Converging towards the optimal path to extinction

Ira B. Schwartz, Eric Forgoston, Simone Bianco, and Leah B. Shaw

Nonlinear Systems Dynamics Section, Plasma Physics Division, U.S. Naval Research Laboratory, Code 6792, Washington, DC 20375, USA
Department of Mathematical Sciences, 1 Normal Avenue, Montclair State University, Montclair, NJ 07043, USA
Department of Bioengineering and Therapeutic Sciences, University of California San Francisco, 1700 4th Street, San Francisco, CA 94158-2330, USA
Department of Applied Science, The College of William & Mary, P.O. Box 8795, Williamsburg, VA 23187-8795, USA

Extinction appears ubiquitously in many fields, including chemical reactions, population biology, evolution, and epidemiology. Even though extinction as a random process is a rare event, its occurrence is observed in large finite populations. Extinction occurs when fluctuations due to random transitions act as an effective force which drives one or more components or species to vanish. Although there are many random paths to an extinct state, there is an optimal path that maximizes the probability to extinction. In this article, we show that the optimal path is associated with the dynamical systems idea of having maximum sensitive dependence to initial conditions. Using the equivalence between the sensitive dependence and the path to extinction, we show that the dynamical systems picture of extinction evolves naturally toward the optimal path in several stochastic models of epidemics.

Keywords: Extinction, Optimal Path, Finite-Time Lyapunov Exponents

1. Introduction

Determining the conditions for epidemic extinction is an important public health problem. Global eradication of an infectious disease has rarely been achieved, but it continues to be a public health goal for polio (World Health Organization, 2009) and many other diseases, including childhood diseases. More commonly, disease extinction, or fade out, is local and may be followed by a reintroduction of the disease from other regions (Anderson & May, 1991; Grassly et al., 2005). Extinction may also occur for individual strains of a multistrain disease (Minayev & Ferguson, 2009), such as influenza or dengue fever. Since extinction occurs in finite populations, it depends critically on local community size (Bartlett, 1957, 1960; Keeling & Grenfell, 1997; Conlan & Grenfell, 2007). Moreover, it is important to know how parameters affect the chance of extinction for predicting the dynamics of outbreaks and for developing control strategies to promote epidemic extinction (Dykman et al., 2008).

The determination of extinction risk is also of interest in related fields, such as evolution and ecology. For example, in the neutral theory of ecology, bio-diversity arises...
from the interplay between the introduction and extinction of species (Banavar & Maritan, 2009; Azaele et al., 2006).

In general, extinction occurs in discrete, finite populations undergoing stochastic effects due to random transitions or perturbations. The origins of stochasticity may be internal to the system or may arise from the external environment. Small population size, low contact frequency for frequency-dependent transmission, competition for resources, and evolutionary pressure (de Castro & Bolker, 2005), as well as heterogeneity in populations and transmission (Lloyd et al., 2007), may all be determining factors for extinction to occur.

Extinction risk is affected by the nature and strength of the noise (Melbourne & Hastings, 2008), as well as other factors, including outbreak amplitude (Alonso et al., 2006) and seasonal phase occurrence (Stone et al., 2007). For large populations, the intensity of internal population noise is generally small. However, a rare, large fluctuation can occur with non-zero probability and the system may be able to reach the extinct state. For respiratory diseases such as SARS, super-spreading may account for these large stochastic fluctuations (Lloyd-Smith et al., 2005). Since the extinct state is absorbing due to effective stochastic forces, eventual extinction is guaranteed when there is no source of reintroduction (Bartlett, 1949; Allen & Burgin, 2000; Gardiner, 2004). However, because fade outs are usually rare events in large populations, typical time scales for extinction may be extremely long.

Stochastic population models of finite populations, which include extinction processes, are effectively described using the master equation formalism. Stochastic master equations are commonly used in statistical physics when dealing with chemical reaction processes (Kubo, 1963) and predict probabilities of rare events occurring at certain times. For many problems involving extinction in large populations, if the probability distribution of the population is approximately stationary, the probability of extinction is a function that decreases exponentially with increasing population size. The exponent in this function scales as a deterministic quantity called the action (Kubo et al., 1973). It can be shown that a trajectory that brings the system to extinction is very likely to lie along a most probable path, called the optimal extinction trajectory or optimal path. It is a remarkable property that a deterministic quantity such as the action can predict the probability of extinction, which is inherently a stochastic process (Schwartz et al., 2009; Dykman et al., 2008).

Locating the optimal path is desirable because the quantity of interest, the extinction rate, depends on the probability to traverse this path, and the effect of a control strategy on extinction rate can be determined by its effect on the optimal path (Dykman et al., 2008). By employing an optimal path formalism, we convert the stochastic problem to a mechanistic dynamical systems problem. In contrast to approaches based on diffusive processes that are valid only in the limit of large system sizes (Näsell, 1999; Lindholm, 2008), this dynamical systems approach can give accurate estimates for the extinction time even for small populations if the action is sufficiently large. Additionally, unlike other methods that are used to estimate lifetimes, this approach enables one both to estimate lifetimes and to draw conclusions about the path taken to extinction. This more detailed understanding of how extinction occurs may lead to new stochastic control strategies (Dykman et al., 2008).

In this article, we show that locating the optimal extinction trajectory can be
achieved naturally by evolving a dynamical system that converges to the optimal path. The method is based on computing finite-time Lyapunov exponents (FTLE), which have previously been used to find coherent structures in fluid flows (Haller, 2000, 2001, 2002; Shadden et al., 2005; Lekien et al., 2007; Branicki & Wiggins, 2010). The FTLE provides a measure of how sensitively the system’s future behavior depends on its current state. We argue that the system displays maximum sensitivity near the optimal extinction trajectory, which enables us to dynamically evolve toward the optimal escape trajectory using FTLE calculations. For several models of epidemics that contain internal or external noise sources, we illustrate the power of our method to locate optimal extinction trajectories. Although our examples are taken from infectious disease models, the approach is general and is applicable to any extinction process or escape process.

2. Stochastic modeling in the large population limit

To introduce our idea of dynamically constructing an optimal path to extinction in stochastic systems, we show its application to a stochastic Susceptible-Infectious-Recovered (SIR) epidemiological model. Details of the SIR model can be found in Sec. 1 of the electronic online supplement. Figure 1 shows the probability density of extinction prehistory in the SI plane. The probability density was numerically computed using 20,000 stochastic SIR trajectories that ended in extinction. Trajectories are aligned at their extinction point. From the extinction point, the prehistory of each trajectory up to the last outbreak of infection is considered. Small fluctuations in the infectious population are not considered in identifying the last outbreak. In this way, we restrict the analysis to the interval between the last large outbreak of infection and the extinction point. The resulting (S,I) pairs of susceptible and infectious individuals are then binned and plotted in the SI plane (Eilers & Goeeman, 2004). The resulting discrete density has been color coded so that the brighter regions correspond to higher density of trajectories. The figure shows that, among all the paths that the stochastic system can take to reach the extinct state, there is one path that has the highest probability of occurring. This is the optimal path to extinction. One can see that the optimal path to extinction lies on the peak of the probability density of the extinction prehistory. It should be noted that extinction for the stochastic SIR model has been studied previously (Kamenev & Meerson, 2008).

The optimal path can be obtained using methods of statistical physics. In figure 1, the numerical prediction of the entire optimal trajectory for the stochastic SIR system has been overlaid on the probability density of extinction prehistory that was found using stochastic simulation. The trajectory spirals away from the endemic state, with larger and larger oscillations until it hits the extinct state. The agreement between the stochastically simulated optimal path to extinction and the predicted optimal path is excellent.

The curve of figure 1 is obtained by recasting the stochastic problem in a deterministic form. The evolution of the probability of finding a stochastic system in a given state X at a given time t is described by the master equation (van Kampen, 2007). Solving the master equation would provide a complete description of the time evolution of the stochastic system, but in general it is a difficult task to obtain explicit solutions for the master equation. Thus, one generally resorts to approxi-
mations to the solution; i.e., one considers an ansatz for the probability density. In this case, since extinction of finite populations is a rare event, we will be interested in examining events that occur in the tail of the probability distribution. Therefore, the distribution is assumed to take the form

$$\rho(X, t) \approx \exp \left(-NS(x)\right),$$

where $\rho(X, t)$ is the probability density of finding the system in the state $X$ at time $t$, $N$ is the size of the population, $x = X/N$ is the normalized state (e.g., in an epidemic model, the fraction of the population in the various compartments), and $S$ is a deterministic state function known in classical physics as the action. Equation (2.1) describes the relationship between the action and the probability density and is based on an assumption for how the probability scales with the population size. The action is the negative of the natural log of the stationary probability distribution divided by the population size. Therefore, the probability (if we normalize the population) is roughly given by the exponential of the action. Intuitively, equation (2.1) expresses the assumption that the probability of occurrence of extreme events, such as extinction, lies in the tails of the probability distribution, which falls steeply away from the steady state.

This approximation leads to a conserved quantity that is called the Hamiltonian \cite{Gang1987}. From the Hamiltonian, one can find a set of conservative ordinary differential equations (ODEs) that are known as Hamilton’s equations. These ODEs describe the time evolution of the system in terms of state variables $x$, which are associated with the population in each compartment. For the SIR example, $x$ is the vector $\langle S, I, R \rangle$. In addition to the state variables, the equations contain conjugate momenta variables, $p_x$. The conjugate momenta, or noise, account for the uncertainty associated with the system being in a given state at a given time due to the stochastic interactions among the individuals of the population. These ODEs can be constructed from information in the master equation about the possible transitions and transition rates in the system. Details can be found in \cite{Appendix A}.

Integration of the ODEs with the appropriate boundary conditions will then give the optimal evolution of the system under the influence of the noise. Boundary conditions are chosen to be fixed points of the system. A typical case is shown schematically in figure 2. Deterministically, the endemic state is attracting and the extinct state repelling. However, introducing stochasticity allows the system to leave the deterministic manifold along an unstable direction of the endemic state, corresponding to nonzero noise. Stochasticity leads to an additional extinct state which arises due to the general non-Gaussian nature of the noise. For the extinction process of figure 1, boundary conditions were the system leaving the endemic steady state and asymptotically approaching the stochastic extinct state.

In general, the optimal extinction path is an unstable dynamical object, and this reflects extinction as a rare event. This has led many authors to consider how extinction rates scale with respect to a parameter close to a bifurcation point \cite{Dorner2005, Kamenev2008, Kamenev2008, Dykman2008}, where the dynamics are very slow. For an epidemic model this means that the reproductive rate $R_0$ should be greater than but very close to 1. However, most real diseases have $R_0$ larger than 1.5, which translates into a faster growth rate from the extinct state. In general, in order to obtain analytic scaling results, one must obtain the ODEs for the optimal path either analytically (using the classical
theory of large fluctuations mentioned within this section and described in detail in Appendix A or numerically (using shooting methods for boundary value problems). This task may be impossible or extremely cumbersome, especially when the system is far from the bifurcation point. In the following section we demonstrate how to evolve naturally to the optimal path to extinction using a dynamical systems approach.

3. Finite-time Lyapunov exponents

We consider a velocity field which is defined over a finite time interval and is given by Hamilton’s equations of motion. Such a velocity field, when starting from an initial condition, has a unique solution. The continuous dynamical system has quantities, known as Lyapunov exponents, which are associated with the trajectory of the system in an infinite time limit, and which measure the average growth rates of the linearized dynamics about the trajectory. To find the finite-time Lyapunov exponents (FTLE), one computes the Lyapunov exponents on a restricted finite time interval. For each initial condition, the exponents provide a measure of its sensitivity to small perturbations. Therefore, the FTLE is a measure of the local sensitivity to initial data. Details regarding the FTLE can be found in Sec. 2 of the electronic online supplement.

The FTLE measurements can be shown to exhibit “ridges” of local maxima. The ridges of the field indicate the location of attracting (backward time FTLE field) and repelling (forward time FTLE field) structures. In two-dimensional (2D) space, the ridge is a curve which locally maximizes the FTLE field so that transverse to the ridge one finds the FTLE to be a local maximum. What is remarkable is that the FTLE ridges correspond to the optimal path trajectories, which we heuristically argue in Sec. 3 of the electronic online supplement. The basic idea is that since the optimal path is inherently unstable, the FTLE shows that, locally, the path is also the most sensitive to initial data. Figure 2: shows a schematic that demonstrates why the optimal path has a local maximum to sensitivity. If one chooses an initial point on either side of the path near the endemic state, the two trajectories will separate exponentially in time. This is due to the fact that both extinct and endemic states are unstable, and the connecting trajectory defining the path is unstable as well. Any initial points starting near the optimal path will leave the neighborhood in short time.

4. Evolving towards the optimal path using FTLE

We now apply our theory of dynamical sensitivity to the problem of locating optimal paths to extinction for several examples. We consider the case of internal fluctuations, where the noise is not known a priori, as well as the case of external noise. In each case, the interaction of the noise and state of the system begins by finding the equations of motion that describe the unstable flow. These equations of motion are then used to compute the ridges corresponding to maximum FTLE, which in turn correspond to the optimal extinction paths (Forgoston et al. 2010).
(a) Extinction in a Branching - Annihilation Process

For an example of a system with internal fluctuations which has an analytical solution, consider extinction in the stochastic branching-annihilation process

\[ A \xrightarrow{\lambda} 2A \quad \text{and} \quad 2A \xrightarrow{\mu} \emptyset, \]

where \( \lambda \) and \( \mu > 0 \) are constant reaction rates (Elgart & Kamenev, 2004; Assaf et al., 2008). Equation (4.1) is a single species birth-death process and can be thought of as a simplified form of the Verhulst logistic model for population growth (N˚ asell, 2001). The mean field equation for the average number of individuals \( n \) in the infinite population limit is given by \( \dot{n} = \lambda n - \mu n^2 \). The stochastic process given by equation (4.1) contains intrinsic noise which arises from the randomness of the reactions and the fact that the population consists of discrete individuals. This intrinsic noise can generate a rare sequence of events that causes the system to evolve to the extinct state. The probability \( P_n(t) \) to observe, at time \( t \), \( n \) individuals is governed by the master equation

\[ \dot{P}_n = \frac{\mu}{2} [(n + 2)(n + 1)P_{n+2} - n(n - 1)P_n] + \lambda [(n - 1)P_{n-1} - nP_n]. \]

The Hamiltonian associated with this system is

\[ H(q, p) = \left( \lambda (1 + p) - \frac{\mu}{2} (2 + p)q \right) qp, \]

where \( q \) is a conjugate coordinate related to \( n \) through a transformation (Assaf et al., 2008), and \( p \) plays the role of the momentum. The equations of motion are given by

\[ \dot{q} = \frac{\partial H}{\partial p} = q[\lambda(1 + 2p) - \mu(1 + p)q], \quad (4.4a) \]

\[ \dot{p} = -\frac{\partial H}{\partial q} = p[\mu(2 + p)q - \lambda(1 + p)]. \quad (4.4b) \]

The mean field is retrieved in equation (4.4a) when \( p = 0 \) (no fluctuations or noise). The Hamiltonian has three zero-energy curves. The first is the mean-field zero-energy line \( p = 0 \) (no fluctuations), which contains two unstable points \( h_1 = (p,q) = (0, \lambda/\mu) \) and \( h_0 = (p,q) = (0,0) \). The second is the extinction line \( q = 0 \) (trivial solution), which contains another unstable point \( h_2 = (p,q) = (-1,0) \). The third zero-energy curve is non-trivial and is

\[ q = \frac{2\lambda(1 + p)}{\mu(2 + p)} \]

The segment of the curve given by equation (4.5) which lies between \(-1 \leq p \leq 0\) corresponds to a (heteroclinic) trajectory which exits, at \( t = -\infty \), the point \( h_1 \) along its unstable manifold and enters, at \( t = \infty \), the point \( h_2 \) along its stable manifold. This trajectory is the optimal path to extinction and describes the most probable sequence of events which evolves the system from a quasi-stationary state to extinction (Assaf et al., 2008).
To show that the FTLE evolves to the optimal path, we calculate the FTLE field using the system of Hamilton’s equations given by equations (4.4a)-(4.4b). Figure 3 shows both the forward and backward FTLE plot computed for the finite time $T = 6$, with $\lambda = 2.0$ and $\mu = 0.5$. In this example, as well as the following two examples, $T$ was chosen to be sufficiently large so that one obtains a measurable exponential separation of trajectories. In figure 3a, one can see that the optimal path to extinction is given by the ridge associated with the maximum FTLE. In fact, by overlaying the forward and backward FTLE fields, one can see all three zero-energy curves including the optimal path to extinction. Also shown in figure 3a are the analytical solutions to the three zero-energy curves given by $p = 0, q = 0$, and equation (4.5). There is excellent agreement between the analytical solutions of all three curves and the ridges which are found through numerical computation of the FTLE flow fields.

It is possible to compute analytically the action along the optimal path for a range of $\lambda/\mu$ values. Using equation (4.5), it is easy to show that the action $S$ is

$$S = 2(1 - \ln 2) \frac{\lambda}{\mu}. \tag{4.6}$$

It is clear from equation (4.6) that the action scales linearly with $\lambda/\mu$.

(b) SIS Epidemic Model - External Fluctuations

We now consider the well-known problem of extinction in a Susceptible-Infectious-Susceptible (SIS) epidemiological model, which is a core model for the basis of many recurrent epidemics. The SIS model is described by the following system of equations:

$$\dot{S} = \mu - \mu S + \gamma I - \beta IS, \tag{4.7a}$$
$$\dot{I} = - (\mu + \gamma) I + \beta IS. \tag{4.7b}$$

where $\mu$ denotes a constant birth and death rate, $\beta$ represents the contact rate, and $\gamma$ denotes the recovery rate. Assuming the total population size is constant and can be normalized to $S + I = 1$, then equations (4.7a)-(4.7b) can be rewritten as the the following one-dimensional (1D) process for the fraction of infectious individuals in the population:

$$\dot{I} = - (\mu + \gamma) I + \beta I(1 - I). \tag{4.8}$$

The stochastic version of equation (4.8) is given as

$$\dot{I} = -(\mu + \gamma) I + \beta I(1 - I) + \sigma \eta(t) = F(I) + \sigma \eta(t), \tag{4.9}$$

where $\eta(t)$ is uncorrelated Gaussian noise with zero mean and $\sigma$ is the standard deviation of the noise intensity. The noise models random migration to and from another population (Alonso et al., 2006; Doering et al., 2005).

Equation (4.9) has two equilibrium points given by $I = 0$ (disease-free state) and $I = 1 - (\mu + \gamma)/\beta$ (endemic state). Using the Euler-Lagrange equation of motion (Goldstein et al., 2001) from the Lagrangian determined by equation (4.9) ($L(I, \dot{I}) = [\eta(t)]^2 = [I - F(I)]^2$) along with the boundary conditions given by the extinct and endemic states, one finds that the optimal path to extinction (as well

*Article submitted to Royal Society*
as its counterpart path from the disease-free state to the endemic state) is given by \( \dot{I} = \pm F(I) \).

As in the first example, one can numerically compute the optimal path to extinction using the FTLE. Figure 3b shows the forward and backward FTLE plot computed for \( T = 5 \), with \( \beta = 5.0 \) and \( \kappa = \mu + \gamma = 1.0 \). Note that we can consider the combination \( \mu + \gamma \) since the Lagrangian depends only on the combination rather than on either parameter separately. In figure 3b, one can see that the optimal path from the endemic state to the disease-free state is given by the ridge associated with the local maximum FTLE. Also shown in figure 3b is the counterpart optimal path from the disease-free state to the endemic state (found by computing the backward FTLE field). In addition, the agreement with the analytical prediction is excellent, as shown in figure 3b.

If one solves equation (4.9) for \( I(t) \) and substitutes both \( I(t) \) and \( \dot{I}(t) \) into the Lagrangian, then one can analytically find an expression for the action along the optimal path. The expression for the action is a function of \( \kappa \) and the reproductive number \( R_0 \) and is given by

\[
S = \frac{2\kappa(R_0 - 1)^3}{3R_0^2}, \tag{4.10}
\]

where \( R_0 = \beta/\kappa \).

(c) SIS Epidemic Model - Internal Fluctuations

We next consider the 1D stochastic version of the SIS epidemic model for a finite population with internal fluctuations using the transition rates found in Sec. 4 of the electronic online supplement. Using the formalism of Gang (1987), one then has the following Hamiltonian associated with this model:

\[
H(I, p) = (\mu + \gamma)I(e^{-p} - 1) + \beta I(1 - I)(e^{p} - 1), \tag{4.11}
\]

where \( I \) is the fraction of infectious individuals and \( p \) is the momentum. Internal fluctuations arise from the random interactions of the population. Although there is no analytical solution for the optimal path to extinction, we can once again determine the optimal path by computing the FTLE flow field associated with this system. Figure 4 shows the forward FTLE plot computed using Hamilton’s equation of motion for \( T = 10 \), with \( \beta = 2.0 \) and \( \kappa = \mu + \gamma = 1.0 \), and as in previous examples, the optimal path to extinction from the endemic state to the disease-free state is apparent. Note that the non-zero momentum corresponding to the extinct state qualitatively agrees with similar boundary conditions found in Dykman et al. (2008) and is associated with non-zero probability flux.

Once again, it is possible to compute the action along the optimal path for a range of values of the reproductive number \( R_0 \). In contrast to the prior two examples, here the action must be computed numerically. Moreover, even the optimal path must be found numerically using the FTLE plot generated for each value of \( R_0 \).

Given an \( R_0 \), we computed the forward FTLE flow field using a grid resolution of 0.005 in both position and momentum space. To find the optimal path corresponding to the ridge of maximal FTLE values, we let the deterministic, endemic steady state be the starting point \( z_0 \) of the path. Then a second point \( z_1 \) on the path

*Article submitted to Royal Society*
was found by locating the maximum of the FTLE values in an arc of radius 20 grid points spanning nearly π radians for negative momentum values and originating at \(z_0\). Subsequent points \(z_{n+1}\) were found by taking the maximum FTLE value on an arc of radius 10 grid points spanning nearly π radians originating at the most recent point \(z_n\) and centered around the vector \(z_n - z_{n-1}\) until the extinction line was reached. The complete optimal path was estimated from the \(z_n\) using cubic spline interpolation. Once the optimal path had been found, the action was computed by numerically integrating the Lagrangian along this path.

Figure 5a shows the numerically computed action versus reproductive number over the range \(1.1 \leq R_0 \leq 20\). The inset of figure 5a shows the portion of figure 5a from \(1.1 \leq R_0 \leq 2\). Also shown in the inset of figure 5a is an analytical, asymptotic scaling result that is valid for values of \(R_0\) close to unity. As can be seen in figure 5a, there is good agreement between the numerically computed action and the analytically computed action near \(R_0 = 1\). Details of the derivation of the analytical scaling law can be found in Sec. 5 of the electronic online supplement.

Figure 5b shows the numerically simulated mean extinction time versus reproductive number for the 1D stochastic SIS model for a finite population (see Sec. 4c). Also shown in figure 5b is the analytical prediction found using the previously mentioned scaling law derived for values of \(R_0\) close to unity. As one can see, the agreement is excellent.

### 5. Conclusions

In all of the examples of Sec. 4, we have shown that the optimal path to extinction is equated with having a (locally) maximal sensitivity to initial conditions. Even though there exist many possible paths to extinction, the dynamical systems approach converges to the path that maximizes extinction. The parameter values chosen for the three examples are such that the extinct and endemic states are far away from each other. Therefore, in general, there will be no possible approximate analytical treatment as performed in Dykman et al. (2008). In addition, we have shown how to constructively compute numerically the action for a wide range of reproductive numbers. Our method allows for the computation of extinction times and can be extended to high-dimensional problems.

Because the method is general, and unifies dynamical systems theory with the probability of extinction, we expect that any system found in other fields can be understood using this approach. Indeed, in problems of general extinction, it is now possible to evolve naturally to the optimal path, and thus predict the path that maximizes the probability to extinction. Future work in this area will include improved sampling methods to find the optimal path more efficiently in higher dimensional models. Specific applications of optimal path location in the future will include spatio-temporal extinction processes such as those that occur in pre-vaccine measles (Keeling & Grenfell 1997; Finkenstädt et al., 2002) and multi-strain extinction in diseases such as dengue (Cummings et al., 2005).

Finally, the optimal path method may lead to novel monitoring and control strategies. In one biological application, knowledge of the most probable path to extinction may help with monitoring an environment and with providing an estimate of the likelihood of extinction based on where the population lies relative to the path. It is known that once a trajectory has left the neighborhood of the
endemic state, most paths to extinction occur near the optimal path. This phenomenon can be seen in figure [1] where the optimal path lies on the peak of the probability density of extinction prehistory. Therefore, the optimal path provides a good location to monitor the system for possible extinction events. Furthermore, although the momenta (noise effects) are not directly observable, it may be possible to infer them dynamically (Luchinsky et al., 2008) from time series data of observable quantities in conjunction with the equations for the time evolution of position and momentum variables. Knowledge of where a system lies in position-momentum space could provide an estimate for how quickly a desired epidemic extinction could occur or could provide the risk of extinction for a species one wishes to conserve.

In yet another application, knowledge of the optimal path to extinction has the advantage that, in the presence of noise (that is estimated from data) and a known population of infectious individuals, it may be possible to develop better vaccine controls that reduce the time to extinction. Figure 2(a) shows a schematic of the path to extinction, where the extinct state is a saddle with stable and unstable directions. For many epidemic models, the extinct state can be shown to have a similar geometry of stable and unstable directions. An approach to the extinct state on the optimal path will lead to the fastest time to extinction. Moreover, since the extinct state has a saddle structure in the presence of noise, it may be controlled with projection methods (Schwartz & Triandaf, 1994) or probabilistic techniques (Schwartz et al., 2004).

One can consider instituting a method of parameter control, where the parameters could be vaccine levels or treatment of infectious individuals. Combined with the above-mentioned monitoring techniques, the control method will allow one to move an existing state that deviates from the optimal path closer to the optimal path as time evolves. By adjusting the parameters, we may target the stable directions of the path when we are close to epidemic extinction (Schwartz & Triandaf, 1994). Comparing observations with model predictions of the optimal path allow us to use controls to minimize the time to extinction.

The authors gratefully acknowledge support from the Office of Naval Research, the Air Force Office of Scientific Research, the Army Research Office, and the National Institutes of Health. I.B.S and L.B.S. are supported by Award Number R01GM090204 from the National Institute Of General Medical Sciences. The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institute Of General Medical Sciences or the National Institutes of Health. We also gratefully acknowledge M. Dykman for helpful discussions.

**Appendix A. Theory of large fluctuations**

Letting $N$ denote the population size, the state variables $X \in \mathbb{R}^n$ of the system describe the components of a population, while the random state transitions which govern the dynamics are described by the transition rates $W(X, r)$, where $r \in \mathbb{R}^n$ is an increment in the change of $X$. In the large population limit without any fluctuations, the mean field equations are given by the system $\dot{X} = \sum_r rW(X, r)$.

Consideration of the net change in increments of the state of the system results in the following master equation for the probability density $\rho(X, t)$ of finding the...
system in state $\mathbf{X}$ at time $t$:

$$
\frac{\partial \rho(\mathbf{X}, t)}{\partial t} = \sum_r \left[ W(\mathbf{X} - \mathbf{r}; \mathbf{r}) \rho(\mathbf{X} - \mathbf{r}, t) - W(\mathbf{X}; \mathbf{r}) \rho(\mathbf{X}, t) \right].
$$

(A 1)

The solution to the master equation (A 1) has a peak around a stable steady state in the limit of large population ($N \gg 1$) with width $\propto N^{1/2}$ (Dykman et al., 2008; Schwartz et al., 2009). However, since we are interested in the probability of extinction, we will consider the tail of the distribution, which gives the probability of having a small number of individuals. The tail of the distribution can be obtained by employing the ansatz given by equation (2.1), which is an assumption for how the probability density scales with population size (Kubo, 1973; Dykman, 1990). Equation (2.1) also implies that a maximum in the extinction probability can be found minimizing the action over a set of extinction paths starting from the stable endemic state. The assumption of this functional form for $\rho$ allows the action to be derived from properties of the master equation.

The density, $\rho(\mathbf{X}, t)$, can be found by substituting the ansatz given by equation (2.1) into the master equation. The resulting equations for the action are given by the Hamilton-Jacobi equation for a Hamiltonian, $H$, given by $\partial S/\partial t + H(\mathbf{x}, \partial S/\partial \mathbf{x}; t) = 0$, where we have normalized the population components and rates respectively as $\mathbf{x} = \mathbf{X}/N, w(\mathbf{x}, \mathbf{r}) = W(\mathbf{X}, \mathbf{r})/N$.

Following the classical mechanics convention, define a conjugate momentum to the state space, $\mathbf{p}$, by letting $p = \partial S/\partial \mathbf{x}$ and where $H(\mathbf{x}, \mathbf{p}; t)$ is the classical Hamiltonian (Kubo et al., 1973). The Hamiltonian, $H$, depends both on the state of the system, $\mathbf{x}$, as well as the momentum, $\mathbf{p}$, which provides an effective force due to stochastic fluctuations on the system. The Hamiltonian equations of motion provide trajectories in time of $\mathbf{x}(t)$ and $\mathbf{p}(t)$, and as such, describe a set of paths going from an initial point at time $t_i$ to some final point at time $t_f$ in $(\mathbf{x}, \mathbf{p})$ space. For a given path, the action is given by $S = \int_{t_i}^{t_f} \mathbf{p}(t) \dot{\mathbf{x}}(t) dt$, and as such determines the exponent of the probability of observing that path. (It should be noted that instead of using the Hamiltonian representation, one could use the Lagrangian representation $L(\mathbf{x}, \dot{\mathbf{x}}; t) = -H(\mathbf{x}, \mathbf{p}; t) + \dot{\mathbf{x}} \cdot \mathbf{p}$, which results in an equivalent solution.)

The action reveals much information about the probability evolution of the system, from scaling near bifurcation points in non-Gaussian processes to rates of extinction as a function of epidemiological parameters (Dykman, 1990; Dykman et al., 2008). As already stated, in order to maximize the probability of extinction, one must minimize the action $S$. The minimizing formulation entails finding the solution to the Hamilton-Jacobi equation, which means that one must solve the $2n$-dimensional system of Hamilton’s equations $[\dot{\mathbf{x}} = \partial_p H(\mathbf{x}, \mathbf{p}), \dot{\mathbf{p}} = -\partial_x H(\mathbf{x}, \mathbf{p})]$ for $\mathbf{x}$ and $\mathbf{p}$, where the Hamiltonian is explicitly given as

$$
H(\mathbf{x}, \mathbf{p}, t) = \sum_r w(\mathbf{x}, \mathbf{r}) [\exp (\mathbf{p}^T \mathbf{r}) - 1].
$$

(A 2)

The appropriate boundary conditions of the system are such that a solution starts at a non-zero state, such as an endemic state, and asymptotically approaches one or more zero components of the state vector, representing a disease free state. Therefore, a trajectory that is a solution to the two-point boundary value problem determines a path, which in turn yields the probability of going from the initial
state to the final state. The optimal path to extinction is the path which minimizes the action in either the Hamiltonian or Lagrangian representation.

We compute the trajectory satisfying the Hamiltonian system that has as its asymptotic limits in time the endemic state as $t \to -\infty$ and the extinct state as $t \to +\infty$. The momentum $p$ represents the force of the fluctuations on the population, and this momentum changes the stability of the equilibrium points. Both the endemic and extinct states have attracting and repelling directions for $p \neq 0$, as shown schematically in Figure 2.

Given optimal path trajectories $(x(t), p(t))$, the action with correct limits is found by $S(x) = \int_{-\infty}^{\infty} p \dot{x} \, dt$ \cite{Goldstein2001}.

References

Allen, L. J. S. & Burgin, A. M. 2000 Comparison of deterministic and stochastic SIS and SIR models in discrete time. Math. Biosci. 163, 1–33.

Alonso, D., McKane, A. J. & Pascual, M. 2006 Stochastic amplification in epidemics. J. R. Soc. Interface 4, 575–582.

Anderson, R. M. & May, R. M. 1991 Infectious diseases of humans. Oxford University Press.

Assaf, M., Kamenev, A. & Meerson, B. 2008 Population extinction in a time-modulated environment. Phys. Rev. E 78, 041123.

Azaele, S., Pigolotti, S., Banavar, J. R. & Maritan, A. 2006 Dynamical evolution of ecosystems. Nature 444, 926–928.

Banavar, J. R. & Maritan, A. 2009 Ecology: Towards a theory of biodiversity. Nature 460, 334–335.

Bartlett, M. S. 1949 Some evolutionary stochastic processes. J. Roy. Stat. Soc. B Met. 11, 211–229.

Bartlett, M. S. 1957 Measles periodicity and community size. J. R. Stat. Soc. Ser. A-G 120, 37–44.

Bartlett, M. S. 1960 The critical community size for measles in the United States. J. R. Stat. Soc. Ser. A-G 123, 37–44.

Branicki, M. & Wiggins, S. 2010 Finite-time Lagrangian transport analysis: Stable and unstable manifolds of hyperbolic trajectories and finite-time Lyapunov exponents. Nonlinear Proc. Geoph., 17, 1–36.

Conlan, A. J. K. & Grenfell, B. T. 2007 Seasonality and the persistence and invasion of measles. P. Roy. Soc. B-Biol. Sci. 274, 1133–1141.

Cummings, D. A. T., Schwartz, I. B., Billings, L., Shaw, L. B. & Burke, D. S. 2005 Dynamic effects of antibody-dependent enhancement on the fitness of viruses. Proc. Natl. Acad. Sci. U.S.A. 102(42), 15259–15264.

de Castro, F. & Bolker, B. 2005 Mechanisms of disease-induced extinction. Ecol. Lett. 8, 117–126.
Doering, C. R., Sargsyan, K. V. & Sander, L. M. 2005 Extinction times for birth-death processes: Exact results, continuum asymptotics, and the failure of the Fokker-Planck approximation. *Multiscale Model. Simul.* 3(2), 283–299.

Dykman, M. I. 1990 Large fluctuations and fluctuational transitions in systems driven by coloured gaussian noise: A high-frequency noise. *Phys. Rev. A* 42, 2020–2029.

Dykman, M. I., Schwartz, I. B. & Landsman, A. S. 2008 Disease extinction in the presence of random vaccination. *Phys. Rev. Lett.* 101, 078101.

Eilers, P. H. C. & Goeman, J. J. 2004 Enhancing scatterplots with smoothed densities. *Bioinformatics* 20(5), 623–628.

Elgart, V. & Kamenev, A. 2004 Rare event statistics in reaction-diffusion systems. *Phys. Rev. E* 70, 041106.

Finkenstädt, B. F., Bjørnstad, O. N. & Grenfell, B. T. 2002 A stochastic model for extinction and recurrence of epidemics: estimation and inference for measles outbreaks. *Biostatistics* 3, 493–510.

Forgoston, E., Bianco, S., Shaw, L. B. & Schwartz, I. B. 2011 Maximal sensitive dependence and the optimal path to epidemic extinction. *B. Math. Biol.* 73(3), 495-514.

Gang, H. 1987 Stationary solution of master equations in the large-system-size limit. *Phys. Rev. A* 36(12), 5782–5790.

Gardiner, C. W. 2004 *Handbook of stochastic methods for physics, chemistry and the natural sciences*. Springer-Verlag.

Goldstein, H., Poole, C. P., & Safko, J. L. 2001 Classical Mechanics (3rd Ed.) Addison Wesley.

Grassly, N. C., Fraser, C. & Garnett, G. P. 2005 Host immunity and synchronized epidemics of syphilis across the United States. *Nature* 433, 417–421.

Haller, G. 2000 Finding finite-time invariant manifolds in two-dimensional velocity fields. *Chaos* 10(1), 99–108.

Haller, G. 2001 Distinguished material surfaces and coherent structures in three-dimensional fluid flows. *Physica D* 149, 248–277.

Haller, G. 2002 Lagrangian coherent structures from approximate velocity data. *Phys. Fluids* 14(6), 1851–1861.

Kamenev, A. & Meerson, B. 2008 Extinction of an infectious disease: A large fluctuation in a nonequilibrium system. *Phys. Rev. E* 77, 061107.

Kamenev, A., Meerson, B. & Shklovskii, B. 2008 How colored environmental noise affects population extinction. *Phys. Rev. Lett.* 101(26), 268103.

Keeling, M. J. & Grenfell, B. T. 1997 Disease extinction and community size: Modeling the persistence of measles. *Science* 275, 65–67.
Kubo, R. 1963 Stochastic Liouville equations. *J. Math. Phys.* **4**, 174–183.

Kubo, R., Matsuo, K. & Kitahara, K. 1973 Fluctuation and relaxation of macrovariables. *J. Stat. Phys.* **9**(1), 51–96.

Lekien, F., Shadden, S. C. & Marsden, J. E. 2007 Lagrangian coherent structures in n-dimensional systems. *J. Math. Phys.* **48**, 065404.

Lindholm, M. 2008 On the time to extinction for a two-type version of Bartlett’s epidemic model. *Math. Biosci.* **212**, 99–108.

Lloyd, A. L., Zhang, J. & Root, A. M. 2007 Stochasticity and heterogeneity in host-vector models. *J. R. Soc. Interface* **4**, 851–863.

Lloyd-Smith, J. O., Schreiber, S. J., Kopp, P. E. & Getz, W. M. 2005 Superspreading and the effect of individual variation on disease emergence. *Nature* **438**, 355–359.

Luchinsky, D. G., Smelyanskiy, V. N., Millonas, M. & McClintock, P. V. E. 2008 Dynamical inference of hidden biological populations. *Eur. Phys. J. B*, **65**(3), 369–377.

Melbourne, B. A. & Hastings, A. 2008 Extinction risk depends strongly on factors contributing to stochasticity. *Nature* **454**, 100–103.

Minayev, P. & Ferguson, N. 2009 Incorporating demographic stochasticity into multi-strain epidemic models: application to influenza A. *J. R. Soc. Interface* **6**, 989–996.

Näsell, I. 1999 On the time to extinction in recurrent epidemics. *J. R. Statist. Soc. B* **61**, 309–330.

Näsell, I. 2001 Extinction and quasi-stationarity in the Verhulst logistic model. *J. Theor. Biol.* **211**, 11–27.

Schwartz, I. B. & Triandaf, I. 1994 Control and Prediction in Seasonally Driven Population Models. In *Predictability and Nonlinear Modelling in Natural Sciences and Economics* (ed. Grasman, J. and van Straten, G.), pp. 216–227. Kluwer Academic.

Schwartz, I. B., Carr, T. W. & Triandaf, I. 1997 Tracking controlled chaos: Theoretical foundations and applications. *Chaos*, **7**(4), 664–679.

Schwartz, I. B., Billings, L. & Boltt, E. M. 2004 Dynamical epidemic suppression using stochastic prediction and control. *Phys. Rev. E*, **70**, 046 220.

Schwartz, I. B., Billings, L., Dykman, M. & Landsman, A. 2009 Predicting extinction rates in stochastic epidemic models. *J. Stat. Mech.-Theory E*. P01005.

Shadden, S. C., Lekien, F. & Marsden, J. E. 2005 Definition and properties of Lagrangian coherent structures from finite-time Lyapunov exponents in two-dimensional aperiodic flows. *Physica D* **212**, 271–304.

Stone, L., Olinky, R. & Huppert, A. 2007 Seasonal dynamics of recurrent epidemics. *Nature* **446**, 533–536.

*Article submitted to Royal Society*
van Kampen, N. G. 2007 *Stochastic processes in physics and chemistry*. Elsevier.

World Health Organization 2009 Progress towards interrupting wild poliovirus transmission worldwide, 2008. *Weekly Epidemiological Record* **84**, 110–116.
Figure 1. Probability density of extinction prehistory and the optimal path to extinction for the stochastic SIR epidemiological model. Colors indicate the probability density (the colorbar values have been normalized, with lighter colors corresponding to higher probability) for 20,000 stochastic realizations. The results were computed using Monte Carlo simulations, and details of the sampling of the trajectories are described in the text. The curve is the numerically predicted optimal path to extinction. Note that the optimal path to extinction lies on the peak of the probability density of extinction prehistory. The population is $3 \times 10^6$ individuals, with $R_0 \approx 15$ (contact rate $\beta = 1500$, recovery rate $\gamma = 100$, birth-death rate $\mu = 0.2$).

Figure 2. a) Schematic showing the fixed points and heteroclinic trajectories (trajectories connecting fixed points). Coordinate axes are dashed lines. The noise coordinate is indicated by the momentum ($p$) coordinate. The deterministic manifold ($p = 0$) is indicated in blue. Deterministically, the extinct state is repelling and endemic state is attracting. However, the endemic state has unstable directions for nonzero noise ($p \neq 0$), and the optimal path (red) is the trajectory carrying the system from the endemic state to a stochastic extinct state. b) Schematic showing the path from the endemic state (blue) to the extinct state (red). The optimal path leaves the endemic point along an unstable manifold, and connects to the extinct state along its stable manifold. The magenta and green dashed lines represent trajectories initially separated by the optimal path. The initial starting distance between trajectories near the endemic state expands exponentially in forward time (shown by the cyan lines). Locally, this shows that the finite-time Lyapunov measure of sensitivity with respect to initial data is maximal along the optimal path.
Figure 3. (a) Forward and backward FTLE flow fields for the branching-annihilation process computed using the Hamiltonian given by equation (4.3) with \( \lambda = 2.0 \) and \( \mu = 0.5 \). The integration time is \( T = 6 \) with an integration step size of \( t = 0.1 \) and a grid resolution of 0.01 in both \( q \) and \( p \) (momentum). The three zero-energy curves are given by the ridges of maximal FTLE and are overlaid with the analytical solution of these curves given by \( p = 0, q = 0, \) and equation (4.5). (b) Forward and backward FTLE flow fields for the SIS epidemic model with external fluctuations. The flow fields were computed using the Lagrangian given by equation (4.9) with \( \beta = 5.0 \) and \( \kappa = \mu + \gamma = 1.0 \). The integration time is \( T = 5 \) with an integration step size of \( t = 0.1 \) and a grid resolution of 0.01 in both \( I \) and \( p \) (momentum). The optimal path to extinction from the endemic state to the disease-free state and its counterpart optimal path from the disease-free state to the endemic state (found by computing the backward FTLE field) are given by the ridges of maximal FTLE and are overlaid with the analytical solution of the optimal paths.

Figure 4. FTLE flow field for the SIS epidemic model with internal fluctuations computed using the Hamiltonian given by equation (4.11) with \( \beta = 2.0 \) and \( \kappa = \mu + \gamma = 1.0 \). The integration time is \( T = 10 \) with an integration step size of \( t = 0.1 \) and a grid resolution of 0.005 in both \( I \) and \( p \) (momentum). The optimal path to extinction is given by the ridge of maximal FTLE.
Figure 5. (a) Numerically computed action (integrated along the optimal path found numerically from the FTLE flow field) versus reproductive number $R_0$ for the SIS epidemic model with internal fluctuations. The inset shows a portion of the graph near $R_0 = 1$. The numerically computed action is given by the black points, while the dashed, red curve shows an asymptotic scaling result that is valid near $R_0 = 1$, and is given by $S(R_0) = (R_0 - 1)^2/[R_0(1 + R_0)]$ (see Sec. 5 of the electronic online supplement).

(b) Numerically simulated (solid curve with black points) mean extinction time versus reproductive number for the SIS epidemic model with internal noise and the analytical prediction (dashed, red curve) found using the asymptotic scaling law that is valid near $R_0 = 1$. 