RESEARCH ARTICLE

An SIS Epidemiology Game with Two Subpopulations

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There is significant current interest in the application of game theory to problems in epidemiology. Most mathematical analyses of epidemiology games have studied populations where all individuals have the same risks and interests. This paper analyzes the rational-expectation equilibria in an epidemiology game with two interacting subpopulations of equal size where decisions change the prevalence and transmission patterns of an infectious disease. The transmission dynamics are described by an SIS model and individuals are only allowed to invest in daily prevention measures like hygiene. The analysis shows that disassortative mixing may lead to multiple Nash equilibria when there are two interacting subpopulations affecting disease prevalence. The dynamic stability of these equilibria is analyzed under the assumption that strategies change slowly in the direction of self-interest. When mixing is disassortative, interior Nash equilibria are always unstable. When mixing is positively assortative, there is a unique Nash equilibrium that is globally stable.

Keywords: infectious disease, multiple populations, mixing patterns, differential inclusions

1. Introduction

The application of game theory to problems at the intersections of economics, public policy, and epidemiology is a relatively young area of study. While a handful of papers addressed the problem before the turn of the century [4, 9, 10, 12, 23], interest in games coupled to epidemiology has spread in the last five years. One of the central research tasks in this young field is an exploration of how different aspects of demography and epidemiology change the nature of solutions to the simple vaccination game and related models.

In the simple vaccination game for an endemic disease in a homogeneous population, there is a unique equilibrium behavior. If the vaccine is expensive or unsafe and risk from infection is small, the equilibrium behavior for individuals is to avoid vaccination. If vaccine is safe and cheap while the risks from infection are large, the equilibrium behavior is to vaccinate at rates almost sufficient to eradicate disease. As the cost of vaccine is varied continuously between these extremes, the equilibrium also varies continuously between these extremes [2, 9]. At the extremes of very cheap and very expensive vaccination, the equilibrium solution for individuals is approximately the same as the utopian ideal where everybody cooperates for the best interest of the community-at-large, but for intermediate costs of vaccine, there can be large differences between the Nash equilibrium and the community’s
ideal. Similar results hold for models where transmission is blocked by changes in behavior [5].

This cartoon of the equilibrium dependence on the parameters can change in important ways as various generalizations are incorporated into the simple vaccination game and its variants to make them more realistic. Age-dependent virulence, for instance, may change the game such that instead of there being a unique equilibrium, there are three equilibria and two of them are locally stable [25]. Both fast and slow rates of vaccination may be equilibria when vaccine provides imperfect protection [7, 20]. Other effects are likely to be discovered in the coming years, as the depth of our explorations of this field grows. One under-explored generalization is the subdivision of populations into two or more interacting subpopulations. Subpopulation structure, and interacting populations in general, will be our focus in this paper.

Several articles have studied epidemiology games with subpopulation structure. Some models have been structured solely in terms of the costs faced by individuals [4, 5, 7, 24]. In these cases, the equilibrium solutions to the games are consistent with the cartoon of the simple vaccination game. Two papers have explicitly addressed epidemiology games where the populations are also biologically structured. In Galvani et al. [11], the authors study an influenza vaccination in an age-structured population but with the age-groups treated as distinct subpopulations rather than life-stages in a single population. Galvani et al. find unique equilibria for the parameter ranges studied, but the numerical results provide limited insight into the general case. A more tractable model is studied by Chen [6]. In that paper, the author studies an SI model with varying degrees of positive assortative mixing, where individuals can invest in self-protective behavior. The author finds that for all the parameter values investigated, there is a unique equilibrium solution to the game.

The results of these papers suggest that population structure, per say, does not qualitatively change the solution set of epidemiology games. But the various extremes of simplicity and complexity in the studies to date leave the reasons for this pattern murky. This paper is intended as a start to clearing the waters regarding the effects of subpopulation structure. First, we describe the generalization of the approach of Reluga et. al [25] for modelling rational decisions of individuals in games with multiple interacting populations. We will then adapt the SIS model studied in [5] to describe two interacting populations with a full range of mixing patterns and study the equilibrium solutions to a population-game of self-protective behaviors. A graphical method for the analysis is presented. Consistent with the results of Chen [6], there is always a unique Nash equilibrium when mixing is positively assortative. However, in the case of disassortative mixing, there may be up to 3 isolated Nash equilibria. Thus, population structure can qualitatively change the behavior of equilibria in epidemiology games. We then propose a differential inclusion for strategy dynamics and show that interior Nash equilibria can be unstable in the case of disassortative mixing, but are stable in the case of assortative mixing.

2. Methods

We begin with the extension of the methods used in Reluga et al. [25] to population games with multiple interacting populations. A population game contains both population-scale components that describe the states of the populations and individual-scale components that describe the states of individuals within these populations. Suppose we have a set of $n$ populations labeled $i = 1 \ldots n$. Within
a population, an individual occupies one of many possible states. The number of individuals in population $i$ occupying each state at time $t$ is represented by a vector $\mathbf{x}_i(t)$ (vectors and matrices will be typeset in bold). We refer to $\mathbf{x}_i(t)$ as population $i$’s state with $x_{ij}(t)$ representing the number of individuals from population $i$ in state $j$ at time $t$. The state of an individual in population $i$ at time $t$ is represented by a probability measure $\mathbf{p}_i(t)$ over the possible states. We refer to $\mathbf{p}_i(t)$ as the individual’s state.

We wish to study the structure of a set of functions

$$ U_i(\pi_i, \pi_1 \ldots \pi_n) $$

(2.1)

representing expected utility to an individual from population $i$ playing strategy $\pi_i$ if all other individuals are playing the resident strategies $\pi_1 \ldots \pi_n$. We call the individual’s strategy $\pi_i$ an invading strategy to distinguish it from each population’s resident strategy $\bar{\pi}_i$.

Before calculating the utilities $U_i$, we must first study the dynamics of the processes of interest. The population-scale and individual-scale dynamics are driven by separate but related processes. Suppose the populations exist in an environment with state vector $\mathbf{e}(t)$. The population-scale dynamics satisfy a system of differential equations where the rates of change in the states of population $i$ depend on the states of the other populations, the resident strategies, the environmental state, and the time;

$$ \frac{d\mathbf{x}_i}{dt} = G_i(\mathbf{x}_1 \ldots \mathbf{x}_n, \pi_1 \ldots \pi_n, \mathbf{e}, t). \quad (2.2) $$

The rate of change in the environment’s state also depends on the states of the populations, the resident strategies, the environment’s current state, and time, or

$$ \frac{d\mathbf{e}}{dt} = G_e(\mathbf{x}_1 \ldots \mathbf{x}_n, \pi_1 \ldots \pi_n, \mathbf{e}, t). \quad (2.3) $$

We assume the populations are so large that the population dynamics are insignificantly affected by the few individuals with invading strategies that differ from the resident strategies.

While the large population sizes let us describe the population-scale dynamics deterministically, the individual-scale dynamics for an individual in population $i$ with invading strategy $\pi_i$ are stochastic. Each individual has risks and opportunities, but these can differ between individuals depending on the course of their life. Suppose that the state of an individual is governed by a continuous-time Markov process. The individual-scale dynamics describe changes in the probability that an individual will be found in any given state using a linear differential equation where the transition rates depend on the population states, the environment’s state, time, and the individual’s strategy, or

$$ \frac{d\mathbf{p}_i}{dt} = Q_i(\mathbf{x}_1 \ldots \mathbf{x}_n, \mathbf{e}, t, \pi_i)\mathbf{p}_i, \quad (2.4) $$

where $Q_i$ is a matrix of transition rates. We have allowed the transition rate to only depend on the resident strategies through the population’s states, but this is easily relaxed.

Individuals may choose behavior strategies based on several different criteria. One way of quantifying an individual’s decision process is to define an “expected utility” function that associates a payoff to each strategy choice and to then assume
individuals make decisions to maximize this payoff. This approach is commonly referred to as “rational expectation” theory in economics. The expected utility for an individual with an invading strategy can be calculated based on the individual-scale dynamics. Given an initial time $t_0 = 0$ and an infinite horizon, the utility of an invading strategy $\pi_i$ to an individual in population $i$ give resident population strategies $\pi_1 \ldots \pi_n$ is

$$U_i(\pi_i, \pi_1 \ldots \pi_n) = \int_0^\infty e^{-h_i t} v_i^T p_i(t) dt$$

(2.5)

where $h_i$ is the discount rate in population $i$ and $v_i$ is a vector of felicities, a.k.a. utility gains, for residence in each state in population $i$. Equation (2.5) can be interpreted as a path integral over all an individual’s possible life-histories and can be derived from the Markov decision process theory developed by Howard [16].

The felicities have two parts: a vector of expected gains per unit time $f_i$ for each state, and a matrix $F_i$ of instantaneous utility gains associated with the transitions between states, where $F_{i,jk}$ is the expected instantaneous gain of an individual in population $i$ jumping from state $k$ to state $j$. Using a result of Howard [16], it is shown that

$$v_i^T = f_i^T + 1^T (F_i \circ Q_i)$$

(2.6)

where $1$ is a vector of 1’s and $\circ$ represents the Hadamard product of two matrices. The Hadamard product of two matrices is the matrix of the product of the components: $(F_i \circ Q_i)_{jk} = F_{i,jk}Q_{i,jk}$.

While population games can be solved and studied in the general form we have so far presented, it is often useful to focus on cases where the population-scale dynamics have a simple attractor. If all dynamics are autonomous and $(x^*_1 \ldots x^*_n, e^*)$ is a stationary solution of the population-scale dynamics, the transition rate matrix

$$Q^*_i = Q_i(x^*_1 \ldots x^*_n, e^*, \pi_i)$$

(2.7)

is also stationary and

$$p_i(t) = e^{-tQ^*_i} p_i(0).$$

(2.8)

If there is a positive discount rate $h_i > 0$ or the Markov process is transient e.g. the dominate eigenvalue $\lambda_0(Q^*_i) < 0$, then over the infinite time horizon, the expected utility has closed form

$$U_i = v_i^T (h_i I - Q^*_i)^{-1} p_i(t_0)$$

(2.9)

where $I$ is the identity matrix.

3. Application in an SIS model

We now study the rational expectation equilibria of utility functions calculated using the methods described above when the dynamics are governed by a standard epidemic model. We will first describe a stochastic model at the individual scale and use the result to motivate a population-scale model that is used in turn to close the equations for the utility calculation. To study the equilibrium structure of an epidemic population game with interacting populations, we will focus on a
2-population SIS model that is a variation of the work of Chen [5]. In this model, individuals can reduce their risk of infection by adopting costly behaviors that prevent transmission and in-turn change the prevalence of the infectious agent.

Take a population of individuals of a single species that is split into two distinct subpopulations. All individuals of this species can be infected by the same pest or pathogen. The situation can be interpreted as representing for example, the transmission of lice within and between two classrooms of students in an elementary school, or the sexual transmission of gonorrhea between men and women. In order to conduct our analysis, we are omitting the biological details like stochasticity needed in practical models of lice or gonorrhea transmission, so our results should not be applied “out-of-the-box” to either of these cases. Rather, the results should guide future research and applications in disease-specific studies.

The duration of infectiousness is assumed to be exponentially distributed with expectation $1/\gamma$, independent of an individual’s subpopulation. When individuals recover, they do not gain any form of immunity against reinfection, so they immediately return to a susceptible state. The risk of infection to a susceptible individual in subpopulation $i$ is proportional to the force of infection $\lambda_i$. Individuals may choose to reduce their personal risk to a fraction $\sigma_i$ relative to the background risk per day by changing behaviors. Throughout the paper, the relative risk $\sigma_i$ will be referred to as the individual’s strategy, since it’s value is determined by an individual’s choices.

Under these assumptions, the changes in an individual’s state are described by a continuous-time Markov process

$$\frac{dp_i}{dt} = Q_i p_i,$$

(3.1a)

where $p_i(t)$ is the probability that an individual in population $i$ is susceptible or infected at time $t$ and the transition-rate matrix

$$Q_i = \begin{bmatrix} -\lambda_i \sigma_i & \gamma \\ \lambda_i \sigma_i & -\gamma \end{bmatrix}.$$  

(3.1b)

Every individual in the population starts in the susceptible state, so

$$p_i(0) = [1 \ 0]^T.$$  

(3.1c)

Eq. (3.1) is non-autonomous and can not be solved in closed form without further assumptions because the force of infection $\lambda_i$ is time-dependent.

The force of infection seen by an individual is determined at the population-scale [22]. In a population with sufficiently strong mixing, we can represent the transmission dynamics using a standard SIS epidemic model. In subpopulation $i$, let $S_i(t)$ represent the number of susceptible individuals at time $t$ and $I_i(t)$ represent the number infected individuals at time $t$. To simplify our analysis, let’s assume that the two subpopulations have the same sizes which we have normalized to 1, so that $S_1 + I_1 = 1$ and $S_2 + I_2 = 1$.

The forces of infection $\lambda_i$ will be modelled as a weighted sum of the number of infected individuals in each subpopulation;

$$\lambda_1 = \beta_{11} I_1 + \beta_{21} I_2,$$

$$\lambda_2 = \beta_{12} I_1 + \beta_{22} I_2,$$

(3.2a)

(3.2b)
where $\beta_{ij}$ represents the background transmission rate from individuals of subpopulation $i$ to individuals of subpopulation $j$. Since there are no temporal changes in the total population sizes, this may be interpreted as either standard-incidence or bilinear-incidence. Unless otherwise specified, we will assume the matrix $[\beta_{ij}]$ is positive.

To scale up the effects of individual choices to the population dynamics, we make the simplest reasonable supposition. In subpopulation $i$, almost all individuals use the behaviors that reduce their infection rate to a fraction $\sigma_i$ of the background infection rate. We call $\sigma_1$ and $\sigma_2$ the resident strategies and for shorthand we write $\sigma = (\sigma_1, \sigma_2)$. Under this postulate, population-scale dynamics of infection are governed by the system of differential equations

\begin{align*}
\dot{S}_1 &= \gamma I_1 - \lambda_1 \sigma_1 S_1, \quad (3.3a) \\
\dot{I}_1 &= \lambda_1 \sigma_1 S_1 - \gamma I_1, \quad (3.3b) \\
\dot{S}_2 &= \gamma I_2 - \lambda_2 \sigma_2 S_2, \quad (3.3c) \\
\dot{I}_2 &= \lambda_2 \sigma_2 S_2 - \gamma I_2. \quad (3.3d)
\end{align*}

The SIS model described by Eq. (3.3) and its variants have been studied extensively in the past [3, 14].

4. Game Equilibria

To formulate a population game, we must determine utilities for various strategies. As described above, we can calculate utilities by inspecting the costs and benefits of isolated events in an individual’s life and adding them all up under appropriate discounting.

For the purpose of our analysis, we propose that individuals must balance two simple costs. One cost, $c_i(I)$, is the cost paid per day of infection by an individual of subpopulation $i$. The other cost, $c_i(B)$, is the cost of self-protecting behaviors per day per percent reduction in infection risk. For a fixed force of infection, we might expect an individual’s risk of infection to be a monotone decreasing function of their investment in self-protection. It will be convenient to assume that the daily risk of infection is a convex function reaching 0 at a finite value of protection investment. There are economic arguments why risk may not be a convex function of pure strategies and good biological reasons why risk may never reach 0. However, these issues would complicate our analysis. In this model, we will assume that an individual must invest $(1 - \sigma_i)c_i(B)$ per day to reduce one’s risk to a fraction $\sigma_i$ of the background risk in population $i$. Thus, the vectors of felicities for the susceptible or infected states in each subpopulation $i$ will be

$$v_i = [- (1 - \sigma_i)c_i(B), -c_i(I)]^T. \quad (4.1)$$

Our model places some important restrictions on the possible applications of the results. For instance, we assume that the prevention is a daily hygiene activity like hand-washing, and does not change the individual’s biological state. Preventative measures like vaccination can not be represented here because they introduce a long-term change in an individual’s state for a one-time cost, which we have not allowed (but see [1, 11, 24]). Similarly, this system does not immediately apply to antiviral prevention, since antivirals change an individual’s state in a way that affects not only susceptibility but also recovery. The costs of infection are also daily
costs that persist for exactly the duration of infection. This does not allow for
treatment, since treatment would involve a choice that could shorten the duration
of infection and affect daily costs. Nor have we allowed for long-term disabilities
incurred from infection because the SIS theory assumes recovered individuals are
indistinguishable from naive individuals. All of these events and others could be
incorporated into our system with the addition of appropriate states, transitions,
costs, and control variables, but would lengthen and complicate our analysis.

Given individual-scale and population-scale models (Eq. (3.1) and Eq. (3.3),
respectively), together with felicities of possible events and behaviors, we can
now investigate the equilibria available. For an equilibrium analysis, we assume
System (3.3) is at the globally attracting stationary solution, with stationary
transition-rates $Q_i^*$ and stationary forces of infection $\lambda_i^*$. Calculated over an in-
finite time-horizon, the utility of individual strategy $\sigma_i$ is

$$U_i(\sigma_i, \sigma_1, \sigma_2) = v_i^T (h\mathbf{1} - Q_i^*)^{-1} p_i(0) = \frac{-c_i(B(1-\sigma_i)(h+\gamma) - c_i(I)\lambda_i^*\sigma_i)}{h(h+\gamma+\lambda_i^*\sigma_i)} \quad (4.2)$$

where $\lambda_i^*$ is an implicit function of $\sigma_1$ and $\sigma_2$.

The discount rate $h = h_1 = h_2$ acts as a catch-all for independent environmental
risks that reduce incentives for planning ahead. In the limit of large discount rates,
the future risks from infection are very small compared to the current costs of
prevention. In the limit of small discount rates, the utility diverges because every
future illness has the same costs as a current illness. Thus, tuning $h$ allows us to
account for some risks external to our epidemic model.

Since the subpopulation sizes equal 1 by assumption, $S_i = 1 - I_i$ at all times.
After this substitution, determining the stationary solutions to Eq. (3.3) reduces
to simultaneously solving the pair of quadratic equations

$$\sigma_1 (1 - I_1^*) (\beta_{11} I_1^* + \beta_{21} I_2^*) - \gamma I_1^* = 0, \quad (4.3a)$$
$$\sigma_2 (1 - I_2^*) (\beta_{12} I_1^* + \beta_{22} I_2^*) - \gamma I_2^* = 0. \quad (4.3b)$$

One stationary solution is the disease-free stationary solution $(S_1, I_1, S_2, I_2) =
(1, 0, 1, 0)$. The disease-free stationary solution is globally attracting so long as the
reproductive number $R < 1$ [14] (see Appendix A). If $R > 1$, there is exactly 1
stationary solution with positive infection prevalence, and it is globally attracting
[3]. We will refer to this solution as the endemic stationary solution.

The role of between-subpopulation transmission relative to within-subpopulation
transmission in asymptotic disease prevalences can be partially understood in terms
of source-sink concepts. A subpopulation is a source of infections if the infection
can persist in that population in isolation. A subpopulation is a sink population
if infection cannot persist in the subpopulation in isolation. Subpopulation $i$ is a
source if $\beta_{ii}/\gamma > 1$ and a sink if $\beta_{ii}/\gamma < 1$. If either subpopulation is a source,
then $R > 1$. But in some cases, between-subpopulation transmission can be strong
enough such that $R > 1$ even when both subpopulations alone are sinks.

The endemic stationary solution has the property that it is decreasing in the
resident strategies $\sigma_i$, and strictly decreasing so long as $R > 1$, $\sigma_1 \beta_{11}/\gamma > 1$, and
$\sigma_2 \beta_{22}/\gamma > 1$ (see Appendix A and Figure 1). The monotonicity of the endemic
stationary solution makes it easier for us to calculate the game equilibria. As the
resident strategies $\sigma$ are varied, there is a region of reachable stationary disease
prevalences. The shape of this region depends on whether the subpopulations are
sources or sinks (Figure 1). For each point in the reachable region of prevalences,
we can uniquely determine the forces of infection exerted on individuals in each
subpopulation. Individual best responses are correspondences\(^1\) depending on these forces. From the individual utilities, we can see that the best responses are step functions. Differentiating the utility with respect to \(\sigma_i\), we find that the best response correspondence is given by

\[
\text{argmax}_{\sigma_i} U_i(\sigma_i, \sigma_1, \sigma_2) = \begin{cases} 
0 & \text{if } \lambda^*_i(\sigma_1, \sigma_2) > \lambda^+_{i}, \\
[0, 1] & \text{if } \lambda^*_i(\sigma_1, \sigma_2) = \lambda^+_{i}, \\
1 & \text{if } \lambda^*_i(\sigma_1, \sigma_2) < \lambda^+_{i}.
\end{cases}
\] (4.4)

where the threshold forces of infection

\[
\lambda^+_{i} = \begin{cases} 
c_i(B)(h + \gamma) & \text{if } c_i(I) > c_i(B), \\
\frac{c_i(I) - c_i(B)}{c_i(B)} + \infty & \text{otherwise}.
\end{cases}
\] (4.5)

The bang-bang form of this best-response correspondence is reminiscent of linear optimal control theory, and is a direct consequence of Theorem B.1 in Appendix B. If the force of infection is below the threshold \(\lambda^+_{i}\) in subpopulation \(i\), individuals prefer no self-protection \((\sigma_i = 1)\). If the force of infection is above the threshold value \(\lambda^+_{i}\), individuals prefer complete self-protection \((\sigma_i = 0)\). If the force of infection is exactly the threshold value, individuals will have the same utility for all self-protection strategy choices. This last condition implies that if the threshold forces of infection intersect within the reachable region, the strategy pair \(\sigma^+ = (\sigma_1^+, \sigma_2^+)\)

\(^1\)A “correspondence” maps each element of the domain to a subset of the range.
Figure 2. Depending on the transmission patterns and the relative costs of protection to infection, there may be one or three solutions to the population game. The threshold forces of infection are plotted as lines over the reachable regions of disease incidence and the dots mark game equilibria. If the costs of protection are sufficiently large for both populations, the endemic prevalences will be an equilibrium state (A). If the costs of protection are lower, but do not cross with the reachable region of prevalences, the costs of protection are solely carried by the population with the lower threshold (B). If the thresholds cross within the reachable region and the direct transmission risks are greater than the indirect transmission risks ($\beta_{11} \beta_{22} > \beta_{12} \beta_{21}$), there is a single internal equilibrium (C). If the thresholds cross within the reachable region and the direct transmission risks are less than the indirect transmission risks ($\beta_{11} \beta_{22} < \beta_{12} \beta_{21}$), there will be three equilibria (D). Parameter values: $h = 0$, $\gamma = 1$, $c_1(I) = c_2(I) = 1$.

for the disease prevalences corresponding to the intersection ($\lambda^*_1 = \lambda^*_2$, $\lambda^*_1 = \lambda^*_2$) must constitute a Nash equilibrium.

In general, a strategy pair ($\sigma_1^*, \sigma_2^*$) is a Nash equilibria of the utility functions $(U_1, U_2)$ if the strategies are best-responses to themselves for all individuals in all subpopulations. The variational-inequality approach described in Cojocaru et al. [8] supplies one approach to solving these equations. Instead, we’ll adopt a graphical approach. The Nash equilibria to the population game can be determined by plotting the threshold forces of infection over the reachable region (Figure 2). The best-response correspondences are step-functions, so all equilibria of the game must lie on the boundaries of the feasible region and at threshold values of the forces of infection. If the stationary prevalences in the absence of self-protection are below both threshold forces of infection for both subpopulations ($\lambda^*_i(\sigma_1 = 1, \sigma_2 = 1) < \lambda^*_i$ for each $i$), then the only Nash equilibrium will be no self-protection in either subpopulation ($\sigma_1^* = \sigma_2^* = 1$) (Figure 2A). If the threshold forces of infection are not positive ($\lambda^*_1 \leq 0, \lambda^*_2 \leq 0$), then complete self-protection $\sigma_1^+ = \sigma_2^+ = 0$ is the best response under all stationary prevalences, and thus is the unique Nash equilibrium.

In between these extremes, one or both subpopulations will prefer some intermediate investment in self-protective behaviors. At this point, it’s convenient to
define some terminology. Let the set

$$T_j = \{(I_1^*, I_2^*): \beta_{1j} I_1^* + \beta_{2j} I_2^* \geq \lambda_j^+\}. \tag{4.6}$$

We say that subpopulation $i$ has a lower threshold than subpopulation $j$ if and only if $T_j \subset T_i$ and that subpopulation $i$ has a higher threshold than subpopulation $j$ if and only if $T_i \subset T_j$.

If the endemic stationary solution in the absence of self-protection ($\bar{\sigma} = (1, 1)$) generates a force of infection greater than the threshold force of infection in at least one subpopulation and no disease prevalences in the feasible region impose forces of infection at threshold levels for both subpopulations (the thresholds plotted in Figure 2 do not cross), there will be a unique Nash equilibrium with some self-protection in the subpopulation with the lower threshold and less-than-complete self-protection in the subpopulation with the higher threshold. To illustrate this, let us assume that, as in Figure 2B, subpopulation 2 has a lower threshold than subpopulation 1. If self-protection within subpopulation 2 alone can reduce the force of infection to subpopulation 2’s threshold value $\lambda_2^+$, then there is a unique Nash equilibrium $\sigma^-$ with $\sigma_1^- = 1$, $0 < \sigma_2^- < 1$, where $\lambda_1^* < \lambda_1^+$ and $\lambda_2^* = \lambda_2^+$. If self-protection within subpopulation 2 alone can not reduce the force of infection to subpopulation 2’s threshold but can reduce the force of infection below subpopulation 1’s threshold, then there is a unique Nash equilibrium $\sigma^-$ with $\sigma_1^- = 1$, $\sigma_2^- = 0$. If self-protection within subpopulation 2 alone can not reduce the force of infection below the thresholds of either subpopulation, then there is a unique Nash equilibrium $\sigma^-$ with $\sigma_2^- = 0$, $0 < \sigma_1^- < 1$, where $\lambda_1^* = \lambda_1^+$.

If there are disease prevalences within the reachable region such that the stationary forces of infection equal their threshold values for both subpopulations simultaneously (as in Figure 2C and Figure 2D), then, as stated above, the corresponding strategy pair $\sigma^+$ is a Nash equilibrium. The uniqueness of this equilibrium depends on the sign of the threshold intersection.

If $\beta_{11}\beta_{22} - \beta_{12}\beta_{21} > 0$, then the mixing pattern for transmission is assortative and self-protective behaviors preferentially reduce transmission within the same subpopulation. When the intersection of the thresholds occurs within the reachable region, self-protection that reduces the force of infection to threshold levels within a single subpopulation is insufficient to reduce the forces of infection below threshold levels in the other subpopulation. There will be no Nash equilibria on the boundaries of the reachable region, and the interior equilibrium is the unique Nash equilibrium (see Figure 2C). This is a minor variation of the special case considered in [6].

If $\beta_{11}\beta_{22} - \beta_{12}\beta_{21} < 0$, then the mixing pattern for transmission is disassortative, and self-protective behaviors preferentially reduce the force of infection in the other subpopulation. When the intersection $\sigma^+$ of the thresholds occurs within the reachable region, self-protection within either subpopulation in isolate can reduce the forces of infection in the both subpopulation below threshold levels (see Figure 2D). The interior equilibrium will not be unique because there will be Nash equilibria on upper left ($\sigma^-$) and lower right ($\sigma^+$) boundaries of the reachable region. Thus, while assortative mixing leads to unique equilibria, disassortative mixing can create multiple Nash equilibria. In degenerate cases when the threshold forces of infection coincide in both populations ($\beta_{11}\beta_{22} - \beta_{12}\beta_{21} = 0$ and $\lambda_1^+\beta_{22} - \lambda_2^+\beta_{21} = 0$), there is a whole segment of Nash equilibria.
5. Stability of the Game Equilibria

A Nash equilibrium is a situation where individuals have no reason to change their minds about strategies if everything starts out just right. But a strategy that lets us “win” the game no matter how things start out would be even more useful. This leads us to the concepts of convergence and invasion potential. In particular, if the resident strategies are near the equilibrium, can individuals do better by using the equilibrium strategies? As Figure 3 shows, when the resident strategies are perturbed from the internal equilibrium $\sigma^+$, the internal equilibrium may or may not be able to re-invade depending on the direction of the perturbation. Local invasion potential can be perturbation dependent for either orientation of the crossings. In fact, an interior Nash equilibrium can only have universal local invasion potential if the response-thresholds for both populations are locally independent. By independent, we mean that if $(\sigma^*_1, \sigma^*_2)$ is an internal Nash equilibrium where

$$U_1(\sigma^*_1, \lambda^*(\sigma^*)) = U_1([0, 1], \lambda^*(\sigma^*)) \quad \text{and} \quad U_2(\sigma^*_2, \lambda^*(\sigma^*)) = U_2([0, 1], \lambda^*(\sigma^*)),$$

then the best response thresholds $\hat{\sigma}_1(\sigma_2)$ and $\hat{\sigma}_2(\sigma_1)$ are locally independent when

$$\frac{\partial \hat{\sigma}_1(\sigma_2)}{\partial \sigma_2} \bigg|_{\sigma_2 = \sigma_2^*} = 0, \quad \text{and} \quad \frac{\partial \hat{\sigma}_2(\sigma_1)}{\partial \sigma_1} \bigg|_{\sigma_1 = \sigma_1^*} = 0.$$

(5.2)
So interior Nash equilibria with different orientations of the threshold crossings are not distinguished simply by the local invasion potential. However, it does seem that there should be some fundamental dynamic difference between assortative and disassortative mixing systems (Figure 2C and Figure 2D respectively).

To formalize our intuition about dynamics, we adopt a variation of the adaptive-dynamics approach. Assume resident strategies change sufficiently slowly to preserve the stability of the endemic disease equilibria and always change in the direction of individuals’ best interests, but the relative rates of change are unknown.

Can we say anything about stability of stationary solutions? This problem is one of establishing stability of a stationary solution in a differential inclusion. Suppose

\[ \dot{\sigma}_1 = G_1(t)[A_1(\sigma_1, \sigma_2) - \sigma_1] \]  
\[ \dot{\sigma}_2 = G_2(t)[A_2(\sigma_1, \sigma_2) - \sigma_2] \]

where \( G_1(t) \) and \( G_2(t) \) are piecewise continuous functions with finite upper bounds and positive lower bounds, and

\[ A_i(\sigma_1, \sigma_2) = \begin{cases} 
0 & \text{if } \lambda_i^*(\sigma_1, \sigma_2) > \lambda_i^+ \\
\sigma_i & \text{if } \lambda_i^*(\sigma_1, \sigma_2) = \lambda_i^+ \\
1 & \text{if } \lambda_i^*(\sigma_1, \sigma_2) < \lambda_i^+ 
\end{cases} \]

Our choice of \( A_i \) is dictated by the assumption that resident strategies will change in the direction of individual self-interest. Under this description of strategy dynamics, the resident strategy of each population changes slowly in direction preferred by the self-interest of individuals but the relative rates of change can be arbitrary.

Substitution shows the Nash equilibria strategy pairs are stationary solutions of the dynamics. The differential equations are non-autonomous and there may be discontinuities in the coefficients, but we can still obtain some stability results for the equilibria.

**Theorem 5.1:** Suppose the Nash equilibrium \( \sigma^+ \) exists for the population game described by Eq. (4.2). If \( \beta_{11} \beta_{22} - \beta_{12} \beta_{21} > 0 \), then \( \sigma^+ \) is a locally stable solution to Eq. (5.3). If \( \beta_{11} \beta_{22} - \beta_{12} \beta_{21} < 0 \), then \( \sigma^+ \) is a locally unstable stationary solution to Eq. (5.3).

**Proof:** It will be helpful to refer to the cartoon shown in Figure 4, where the thresholds divide the resident-strategy phase-space up into quadrants numbered 1 to 4 in the neighborhood of an interior Nash equilibrium \( \sigma^+ \). First, we prove that the orientations of the thresholds in strategy space are the same as orientations in incidence space. The we study stability based on the threshold orientations in strategy space.

For each pair of resident strategies and each time, the velocity of change in System (5.3) takes on one of a cone of values bounded away from 0 except at the Nash equilibria. These cones can be characterized based on quadrant, as in the examples in Figure 4. The nullclines of System (5.3) solve \( \lambda_i^*(\sigma_1, \sigma_2) = \lambda_i^+ \). Since, the stationary incidences \( I_1^* \) and \( I_2^* \) are decreasing functions of \( \sigma_1 \) and \( \sigma_2 \) (see Appendix A) and \( \lambda_i^* \) is a positively weighted sum of the stationary incidences, then the force of infection \( \lambda_i^* \) is also a decreasing function of the resident self-protection probabilities. It follows that the thresholds in Figure 4 must have non-positive slopes. In addition, the Jacobian determinant of the coordinate transform between
Figure 4. Cartoon of phase-plane for the differential inclusion described by Eq. (5.3) in the neighborhood of the $\sigma^+$ (large dot) when (A) $\beta_{11}\beta_{22} - \beta_{12}\beta_{21} > 0$ and (B) $\beta_{11}\beta_{22} - \beta_{12}\beta_{21} < 0$. The solid line is the threshold for population 1. The dotted line is the threshold for population 2. The arrows clusters represent the cones of available directions within each of four quadrants labeled 1 – 4 and along the thresholds.

\[
\frac{d\sigma}{dI} = \gamma^2 \left( 1 + \frac{\beta_{12}I_1^*}{(\lambda_1^*)^2} + \frac{\beta_{21}I_2^*}{(\lambda_2^*)^2} \right) > 0
\]  

(5.4)

everywhere when $I^*$ is strictly positive. This ensures that the sign of the threshold crossing in $(\bar{\sigma}_1, \bar{\sigma}_2)$ space is the same as the sign in $(I_1^*, I_2^*)$ space. Thus, the transforms of plots like those in Figure 2 to resident strategy space like the plots in Figure 3 consists only of stretchings (no flips or folds).

Since the threshold curves have non-positive slopes, any trajectories starting initially in quadrants 3 or 4 must eventually be absorbed by quadrants 1 and 2. By inspecting the vector-field cones on the boundaries of quadrants 1 and 2, we find that quadrants 1 and 2 are invariant. Within quadrants 1 and 2, the dynamics are monotone. If $\beta_{11}\beta_{22} - \beta_{12}\beta_{21} > 0$, as in Figure 4A, the dynamics converge to the interior Nash equilibrium $\sigma^+$. But if $\beta_{11}\beta_{22} - \beta_{12}\beta_{21} < 0$, as in Figure 4B, the dynamics will diverge from the internal Nash equilibrium $\sigma^+$. □

A similar argument can be used to show that $\sigma^-$ and $\sigma^-$ are locally dynamically stable under Eq. (5.3) when they exist. Thus, we find stability properties very reminiscent of the Lotka competition models. If transmission is stronger between populations than within populations, there may be two boundary Nash equilibria that are locally stable, but if transmission is stronger within populations, a unique interior Nash equilibrium is always globally stable.

6. Discussion

In this paper, we have described a modelling approach to games in structured populations, and applied the approach to study the rational-expectation equilibria in a two-population game where choices reduce the risk of infection. We find that, depending on the mixing patterns and relative costs, there may not be a unique Nash equilibrium. Nash equilibria appearing on the interior of the reachable region of stationary incidences are stable against slow changes in the resident strategies when mixing is assortative, but unstable against slow changes in the resident strategies when mixing is disassortative. If transmission is stronger between populations than within populations, the game may have an unstable equilibrium and two stable equilibria satisfying the rational-expectations criteria. We conjecture that all Nash equilibria on the boundary of the reachable region are locally stable if there
is an unstable interior Nash equilibrium. The graphical analysis we've used to illustrate equilibria solutions is reminiscent of the phase-plane analysis of Lotka's competition model from biology [17] and Cournot's duopoly model from economics [13]. To some extent, this similarity is superficial, since the underlying mechanisms are different from either of these two classical works. The option of having a graphical shorthand supplementing mathematical and verbal reasoning will hopefully be useful to some readers.

In these analyses, we have avoided special cases for the sake of the larger picture. In cases where the transmission rate matrix $[\beta_{ij}]$ is non-negative but not strictly positive, Perron–Frobenius theory and related results should lead to a natural dissection of the games equilibrium structure. For instance, if there is no cross-transmission between subpopulations, the subpopulations can be treated as independent populations. Cases where $R_0 = 1$ or $\lambda^* = 0$ that appear in various calculations can be dealt with piecemeal. Also, if the threshold forces of infection are the same in both populations e.g. the lines in Figure 2 lie on top of each other, then there is a continuous interval of weak Nash equilibria. If the threshold forces of infection are proportional between the populations e.g. the lines in Figure 2 are parallel, then the problem reduces to one of heterogeneity in costs [4]. In some circumstances, special cases like these may be very useful as illustrations of specific phenomena, and care should exercised when studying systems near these special cases. Similarly, analysis in cases of unequal subpopulation sizes should be straightforward.

The practical importance of multiple equilibria in this example of an epidemiological game is unclear. The regions of parameter space for which multiple equilibria occur are relatively narrow because, as the transmission dynamics become more strongly disassortative, the reachable region of forces of infection becomes narrower. If the precision of information used in individual decisions is too low, actors will not be able to distinguish among the various equilibria, and analysis of the distinction in this particular model will be moot. More research is needed to establish whether this narrowness is common or unusual, depending on the economic, demographic, biological, and epidemiological structures involved. In addition, multiple equilibria may be created by other mechanisms. Our results depend strongly on the monotonicity result of Appendix A. Monotonicity does not necessarily hold for generalizations of this model, since there may be multiple stationary solutions for given parameter values and these may not be monotone functions of the resident self-protection strategies. The generalization of our results to other behaviors and mechanisms is also unclear. It seems reasonable to conjecture that this analysis will hold under other control mechanisms like vaccination, but other game-theory results like those of [18] suggest there may be a great diversity of results that depend on the specifics of the models.

This analysis has focused on the very constrained problem managing infectious disease risk, while neglecting external factors. To some extent, independent external risk factors like car crashes are accounted for by the discount rate $h$. When there are few external risks, discounting is small ($h \approx 0$), but when external risks are many, the discount rate is large. However, discounting can not account for additional external effects correlated to internal states and behaviors. Greater model specificity would be needed to incorporate such additional external effects. A related issue is the relative sizes of costs incurred from prevention and illness. The importance of a rational-expectation equilibrium depends greatly on the size of the costs incurred under sub-optimal behavior. If the expected costs of sub-optimal behavior are small compared to the variance of internal and external costs, it seems unreasonable to expect individuals to put much thought into optimizing behavior.
The results should generalize to some extent when considering arbitrary numbers of subpopulations, but the best approach to multi-group problems is not obvious. The results in the case of positive assortative mixing [6] rely on a convenient mathematical form for the force of infection from which a generalization is not immediately clear. It seems reasonable to conjecture that multiple equilibria become more likely as the number of subpopulations increases, given May’s observations regarding $n$-species competition models [21], but narrowing of the windows of reachable disease prevalences suggests that internal equilibria become less likely as the number of subpopulations increases. Another open problem is to investigate the trade-off between these effects.

Our suppositions in the derivation of the population-scale model from the individual-scale can be weakened or reinterpreted in several ways. For instance, $\bar{\sigma}_i$ can be interpreted as an average, or aggregate rate of self-protection in a heterogeneous population. This broadens the application of our results, but doesn’t change any of the fundamental results of the analysis. The “hazard” strategy-space considered in this paper is a subset of larger strategy spaces. A mixed strategy space might be a set of probability measures defined on a subset of functions $\mathbb{R} \mapsto [0,1]$ mapping a point in time to a specific behavior. This could include strategies where individuals change their behaviors systematically from day-to-day even when their state has not changed. Correlations among individual behaviors could then lead to significant effects at the scale of the population dynamics, for which we would need to account. But despite it’s limitations, consideration of a “hazard” strategy-space seems to yield useful results.

While we have found some insight into how population structure can change Nash equilibria in epidemiology games, clearly there are opportunities for further study.

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References


Appendix A. Uniqueness and monotonicity

For the SIS model considered in this paper, the basic reproductive number of the disease-free state in the absence of self-protective behavior is

$$R_0 = \rho_0 \left( \begin{bmatrix} \frac{\beta_{11}}{\gamma} & \frac{\beta_{21}}{\beta_{22}} \\ \frac{\beta_{12}}{\gamma} & \frac{\beta_{22}}{\gamma} \end{bmatrix} \right)$$  \hspace{1cm} (A1)

The disease-free reproductive number with possible self-protective behavior is

$$\mathcal{R} = \rho_0 \left( \begin{bmatrix} \frac{\sigma_1}{\beta_{11}} & \frac{\beta_{21}}{\beta_{22}} \\ \frac{\sigma_2}{\beta_{12}} & \frac{\beta_{22}}{\gamma} \end{bmatrix} \right),$$  \hspace{1cm} (A2)

which is the largest solution of the quadratic equation

$$\left( \mathcal{R} - \frac{\sigma_1}{\gamma} \frac{\beta_{11}}{\gamma} \right) \left( \mathcal{R} - \frac{\sigma_2}{\gamma} \frac{\beta_{22}}{\gamma} \right) - \sigma_1 \sigma_2 \frac{\beta_{12} \beta_{21}}{\gamma^2} = 0.$$  \hspace{1cm} (A3)

The reproductive number is bounded,

$$R_0 \geq \mathcal{R} \geq \max \left\{ \frac{\sigma_1}{\gamma}, \frac{\sigma_2}{\gamma}, \sqrt{\frac{\sigma_1 \sigma_2 \beta_{12} \beta_{21}}{\gamma^2}} \right\}.$$  \hspace{1cm} (A4)

We will now provide an alternative proof of the uniqueness of a positive stationary solution when $\mathcal{R} > 1$.

**Theorem A.1:** If $[\beta_{ji}]$ is a positive matrix with a unique inverse and $\mathcal{R} > 1$, then Eq. (3.3) has a unique endemic stationary solution. If $\mathcal{R} < 1$, there is no endemic stationary solution.

**Proof:** An endemic stationary solution is a stationary solution with a positive number of infecteds ($I_1 I_2 \geq 0, I_1 + I_2 > 0$). Eq. (4.3), supplying the stationary
solution condition, can be rewritten as the matrix equation

\[
\begin{bmatrix}
\beta_{11} & \beta_{21} \\
\beta_{12} & \beta_{22}
\end{bmatrix}
\begin{bmatrix}
I_1 \\
I_2
\end{bmatrix}
= 
\begin{bmatrix}
\frac{\gamma}{\sigma_1} I_1 \\
\frac{\gamma}{\sigma_2} I_2
\end{bmatrix}.
\]

(A5)

The inverse of \([\beta_{ij}]\) is an M-matrix so long as it exists. Eq (A5) can be rewritten as the equation

\[H(I) - I = 0\]  (A6a)

where

\[H_i(I) = \frac{\sigma_i}{\gamma} \sum_j \beta_{ji} I_j.\]  (A6b)

We know apply the theorem of Kennan [19]. The vector-valued function \(H(I)\) satisfies 's quasi-increasing condition because for every \(i\) and \(j \neq i\),

\[
\frac{\partial H_i}{\partial I_j} \geq 0.
\]

(A7)

Next, the vector valued function \(H(I) - I\) is strictly radially quasiconcave. Suppose there exists a solution \(I^*\) to Eq. (A6). Let \(k \in (0, 1)\).

\[
H_i(kI^*) - kI_i^* = \frac{\sigma_i}{\gamma} k \sum_j \beta_{ji} I_j^* - kI_i^* = k \left( \frac{\sigma_i}{\gamma} \sum_j \beta_{ji} I_j^* - I_i^* \right)
\]

(A8)

\[
= k \left( \frac{\sigma_i}{\gamma} \sum_j \beta_{ji} I_j^* - I_i^* \right)
\]

(A9)

By inspection,

\[
\frac{\sigma_i}{\gamma} \sum_j \beta_{ji} I_j^* - I_i^*
\]

(A10)

vanishes at \(k = 1\) and is decreasing in \(k\). Thus, for all \(k \in (0, 1)\) and any \(i\),

\[
\frac{\sigma_i}{\gamma} \sum_j \beta_{ji} I_j^* - I_i^* > 0,
\]

(A11)

\[
k \left( \frac{\sigma_i}{\gamma} \sum_j \beta_{ji} I_j^* - I_i^* \right) > 0.
\]

(A12)

And so, \(H(I) - I\) is strictly radially quasiconcave. By Corollary 1 of [19], there is no more than one positive solution.

Near \(I = 0\),

\[
H(I) \approx \begin{bmatrix}
\frac{\sigma_1}{\gamma} \frac{\beta_{11}}{\beta_{12}} & \frac{\sigma_2}{\gamma} \frac{\beta_{21}}{\beta_{22}} \\
\frac{\sigma_1}{\gamma} \frac{\beta_{12}}{\beta_{11}} & \frac{\sigma_2}{\gamma} \frac{\beta_{22}}{\beta_{21}}
\end{bmatrix} I.
\]

(A13)
The matrix in this equation is positive, so it has a positive eigenvalue (namely $R$) with a corresponding positive eigenvector. Thus, if $R > 1$, then $H(I) - I$ is initially strictly increasing for $I$ lying in this eigenspace. We can also see from inspection that for all positive $I$,

$$H_i(I) < \frac{\sigma_i}{\gamma}$$  \hspace{1cm} (A14)

By Tarski’s theorem [19], there exists one or more positive fixed points.

Thus, $R > 1$ implies there exists a unique positive fixed point. If $R < 1$, then $H(I)$ is bounded above by it’s linearization in Eq. (A13) which is a contraction to the origin (since both eigenvalues have modulus less than 1), so the only fixed point is $I = (0, 0)$.

Hethcote and Yorke [14] show that the unique positive stationary solution is globally attracting when $R > 1$ and the disease-free stationary solution is globally attracting when $R \leq 1$. We now show that more self-protection always decreases disease prevalence. Although this seems intuitive, I am not aware of an earlier proof. The identification of game equilibria depends critically on this result.

**Theorem A.2**: If $R(\sigma) > 1$, then the unique positive stationary disease prevalences $I_1^*, I_2^*$ are strictly increasing in both $\sigma_1$ and $\sigma_2$.

**Proof**: From Theorem A.1, we know that the vector-valued function

$$I^*(\sigma) = (I_1^*(\sigma_1, \sigma_2), I_2^*(\sigma_1, \sigma_2))$$  \hspace{1cm} (A15)

that maps resident strategies to the disease prevalences of the globally attracting stationary solution is a well-defined function. $R(\sigma) > 1$ implies the stationary solution $I_1^*(\sigma) > 0$ and $I_2^*(\sigma) > 0$. The inverse relation

$$\sigma_i = \gamma I_i^* \left(1 + \frac{1}{\sum_j \beta_{ij} I_j^*}\right)$$  \hspace{1cm} (A16)

derived from Eq. (A6) is a smooth function for positive $I^*$. The derivative only becomes unbounded as $R$ approaches 1, corresponding to a loss of uniqueness. Thus, $I^*(\sigma)$ is a bijective function on the domain

$$\{(\sigma_1, \sigma_2) : \sigma_i \in [0, 1] \text{ and } R(\sigma_1, \sigma_2) > 1\}$$  \hspace{1cm} (A17)

If follows from the bijective property that the Jacobian matrix $\partial I^*/\partial \sigma$ exists within the domain and is non-singular. When we differentiate Eq. (A16), we find that the Jacobian matrix solves

$$\frac{\partial \sigma}{\partial I} = \gamma \left[\begin{array}{cc} 1 + \frac{1}{\lambda_1} & 0 \\ 0 & 1 + \frac{1}{\lambda_2} \end{array}\right] - \gamma \left[\begin{array}{ccc} 1 & \beta_{12} \gamma & I_2^* \gamma \\ \beta_{12} I_1^* \gamma & (\lambda_1)^2 & I_1^* \gamma \\ I_1^* \gamma & (\lambda_2)^2 & I_2^* \gamma \end{array}\right]$$  \hspace{1cm} (A18)

$$= \gamma \left[\begin{array}{cc} 1 & 0 \\ 0 & 1 \end{array}\right] + \gamma \left[\begin{array}{ccc} \frac{1}{\lambda_1} & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{array}\right]$$  \hspace{1cm} (A19)

The eigenvalues are $\gamma$ and $\gamma(1 + \frac{I_2^* \beta_{12} \gamma}{(\lambda_1)^2} + \frac{I_1^* \beta_{12} \gamma}{(\lambda_2)^2})$ which are both positive, so the Jacobian matrix $\frac{\partial \sigma}{\partial I}$ is an M-matrix. [15, p. 114]. From M-matrix theory, it imme-
Immediately follows that $\frac{\partial I^*}{\partial \sigma}$ is non-negative. The extra condition that $\beta_{ij} > 0$ for all $ij$ ensures $\frac{\partial \sigma}{\partial I^*}$ is irreducible and $\frac{\partial I^*}{\partial \sigma}$ is positive. We conclude the unique endemic stationary solution is strictly increasing in the population’s resident strategies $\sigma$.

\section*{Appendix B. On linear-fractional forms of the utility function}

One result that follows for the form of utility function derived in Section 2 is that linearity in a strategy parameter $\sigma$ implies that the utility function is a linear-fractional transform, a.k.a. Mobius transform, in said strategy parameter.

\textbf{Theorem B.1:} Suppose $\sigma$ is a component of the strategy vector $\pi$ appearing in exactly one row of the augmented matrix $[(hI - Q^*)^T v]$ and that that row is an affine function of $\sigma$. Also, assume that the initial state $p(0)$ is independent of $\sigma$. Then the utility function has the Mobius transform form

$$U(\sigma) = \frac{a\sigma + b}{d\sigma + e}$$

(B1)

and

$$\frac{\partial U(\sigma)}{\partial \sigma} = \frac{ae - bd}{(d\sigma + e)^2}.$$  

(B2)

where $a, b, d$, and $e$ are independent of $\sigma$.

\textbf{Proof:} For a square $n \times n$ matrix $A$,

$$\det A = \sum_{\rho} \text{sign}(\rho) \prod_j A_{\rho(j), j}$$

(B3)

where each $\rho$ is a different permutation of the integers $1 \ldots n$. Since permutations are bijections, components from each row appear exactly once in each monomial term of the determinant. Thus, if $\sigma$ occurs only in one row of $A$, and occurs linearly in that row, then $\det A$ must be linear in $\sigma$ also.

Now, let $w$ be a solution of $(hI - Q^*)^T w = v$. We will use Cramer’s rule to solve this. For every $k$,

$$w_k = \frac{\det M_k}{\det hI - Q^*}$$

(B4)

where $M_k$ is the same as $(hI - Q^*)^T$ accept that the $k$th column has been replaced by $v$. From our rule for determinants, both the numerator and denominator must be linear in $\sigma$, making $w_k$ a linear-fractional transform in $\sigma$. Since $U = w^T p(0)$ and $p(0)$ is independent of $\sigma$, then $U$ is a sum of linear-fractional transforms with the same denominator, so $U$ itself is a linear-fractional transform in $\sigma$. Eq. (B2) is obtained by direct differentiation.

This may raise several interesting mathematical ideas for the reader, which, alas, we do not have space to address. Obvious cases where Theorem B.1 will fail are cases those individuals have imperfect knowledge about their own state so strategies appear in multiple columns of $Q^*$ and cases where payoffs are nonlinear functions of strategy.