + \gamma_a)S_a - \beta_d S_a I_a - \beta_j S_j I_j - \mu_j S_j - \beta_a S_a I_a - c S_a - \\
\dot{I}_a &= \beta_d S_a I_a - (\mu_a + \gamma_a)I_a - c I_a - \\
\dot{S}_j &= \rho S_j - \mu_j S_j - \beta_d S_j I_j - c S_j - \\
\dot{I}_j &= \beta_d S_j I_j - (\mu_j + \gamma_j)I_j - c I_j - \\
\dot{R}_a &= \delta_a(a) - \mu_j R_j - \beta_j S_a R_j - c R_a - \\
\dot{R}_j &= \delta_j(a) - \mu_j R_j - \beta_j S_j R_j - c R_j - \\
\end{align*}
\]

where \( S(a), I(a), \) and \( R(a) \) refer to susceptible juveniles (adults), infected juveniles (adults), and recovered adults, respectively. The system parameters \( v, \mu, \rho, \) and \( \gamma \) represent the host birth rate, the juveniles mortality rate at low population density, the adults mortality rate, and the rate at which juveniles pass into adulthood, respectively. We assume that host population is self-regulating with density-dependent mortality in juveniles (\( \gamma \)) affected by total adults density \( (A = S_a + I_a + R_a) \). The force of infection for susceptible juveniles (adults) individuals is \( \lambda_{ju} = \beta_j(a) I_j \) where \( I = I_a + I_j \) is the total infectious density in the population. Parameters \( \alpha \) and \( \delta \) represent the disease-induced mortality in juveniles, and the adult recovery rate, respectively. Finally, \( \epsilon \) is the control parameter and represents the culling effort over the adults population. Only adult and sub-adult individuals are here assumed to be culled either because of conservation measures or, in the case of hunting, because of the preference for large trophies. Moreover, culling is not usually allowed in spring when peak fertility occurs and when most juveniles are born. By the onset of the hunting season, at the beginning of autumn, juveniles have already moved into the sub-adult age class.

Contact rate among adults increases dramatically during mating season (when males roam considerable distances in search of reproductive females). Consequently, within adult transmission rate is assumed to be larger than within juvenile transmission. It is this variation in the within-class transmission rates that forms the basis of transmission heterogeneity. We assign the magnitude of this heterogeneity in age-specific transmission as a new variable, \( \Delta\beta = \beta_a - \beta_j \geq 0 \) assessing the marginal increase of adult transmission rate \( \beta_a \) relative to that of juveniles \( \beta_j \).

SUPPORTING INFORMATION

Protocol S1 Parameter setting for CSF. Details about the setting of epidemiological and demographic parameters in model (1) for classical swine fever in wild boar. Found at: doi:10.1371/journal.pone.0000747.s001 (0.04 MB DOC)

Protocol S2 Basic reproduction number computation. Details about calculation of the basic reproduction number for model (1). Found at: doi:10.1371/journal.pone.0000747.s002 (0.05 MB DOC)

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Author Contributions

Conceived and designed the experiments: LB GD LR. Performed the experiments: LB GD LR. Analyzed the data: LB GD LR. Wrote the paper: LB GD LR.

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