Periodically varying externally imposed environmental effects on population dynamics

M. Ballard,1,2 V. M. Kenkre,1 and M. N. Kuperman1,2

1Consortium of the Americas for Interdisciplinary Science and Department of Physics and Astronomy, University of New Mexico, Albuquerque, New Mexico 87131, USA
2Centro Atómico Bariloche and Instituto Balseiro, 8400 S. C. de Bariloche, Argentina

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Effects of externally imposed periodic changes in the environment on population dynamics are studied with the help of a simple model. The environmental changes are represented by the temporal and spatial dependence of the competition terms in a standard equation of evolution. Possible applications of the analysis are on the one hand to bacteria in Petri dishes and on the other to rodents in the context of the spread of the Hantavirus epidemic. The analysis shows that spatiotemporal structures emerge, with interesting features which depend on the interplay of separately controllable aspects of the externally imposed environmental changes.

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I. INTRODUCTION

The mathematical study of population dynamics has been a subject of great interest in recent years, with application widely spread among different fields. An example is the description of emerging patterns in bacterial colonies [1–8]. Other studies describe the behavior of the populations of superior organisms such as insects and rodents [9–12].

If, in basic models for the description of the evolution of a given population, we focus attention on reproduction, competition for resources, and diffusion, the Fisher equation [12,13] appears to be a useful mathematical tool:

\[
\frac{\partial u(x,t)}{\partial t} = D \frac{\partial^2 u(x,t)}{\partial x^2} + au(x,t) - bu^2(x,t).
\]

Diffusion with coefficient \( D \) is considered here as well as the growth of the population at rate \( a \) and a competition process weighted by \( b \), also containing environmental features. In the present paper, we consider effects of spatiotemporal dynamics in the nonlinear term as a representation of externally imposed or seasonal environmental variations. We thus take \( b \) in Eq. (1) to be space and time dependent, \( b(x,t) \). Environmental variations could alternatively be considered as affecting \( a \) (such that \( a \) is dependent on time and space rather than \( b \)) as has been done in some earlier studies [3,4,8]. However, we restrict our attention to a constant \( a \) and varying \( b(x,t) \). This allows us to define, in a straightforward manner, two characteristic quantities, a Fisher velocity and a Fisher length in terms of a constant \( a \). Following Ref. [8] we consider a bounded region which will be referred to as a bubble of favorable conditions suitable for the survival of a given species. Outside of this region the environmental conditions are so harsh that there is little or no possibility of survival.

Associated with the Fisher equation [13] there is a natural velocity and a natural length, \( 2\sqrt{D/a} \) and \( \pi \sqrt{D/a} \), respectively. The former is known as the Fisher velocity and is the velocity at which fronts tend to travel after adjustments from most initial conditions [12]. Thus an arbitrary shape of the initial population with compact support will have fronts that move, after initial transients, with the Fisher velocity. It is an important quantity in experiments on bacteria with moving masks [3,4] as it represents the minimum mask velocity at which bacteria tend to extinction, rather than being able to follow the bubble through the combined effect of growth and diffusion. The Fisher length \( \pi \sqrt{D/a} \) arises [8,14] in the bacterial context as the minimum length of the mask below which the steady-state population of the bacteria vanishes as the bacteria diffuse out of the masked area quicker than they can grow to any finite saturation value. The Fisher length is thus the “diffusion length” within the growth time (reciprocal of the growth rate \( a \)).

Consideration of these two quantities suggests that it is natural to envisage two observations involving externally imposed periodic variation of the environment. The first is one in which experiments of the kind reported in Refs. [3,4] are carried out with the velocity varying periodically, in other words with the bubble repeatedly returning to its original position. This is the oscillating bubble case. The second experiment is one in which the bubble extent is made to vary from a large size to one below the Fisher length (i.e., the extinction size). The objective of the latter (breathing bubble) experiment is to analyze extinction tendencies under periodic variations of the bubble size. We analyze these two hypothetical experiments in this paper through numerical calculations based on a Crank-Nicholson scheme.

The present theoretical investigation has been motivated by two quite different experimental situations: on the one hand by a series of observations in the context of mice population dynamics of the Hantavirus [15], and, on the other, by an easily realizable modification of experiments [3,4] in the context of bacteria in a Petri dish. The population of rodents on mountain slopes involved in the spread of the Hantavirus epidemic undergoes spatiotemporal changes with change of seasons [15]. The habitable zone of the rodents moves periodically in time as a direct consequence of the seasonal variation of the environment. In the bacterial context, the externally imposed variation is provided simply by moving a mask over a Petri dish through attachment to an electric motor. In experiments reported thus far [3,4] the mask is moved unidirectionally at a constant velocity. In easily realizable modifications of the experiment that we suggest in the light of our present investigation, the mask would be made to move periodically. The vastly different realizations of peri-
II. TRAVELING BUBBLE

Solutions of the Fisher equation for bounded domains, corresponding to static bubbles, have been already discussed in Ref. [8] and references therein. A natural change, to be considered as a first approximation to the problem addressed here, is to allow a translation in the bubble. We have performed a brief analysis of situations where the velocity is constant and the system’s behavior corresponds to that of a moving front, as well as others where the velocity of the bubble changes continuously. For the former case, and as mentioned in Ref. [12], we recover a critical velocity beyond which the front will no longer survive; see bold curve of Fig. 1. As \( b \to \infty \) at the boundaries of the bubble, the curves will abruptly go to zero, rather than asymptotically as above. The analysis of this situation can be done by making a change of variables on Eq. (1). If we set \( \xi = x - ct \) we obtain the following equation which is valid in the moving frame accompanying the bubble:

\[
D \frac{\partial^2 u(\xi)}{\partial \xi^2} + au(\xi) - bu^2(\xi) + \epsilon \frac{\partial u(\xi)}{\partial \xi} = 0. \tag{2}
\]

By measuring the maximum population density \( u_{\text{max}} \) of the front solution or the steady solution in a moving frame, we obtain a curve that will be useful in interpreting the following results. As the bubble accelerates in the moving frame, the system is not allowed to relax into a steady state; see the accelerating curve of Fig. 1. The time involved in the changes induced by the acceleration of the bubble compete with the relaxation time of the system. We will leave these results for now and recall them later to understand the following analysis.

III. OSCILLATING BUBBLE

Here, the habitable bubble of fixed width oscillates as a whole. We study two cases: sinusoidal velocity, and a velocity which is constant but suddenly changes direction at a given distance away from the original position. All points within the amplitude of oscillation will thus be covered periodically for varying amounts of time. For the purpose of comparison, we will consider the velocity to be an average velocity for the sinusoidal case. The parameters involved are the average velocity of the bubble \( v \), the amplitude of oscillation characterized by the distance between the center of the bubble at the turning points \( A \), the length of the bubble, \( W \), and the Fisher parameters: the diffusion coefficient \( D \), the growth rate \( a \), and the competition term \( b(x,t) \). For the following analysis, \( b(x,t) \) is a step function that defines the shape of the bubble. We have examined the following aspects of Eq. (1):

(i) The role of each of the three parameters, \( v \), \( A \), and \( W \), in the dynamics of the population.

(ii) How the two cases (sinusoidal velocity and constant velocity) differ from each other.

(iii) Qualitative characteristics and comparisons between solutions for various parameter sets.

(iv) The evolution of the population as diffusion goes to zero.

(v) The appearance of a curious dipping phenomenon.

A. Numerical results for \( D > 0 \)

It is helpful to divide the behavior into two regimes which correspond to no extinction and extinction. The parameters \( A \), \( W \), and the critical bubble width, \( W_c = \pi(D/a) \) \([8,14]\), are sufficient to define these regimes. In the steady state, bubble widths smaller than the critical value are unable to support a population density. This feature now manifests itself in the amount of overlap between the two extremes of an oscillating bubble. For \( A < W \) there will be an area that is permanently covered by the bubble. When this covered area, from now on referred to as the overlap, is greater than the critical width, the population will always survive to some degree. This regime is bounded by \( 0 < A < (W-W_c) \).

In the other regime, extinction may occur when there is an overlap that is less than the critical value, \((W-W_c) < A \). The distance between the inner edges of the bubble when \( A > W \) will be called the separation \( A-W \). Naturally, when there is a separation, no single location will receive continuous coverage by the bubble.

For either regime, if the velocity is below the Fisher velocity, \( v_f \), the population will be able to follow the oscillating bubble. As the velocity increases it becomes more difficult for the population density to keep up with the moving bubble (via diffusion and new growth). Some of the population falls prey to the harsh environment which quickly reduces the density. If \( b(x,t) \) is infinite for \(|x| > W/2 \), the exposed population will go extinct, never to return. One might base the analysis on a representation of the adverse conditions outside the protection provided by the mask by taking \( b \) to be infinite outside the bubble. However, this would neither be realistic nor possible in a numerical computation. Therefore we have considered \( b \) outside the bubble to be a large but finite number. Most of the phenomena analyzed in this paper appear as transients which are both realistic and interesting but would
to consider arriving at this minimum outside of the mask, in our case 10−5 in the units used, introduced of discrete entities. The value we have chosen for population as effectively extinction for populations comprised with the amount of overlap or separation. 

oscillates spatiotemporally as the velocity increases and varies with the amount of overlap or separation.

For the zero extinction regime, high velocities effectively wash out the oscillatory behavior and the peaks merge so that the length of permanent overlap then determines the population density. The behavior of the population in the extinction regime, \((W−W_c)<A\), is the same except for one crucial characteristic. Again, the peaks at the endpoints come closer together for higher velocities, but they never merge. The population dies out before a constant density forms in the center. As \(A\) increases, extinction occurs for smaller and smaller velocities.

In the discussion above an average velocity of the bubble was assumed. We now make a distinction between two types of velocities and their relative effects on the population density. A bubble moving with sinusoidal velocity lingers for longer time over the end points of its trajectory and for shorter time in the center as compared to the constant velocity bubble. For a single location, a larger population density is permitted to grow if more time is spent under the bubble. The basic qualitative differences of the population density thus follow from these two aspects. Though the two velocities produce the same essential behavior, we find that there indeed exists a difference between the two after long times. This will be addressed in the final section.

To further understand the structural changes of the population, we have examined the maximum population density as it oscillates parametrically in time. The shape that is outlined can be roughly characterized by a bowtie, Fig. 3. The densities of the maxima, minima, center, and the spatial extremes are the four features that comprise the qualitative differences between bowtie morphologies. As the average velocity increases these four points of comparison within the bowtie undergo changes in proportion. The maxima of the bowtie move closer to the center (recall that in the no-extinction regime these maxima eventually merge). The minima move farther from the center. The spatial spread of the bowtie is smaller and the overall values of the population density decrease for all positions. The difference between the maxima and minima of the bowtie, \(A_b\) (the amplitude of oscillation of the maximum population density) initially increases and then decreases as velocity increases. Figure 4 depicts the abrupt appearance and then gradual decay of the
diffusion is thus limited by the time a single location is covered by the bubble and has little effect when velocity is high. A solution thus exists in the high velocity limit; refer to Fig. 4.

FIG. 4. The amplitude of the bowtie is plotted against the average velocity (which here varies sinusoidally). The inset shows the average \( u_{\text{max}} \) after long time. The combination of the both show the validity of the analytic solution in the high velocity regime when the curves for \( D=aW^2 \) and \( D=0 \) converge.

The bowtie as a function of velocity. The bowtie appears in both nondiffusive systems and diffusive systems. Bowties of the former develop amplitudes larger than those of the latter. Also, bowties are formed at lower velocities for the nondiffusive systems and diffusive systems. Bowties of the bowtie as a function of velocity. The bowtie appears in both validity of the analytic solution in the high velocity regime when

\[
\frac{\partial u(z,t)}{\partial t} = au(z,t) - b \left( z + \int_0^t v(s)ds \right) u(z,t)^2 + D \frac{\partial^2 u(z,t)}{\partial z^2}.
\]

The transformation \( \phi=1/u \) yields an equation of the form

\[
\frac{\partial \phi(z,t)}{\partial t} + a \phi(z,t) = b \left( z + \int_0^t v(s)ds \right) + D \left[ \frac{\partial^2 \phi(z,t)}{\partial z^2} - \frac{2}{\phi} \left( \frac{\partial \phi}{\partial z} \right)^2 \right].
\]

By allowing \( D=0 \), we recover the equation for a highly damped linear oscillator that is driven by an external force proportional to \( b \left[ z + \int_0^t v(s)ds \right] \). The population will thus follow the periodic forcing function with a lag. This oscillator interpretation clarifies the structure of the bowtie. Each location grows periodically, with a temporal lag, at the behest of the driving term. The locations at the edges of the bubble’s spatial trajectory will have higher population densities because the driving oscillator lingers for longer time at a lower value (less harsh conditions).

The solution, for \( D=0 \), as derived in Ref. [16], was found to be

\[
\phi(x,t) = e^{-at} \phi_0 \left[ x - \int_0^t v(s)ds \right] + \int_0^t e^{-a(t-t')} b \left[ x - \int_0^t v(s)ds \right] dt'.
\]

and thus \( u(x,t) \) explicitly as

\[
u(x,t) = \frac{u_0 \left[ x - \int_0^t v(s)ds \right]}{e^{-at} + u_0 \left[ x - \int_0^t v(s)ds \right] \int_0^t e^{-a(t-t')} b \left[ x - \int_0^t v(s)ds \right] dt'}.
\]

As the bubble’s velocity greatly surpasses the Fisher velocity, the effect of diffusion becomes negligible. The diffusion length per time competes with the velocity of the bubble. Diffusion is thus limited by the time a single location is covered by the bubble and has little effect when velocity is high. A solution thus exists in the high velocity limit; refer to Fig. 4.

When the velocity is relatively low, \( v < v_f \), population migration occurs via diffusion and new growth. By letting \( D \to 0 \) (correspondingly, \( v_f \to 0 \) and critical overlap \( \to 0 \)) one might conclude that a population cannot maintain itself unless there is a bit of overlap. This is true when the velocity is such that growth cannot occur within the short period of coverage by the bubble. However, when these two rates (bubble velocity and \( aW \)) are comparable, oscillations in the population density again arise in the diffusionless case. The bowties in these circumstances vaguely resemble those of before. When \( A < W \) the maximum population density is always saturated; there are no means by which to leave a sheltered region without diffusion. When \( A > W \), the bowtie limits to a U shape.

IV. BREATHING BUBBLE

The breathing bubble corresponds to a situation when the width of a anchored bubble varies in time. As before, we consider several cases and for each of them a family of values for the involved parameters. These parameters are the velocity of the variation of the width, the mean width of the bubble, the amplitude of the breath, and again, the Fisher
parameters of Eq. (1). We are interested in analyzing the relationship between the existence of critical values and the response of the population density to a changing environment. We first take the case when the bubble width oscillates at a constant speed but always preserves a size above the critical one. The interesting aspect of this situation occurs at the boundaries of the bubble where population fronts are formed. The fronts can follow the movement of the bubble when the velocity of the breathing motion is under a critical value. Above this critical value, the population is confined to the minimum size of the bubble, which indicates that the population front can no longer follow the changes of the bubble. To show this we display in Fig. 5 the relative amplitude of the oscillation of the population front, $A_f/A$ vs the speed of the bubble breath, where $A$ is the amplitude of breathing.

We observe that at a critical value this amplitude goes to zero. That is, in spite of the changing size of the bubble, the population density remains bounded statically. Instead of a constant value for the velocity of breathing, we can take a fixed frequency for a sinusoidal behavior. The frequency is associated with the breathing motion of the bubble boundaries. The results are analogous to those displayed in Fig. 5. No qualitative differences were found.

![FIG. 5. Amplitude of the front oscillations for several velocities of the bubble where $D=aW_c^2$.](image1)

![FIG. 6. Amplitude of the front oscillations for several values of the inverse of the frequency of breathing of the bubble where $D=aW_c^2$.](image2)

![FIG. 7. Maximum value adopted by the population as a function of the instantaneous width of the bubble. The velocity of breathing is: (a) $0.1aW_c$, (b) $1aW_c$, (c) $10aW_c$, (d) $40aW_c$, and $D=aW_c^2$.](image3)
When the minimum size attained by the bubble during the breathing is under the critical size, the situation is different. It is not enough to plot, as in previous examples, the amplitude of the oscillation of the front because some variations in the maximum density of the population are also observed. To visually understand what is happening we again plot the temporal behavior of maximum value of the population, \( u_{\text{max}} \), as a function of the momentary width of the breathing bubble. This is what is shown in Fig. 7, where the trajectories are traversed clockwise. As observed in the plots, if the velocity is low enough, the size of the bubble will be under the critical size long enough to provoke the extinction of the population. On the contrary, for higher values of the velocity there is a range of values at which the population can survive. The response of the population to changes is much slower than the dynamics of the environment. Each of the results obtained in this case are essentially the same as the corresponding results from the oscillating bubble. In particular, the results displayed in Figs. 5 and 6 can be associated with the discussion included in the previous section, those included in Fig. 7 are analogous to the bowtie effect.

V. CONCLUSIONS

In the present work, we have numerically studied the behavior of a population whose dynamics are described by a modified Fisher equation with additional environmental variations. Analytic solutions were also considered for simplified examples. The results presented here show that besides the expected behavior some surprising features arise. There exist two main parameters that characterize the evolution of a given population: the Fisher length and Fisher velocity. A given group of individuals will not survive if the habitat size is smaller than the Fisher length. At the same time, the population will not be able to maintain saturation when the habitat moves with a velocity higher than the Fisher velocity. With these two considerations we have examined two types of periodically varying environments. Though extinction of the population was verified in conditions corresponding to the two situations mentioned above, it was also found that some nontrivial scenarios can also arise. Situations in which extinction was expected exhibited what we called the bowtie effect. An interplay between the time associated to the transient towards extinction and the periods of the changing environments is the cause of unexpected behavior of the population. We have also verified that there is a correspondence between the results obtained when considering oscillatory or breathing bubbles, though the situations are different.

For completeness, the case when the temporal variation of the external conditions is included in the linear term was also analyzed. Two cases were considered, the first one associated with very harsh conditions outside the bubble, leading to extinction, and the second one corresponding to an environment turning more hostile, yet still habitable. The coefficient \( a \) takes negative values in the first case and positive but small \((a \ll 1)\) in the second situation. We have found results analogous to the previous ones provided that the relaxation time to extinction is of the order of the corresponding relaxation times in the cases presented in Secs. II–IV \([17–21]\). Values greater than \(-1\) warrant this condition. All the phenomena reported in this paper for changing \( b \) were also obtained by keeping \( b \) constant and varying \( a \).

A diffusionless approximation helped us to understand the origin of the observed phenomena in terms of oscillating perturbations. Finally, it should be mentioned that a curious effect arises the presence of constant bubble velocity and diffusion. Probing longer times reveals an overlying oscillation in the population density profile. For diffusive systems, \( D > 0 \), and constant velocities, the entire bowtie dips down and then back up again in a periodic manner with constant amplitude. For different velocities the bowtie dips and rises with different periods and amplitudes; see Fig. 8. As velocity increases, the frequency nor the amplitude of the alternative dip do not necessarily vary accordingly. Sinusoidal velocities do not produce this effect. Further analysis is necessary to completely understand the origin of this behavior and its relation to the two required conditions, diffusion and constant bubble velocity.

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