LIFE-HISTORY TRADE-OFFS INFLUENCE DISEASE IN CHANGING CLIMATES: STRATEGIES OF AN AMPHIBIAN PATHOGEN

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Abstract. Life-history trade-offs allow many animals to maintain reproductive fitness across a range of climatic conditions. When used by parasites and pathogens, these strategies may influence patterns of disease in changing climates. The chytrid fungus, Batrachochytrium dendrobatidis, is linked to global declines of amphibian populations. Short-term growth in culture is maximal at 17°–25°C. This has been used in an argument that global warming, which increases the time that amphibians spend at these temperatures in cloud-covered montane environments, has led to extinctions. Here we show that the amphibian chytrid responds to decreasing temperatures with trade-offs that increase fecundity as maturation rate slows and increase infectivity as growth decreases. At 17°–25°C, infectious zoospores encyst (settle and develop a cell wall) and develop into the zoospore-producing stage (zoosporangium) faster, while at 7°–10°C, greater numbers of zoospores are produced per zoosporangium; these remain infectious for a longer period of time. We modeled the population growth of B. dendrobatidis through time at various temperatures using delayed differential equations and observational data for four parameters: developmental rate of thalli, fecundity, rate of zoospore encystment, and rate of zoospore survival. From the models, it is clear that life-history trade-offs allow B. dendrobatidis to maintain a relatively high long-term growth rate at low temperatures, so that it maintains high fitness across a range of temperatures. When a seven-day cold shock is simulated, the outcome is intermediate between the two constant temperature regimes, and in culture, a sudden drop in temperature induces zoospore release. These trade-offs can be ecologically important for a variety of organisms with complex life histories, including pathogenic microorganisms. The effect of temperature on amphibian mortality will depend on the interaction between fungal growth and host immune function and will be modified by host ecology, behavior, and life history. These results demonstrate that B. dendrobatidis populations can grow at high rates across a broad range of environmental temperatures and help to explain why it is so successful in cold montane environments.

Key words: amphibian; Batrachochytrium dendrobatidis; Chytridiomycosis; climate; disease; life history; population decline; temperature; trade-off.

INTRODUCTION

Rapid declines in many amphibian populations are occurring globally (Stuart et al. 2004). Some of these declines are associated with the amphibian chytrid fungus, Batrachochytrium dendrobatidis, the agent of chytridiomycosis, an emerging infectious disease of amphibians (Berger et al. 1998, Daszak et al. 2003). High infection prevalence and outbreaks of chytridiomycosis occur most frequently at upland sites and during the cold season (Berger et al. 1998, 2004, Bradley et al. 2002, Retallick et al. 2004, Woodhams and Alford 2005, Bosch and Martinez-Solano 2006, Lips et al. 2006, Rachowicz et al. 2006). Analysis of 56 sites in Australia indicated that the amphibian chytrid’s presence was significantly related to temperature but not elevation or rainfall (Drew et al. 2006). Ecological niche modeling based on known localities of occurrence predicted an extensive range of suitable pathogen habitat, with a mean annual temperature of 13.1°C (Ron 2005). However, the density of the amphibian chytrid grown in culture increased most quickly (or had maximal short-term growth) at temperatures between 17° and 25°C (Longcore et al. 1999, Piotrowski et al. 2003).
This growth pattern in culture has led to the suggestion that global warming is creating an optimal climate for chytridiomycosis in high-elevation regions of Central America, where increasing cloud cover may be modulating daily minimum and daily maximum temperatures to converge around a “chytrid thermal optimum” (Pounds et al. 2006). Piotrowski et al. (2004) demonstrated that after 23 days in culture at 10°C, the density of *B. dendrobatidis* in culture converged with that in cultures grown at higher temperatures. Here we investigate whether this effect is due to nutrient limitation in culture or if there are life-history trade-offs that compensate for slower initial population growth at lower temperatures. We examine the hypothesis that the amphibian chytrid adapts to temperatures below 17°C to 25°C through life-history trade-offs.

The amphibian chytrid develops through multiple stages in a single host. The life stages include a mobile infectious zoospore stage, a sessile developing thallus, and a zoospore-releasing zoosporangium (Fig. 1; also see Plate 1). Growth of the thallus may be colonial or a single unit termed monocentric (Longcore et al. 1999, Berger et al. 2005). In nutrient-rich medium, such as 1% tryptone, amphibian chytrid thalli mature extremely quickly through an exponential growth phase until the population levels off at a nutrient-limited stationary phase that releases few infectious zoospores. The stationary phase is reached in approximately 6–7 days at temperatures between 17°C and 28°C (Longcore et al. 1999, Piotrowski et al. 2004). This rapid limitation of growth is not likely to occur within living amphibian skin, and longer-term growth is investigated here.

The zoological literature suggests that trade-offs exist between fecundity and rate of maturation, particularly in animals with complex life histories. Mortality rate and maturation rate can vary with temperature, density, or other factors. Growth rate and fecundity can compensate for these changes to maximize reproductive fitness in marine invertebrates (Llodra 2002), insects (Howe 1967, Sisodia and Singh 2002), amphibians (Smith-Gill and Berven 1979, Werner 1986), fish (Kavanagh and Alford 2003, Savage et al. 2004), reptiles (Angilletta and Dunham 2003), and birds (Spencer and Bryant 2002). Trade-off strategies may be used by other organisms (Savage et al. 2004), including parasites and pathogens, and provide a framework for understanding disease emergence in the context of global change. Applying this framework to the amphibian chytrid fungus, which shows development through multiple life-history stages within a single host, suggests several a priori hypotheses. Specifically, this pathogen may adapt to changing temperatures by trading fecundity for maturation rate. We hypothesized that at relatively low temperatures, chytrid thalli would develop more slowly and mature at a larger size. Zoosporangia may thus produce more propagules at low temperatures than they do at higher temperatures. Infectivity, dependent upon the life span of the zoospore stage, may also be traded for rapid encystment and growth. That is, at lower temperatures, zoospores may survive longer. *Batrachochytrium dendrobatidis* grows well in culture and develops through all of the same life stages as on the keratinized skin of amphibians and the mouthparts of tadpoles (Berger et al. 1998, Longcore et al. 1999). Therefore, experiments in culture may be a good surrogate for modeling in vivo pathogen population changes while eliminating the effects of variation in host responses (Woodhams et al. 2006, 2007).

Here we present a series of laboratory experiments designed to quantify components of *B. dendrobatidis* life history. These experiments provide temperature-dependent parameter estimates that we incorporate into a population model that determines their net effects on long-term population growth rate.

**METHODS**

**Quantifying zoospore survival and encystment**

We quantified *B. dendrobatidis* zoospore survival rate and encystment rate at different temperatures. For this experiment, *B. dendrobatidis* isolated from an eastern banjo frog, *Limnodynastes dumerili*, Victoria, Australia (isolate number 98-1469/10), was maintained at James Cook University, Townsville, Australia, in TGH-L liquid medium (1.6% tryptone, 4% gelatin hydrolysate, 0.5% lactose) in 25-cm² cell culture flasks. After 4 d incubation at 23°C, the culture flasks were scraped to remove the attached cells and centrifuged at 980 × g (or 9600 m/s²) for 5 min to purify the zoospores in the supernatant. Zoospores were checked for activity and
counted using a hemocytometer slide at 400× magnification. Zoospores were prepared at a concentration of $162 \times 10^4$ zoospores/mL, and 5 mL were placed in each of 12 sterile glass vials with TGHl liquid medium. Three vials were placed at each of four temperatures in incubators or in controlled-temperature rooms: (1) 28°C, (2) 23°C, (3) 14.5°C, and (4) 4°C. After placement at each temperature for 0, 2, 5, 22, and 45 h, the numbers of zoospores were estimated by removing 10 μL of solution from gently mixed cultures and taking the mean zoospore count from two squares on a standard hemocytometer slide. Zoospores were classified as either active (swimming or moving) or inactive. Since zoospores move rapidly, counts of active zoospores were made at a single time point for each small square of the hemocytometer.

Over this short time period, the only processes affecting zoospore abundance are mortality and encystment (no production of new zoospores). Zoospores encysted on the sides of the glass flasks. We used motility as a surrogate measure for viability (Berger 2001) because nonmotile zoospores that did not encyst did not appear to develop into thalli in solution. *Batrachochytrium dendrobatidis* does not appear to produce a resting stage arising from zoospores (Longcore et al. 1999, Berger et al. 2005, Di Rosa et al. 2007). With these assumptions, the rate of change of live zoospores in the experiment is predicted to be: $dZ(t)/dt = -(d_Z + m_Z)Z(t)$, where $Z(t)$ is the density of live zoospores, $d_Z$ is the per capita zoospore death rate, and $m_Z$ is the per capita encystment rate. Dead zoospores accumulate at: $dD(t)/dt = d_Z Z(t)$, where $D(t)$ is the density of dead zoospores. Therefore, the predicted densities of live and dead zoospores are

$$Z(t) = Z(0) \exp[-(d_Z + m_Z)t]$$

and

$$D(t) = D(0) + d_Z Z(0) \left[1 - \exp[-(d_Z + m_Z)t]\right]/(d_Z + m_Z)$$

where $Z(0)$ and $D(0)$ are the initial densities of live and dead zoospores, respectively.

The parameter combination $(d_Z + m_Z)$ was estimated for each temperature as the slope of the trajectory of the log-transformed live zoospore density through time. The initial zoospore density (live + dead) was $162 \times 10^4$ zoospores/mL; however, the initial proportion of live zoospores in this sample was not measured directly. Therefore, $Z(0)$, which should be the same for all temperature treatments, was calculated as the average of the initial densities of live zoospores across the four temperature treatments, which were estimated for each treatment as the intercept of the regression ($\ln(Z(t)) = \ln(Z(0)) - (d_Z + m_Z)t$). The parameter combination $(d_Z + m_Z)$ was then calculated for each temperature with $Z(0)$ set to this common value. The initial density of dead zoospores, $D(0)$, was assumed to be the same for all temperature treatments and was calculated as $162 \times 10^4$ zoospores/mL $- Z(0)$. Parameter $d_Z$ was calculated for each temperature by least-squares fitting of Eq. 2 to the observed time series of dead zoospores, using the estimated value of $(d_Z + m_Z)$ for that temperature.

**Quantifying thallus maturation and zoospore production**

The life-history traits of thallus maturation (time to sporulation) and fecundity (number of zoospores produced) were experimentally quantified and examined for the effects of temperature and initial density. We grew the amphibian chytrid type isolate (JEL 197) under variable temperature regimes at Vanderbilt University. Eight culture flasks containing 10 mL 1% tryptone broth with 1% penicillin/streptomycin were each inoculated with $5 \times 10^5$ zoospores/mL. We obtained zoospores by the following method: We scraped culture flasks of *B. dendrobatidis* and applied ∼1.5 mL to 1% tryptone agar plates. Plates were dried for ∼1 h in a sterile hood, incubated for 7 d at 23°C, and rinsed with 3 mL broth once and again after soaking for 20 min. Thalli were removed by filtering broth through a 20-μm Spectra/Mesh nylon filter (Spectrum Laboratories, Rancho Dominguez, California, USA). Gently mixed zoospores in broth were counted on a hemocytometer slide as described in Quantifying zoospore survival and encystment and diluted in broth to the desired concentration. We placed four flasks into each of two treatments: $10^6 \pm 0.5^\circ C$ and $23^\circ C \pm 0.5^\circ C$. Two counts were made from each flask initially and each day for 14 d. After gently mixing the culture flasks, 10 μL was added to one side of the hemocytometer slide, and to the other side of the slide 10 μL of zoospores diluted 1:10 in Lugol solution (Sigma Chemical, St. Louis, Missouri, USA) was added to kill and stain the zoospores. Numbers of active, inactive, and total stained zoospores were counted. The proportion of mean active to mean total (active plus inactive zoospores) was calculated at each time point. In addition, the mean live zoospores per initial live zoospores (at day 0) was calculated at each time point.

When *B. dendrobatidis* thalli reach maturity they become zoosporangia, release their zoospores, and then die (Longcore et al. 1999). In each flask, no zoospores remained active from the initial inoculum after the first day; however, these initial zoospores resulted in an observable pulse of zoospores a number of days later. We assumed that the zoospores in this first pulse were the first generation offspring of the initial zoospore inoculum. At each temperature we calculated the average time to sporulation ($t$) and the total number of viable zoospores released per initial live zoospores present ($F$) by least-squares fitting of the time series of the average number of live zoospores per initial live zoospore (from time $t = 0$ through the end of the first pulse) to the output of the model:

$$dZ/dt = R(t) - (m_Z + d_Z)Z(t)$$

where the parameter combination $(m_Z + d_Z)$ at each
temperature was taken to be the value calculated in the experiments described in Quantifying zoospore survival and encystment, above (for 10°C, the value was interpolated from the experimental results at 4°C and 14.5°C). \( R(t) \) is the recruitment of new active zoospores representing the progeny of the initial inoculum of zoospores at time \( t = 0 \). A delayed gamma distribution was used for \( R(t) \), to allow for the observed variability in the time to sporulation. The delayed gamma distribution has density function

\[
R(t) = F (t - \phi) \gamma(a-1) \exp \{- (t - \phi)/s\}/(s^a \Gamma(a)),
\]

where \( a \) is the shape parameter, \( s \) is the scale parameter, \( \phi \) is the lag, and \( \Gamma(a) \) is a gamma function. \( F \) is the total number of viable zoospores produced per initial zoospore, and \( \tau = \phi + a \times s \) is the mean time to sporulation.

The fitting was performed using the optim function in the program R (R Development Core Team 2007) to determine the values of \( F, \phi, a, \) and \( s \) that minimized the sum-of-squares difference between the output of Eq. 3 and the observed values.

To examine the effect of changing temperature on the quantity and timing of zoospore production, 12 flasks were inoculated with zoospores (JEL 197) in 1% tryptone broth with 1% penicillin/streptomycin antibiotic, and zoospores were counted daily as described in Quantifying zoospore survival and encystment. All flasks were incubated for 5 d in a constant 26.5°C incubator at Vanderbilt University. Flasks were then placed at: (1) 7.5°C, (2) 23.5°C, or (3) 26.5°C and monitored for 10 d.

Thallus maturation was also measured by examining cultures (JEL 197) grown in 35-mm glass bottom culture dishes (Mat Tek, Ashland, Massachusetts, USA) under 400× magnification with an inverted microscope (Olympus CK40; Olympus, Melville, New York, USA). Living colonies attached to the glass slide were monitored daily for motile zoospores. Time to sporulation was recorded at the first time point motile zoospores were observed. Culture dishes were inoculated with 10, 100, 500, or 1000 zoospores/µL in 1% tryptone broth and grown at either 10°C or 23°C. We analyzed digital photographs with ImageTool version 2.0 (University of Texas Health Science Center, San Antonio, Texas, USA) and quantified the effect of initial density on development (time to sporulation and size of thalli). Time to zoospore release could not be used to estimate model parameter \( \tau \) in this experiment because zoospores were released over a period of days and were not quantified.

To measure the effects of density on \( B. \) dendrobatidis growth, zoospores were diluted with TGhL liquid medium to concentrations between 1 × 10^4 and 1 × 10^2, and 100 µL were pipetted into five replicate wells on 96-well plates. Plates were held in an incubator at 23°C for 7 d. Growth within the wells was measured daily by optical density at 490 nm on an MRX Microplate Reader (DYNEX Technologies, Chantilly, Virginia, USA). We quantified the recruitment of initial zoospore density on culture growth. Density effects were not included in the model describing temperature-related trade-offs.

Building a model of long-term population growth rate

We developed a stage-structured model of \( B. \) dendrobatidis population growth, using the experimentally determined parameter values. The model incorporates the temperature-dependent life-history strategies observed in the experiments and determines their net effects on long-term population growth rate. The model divides the population of \( B. \) dendrobatidis into a mobile infectious zoospore stage, \( Z \), and a sessile developing sporangium stage, \( S \). The zoospore-releasing zoosporangium stage is short-lived relative to the other stages; it is assumed that zoosporangia release zoospores after a fixed (but temperature-dependent) development time. For simplicity, this model assumes a fixed lag between the time that zoospores encyst and the time at which the resulting sporangia start to produce new zoospores, although some variability in this time lag was observed in the experimental results in Quantifying thallus maturation and zoospore production, above. The developing sporangium stage, \( S \), incorporates both the thallus and zoosporangium stages until zoospores are released. Therefore the zoospore-releasing stage does not need to be modeled explicitly.

The growth of the population is modeled as a linear system of delayed-differential equations:

\[
\frac{dZ(t)}{dt} = fZ(t) - mZ(t) - dZ(t)
\]

(4)

\[
\frac{dS(t)}{dt} = mZ(t) - MS(t) - dS(t)
\]

(5)

\[
MS(t) = mZ(t - \tau)\exp\{-dS(t)\}
\]

(6)

where \( MS(t) \) is the maturation rate out of the developing thallus stage.

The first term in Eq. 4 represents the instantaneous rate of production of mobile zoospores. Each developing thallus produces new zoospores at the time that it matures into the zoosporangium stage, which occurs at rate \( MS(t) \). The second term in Eq. 4, which is equal to the first term of Eq. 5, represents the rate at which zoospores settle out of the mobile zoospore stage and become developing thalli. The variable \( mZ \) is the per capita rate of encystment of zoospores per hour. This constant rate of encystment means that the model assumes that the time spent in the mobile zoospore stage is exponentially distributed, with an average duration of 1/mZ (in the absence of loss due to other causes). Variables \( dZ \) and \( dS \) are density-independent per capita death rates of zoospores and developing thalli, respectively. \( \tau \) is the time from zoospore encystment until production of zoospores from the zoosporangium. Therefore, the maturation rate out of the developing thallus stage, \( MS(t) \) in Eq. 6 is equal to the rate that developing zoospores enter that stage \( \tau \) hours ago, multiplied by the probability of surviving through those \( \tau \) hours. We modeled the populations of active
zoospores and living thalli through time at constant 10°C and constant 23°C. Simulations of the model at constant temperatures were performed using a fourth/fifth-order Runge-Kutta algorithm, using the Solver software (Gurney et al. 1998), which handles time lags efficiently. Initial conditions for the simulations were 100 zoospores and no sporangia in any age class (i.e., \( Z(0) = 100 \) and \( S(t) = 0 \) for all \( t \leq 0 \)).

The long-term intrinsic rate of increase of the \( B. \ dendrobatidis \) population (actually the eigenvalue, \( \lambda \)) can be calculated from the characteristic equation:

\[
FmZ \exp\{-\lambda t\} - \lambda - \mu = 0
\]

where \( F = f \exp\{-dZ\} \) is the effective number of zoospores produced per individual entering the thallus stage and \( \mu = mZ + dZ \). Unfortunately it is not possible to produce an analytical expression for \( \lambda \), but it can be solved for numerically.

Natural thermal conditions experienced by \( B. \ dendrobatidis \) in a complex environment include temperature fluctuations and host behavioral adjustments. We considered populations developing through changing temperatures, or “cold shock.” When temperatures changed, we used the weighted average of the model parameters considering the time spent at each temperature, rather than locking the zoospores or thalli into development at the rates of the initial temperature. In order to simulate the population in a fluctuating temperature environment, we discretized the model on an hourly time step and calculated parameters based on the temperatures experienced by each hourly cohort (see Appendix). We simulated a population at 23°C experiencing a one-week long cold shock to 10°C (similar to the experiment shown in Fig. 2B) to determine its effect on long-term population growth rate.

In this model we have not included the observed effects of inoculum density on the rate of thallus maturation (Fig. 3), because any resource limitation resulting in density-dependent demographic rates on amphibian skin under natural conditions are likely to be different from those observed in culture.

**RESULTS**

**Quantifying zoospore survival and encystment**

We predicted that at low temperatures zoospores may trade rapid encystment for increased survival. Zoospores...
were slowly lost from solution at 10°C due to encystment (4% of zoospores per hour). At 23°C, zoospores were lost from solution more quickly due to encystment (17.8% of zoospores per hour). Zoospore mortality, including loss of infectivity, was 0.3% per hour at 10°C and 1.61% per hour at 23°C. Thus, as predicted, zoospores died and encysted more quickly as temperature increased (Fig. 2A, Table 1).

Quantifying thallus maturation and zoospore production

At low temperatures thalli took longer to mature into zoosporangia, but more zoospores were produced per zoosporangium than at higher temperatures (Fig. 2B). The maturation time of thalli was ~87.7 h at 23°C, but 222.7 h at 10°C, given an inoculum of 500 zoospores/μL. When mature, the number of zoospores produced (free zoospores counted in culture) per initial zoospore inoculated was ~64.8 zoospores per initial zoospore at 23°C and 160.5 zoospores per initial zoospore at 9°C.

The amphibian chytrid is temperature sensitive and can be killed by temperatures exceeding 30°C in vitro and in vivo (Longcore et al. 1999, Woodhams et al. 2003, Berger et al. 2004, Piotrowski et al. 2004). Close to the maximal temperature tolerance for B. dendrobatidis few zoospores were released (Figs. 2 and 4). However, when temperatures were suddenly reduced from 26.5°C to 7.0°C, zoospore production greatly increased within 3 d (Fig. 2C). This increase in zoospore production was not observed when temperatures were suddenly reduced from 26.5°C to 23.5°C (Fig. 2C).

Batrachochytrium dendrobatidis cultures grew more quickly when inoculated with larger numbers of zoospores (Fig. 3A). Zoospores alone could not be quantified by optical density. At 23°C, motile zoospores were observed after 4–5 d regardless of initial density. At 10°C, the initial zoospore density had a large effect on time to zoospore production; the time to thallus maturation was longer in cultures with lower initial density (Fig. 3B).
When grown for 5 d at constant 23°C, the size of thalli was smaller at higher densities (Pearson correlation, \( r_{35} = -0.344, P = 0.04 \); Fig. 3C). At higher densities, thalli grew and matured more quickly than at lower densities; they also grew to a smaller size. The number of zoospores produced per zoosporangium grown under different density conditions was not measured.

Growth form in culture was also affected by temperature (Fig. 4). Very large colonial growth structures (30–40 \( \mu \text{m} \) diameter) were observed in cultures grown at 23°C (Fig. 4B) but not at 10°C or 26°C (Fig. 4A, C). Although normally spherical (Long-core et al. 1999), thalli became angular at 10°C. Thalli at 10°C and 26°C were more often monocentric, less clumped, and more evenly distributed than at 23°C.

**Modeled long-term population growth rate**

For the best estimates of the parameters from our data (Table 2), the population model predicts that *B. dendrobatidis* will have a higher intrinsic rate of increase at 23°C (\( \lambda = 0.044 \text{ h}^{-1} \)) than at 10°C (\( \lambda = 0.021 \text{ h}^{-1} \)). Fig. 5 illustrates the effects on the long-term population growth rate of the trade-off observed in this species between fecundity (\( F \), the number of zoospores produced per zoosporangium) and time to sporulation (\( t \)). The two surfaces in this figure show this relationship when the other two parameters in the model, the zoospore mortality (\( m_Z \)) and encystment (\( d_Z \)) rates, are set to the default values for either 10°C or 23°C. In terms of their net effects on long-term population growth rate, the increased fecundity at 10°C is not sufficient to offset the increased time to sporulation at this temperature, such that overall population growth rate is higher at the higher temperature (Fig. 5).

Fig. 6 shows simulations of the growth of populations of *B. dendrobatidis* over 50 d starting from 100 zoospores, using the default parameter values at 10°C and 23°C (Table 2). In the long term, the population density at the lower temperature remains below that at the higher temperature, but maintains a relatively high growth rate. Starting from 100 zoospores, after 50 d at 10°C, the model predicts almost \( 10^{12} \) live sporangia and \( 10^{24} \) live sporangia at 23°C (Fig. 6). Assuming an infection threshold of \( 10^7–10^8 \) sporangia beyond which amphibian mortality occurs (Carey et al. 2006), this model predicts mortality at \( \sim 360 \) h at 23°C and \( \sim 700 \) h at 10°C (Fig. 6), in the absence of host defenses. For susceptible amphibian species, mortality of infected hosts from chytridiomycosis can occur as fast as 18 and 48 d post-infection and varies with species and environmental conditions (Woodhams et al. 2003, 2007, Berger et al. 2004, Blaustein et al. 2005). Infection by even one zoospore can lead to mortality in some species (Carey et al. 2006).

Fig. 6 also demonstrates the effects of a cold shock on the growth of a *B. dendrobatidis* population over 50 d. This simulation was started at 23°C, and after 400 h the temperature was reduced to 10°C for a period of 7 d and then brought back up to 23°C. The cold shock led to a

**Table 1. Rates of zoospore mortality and encystment per hour under four temperature conditions for *Batrachochytrium dendrobatidis* (a chytrid fungus) isolated from an eastern banana frog *Limbodynastes dumerilli* (98-1469/10).**

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Zoospore death rate, ( d_Z ) (h(^{-1}))</th>
<th>Zoospore encystment rate, ( m_Z ) (h(^{-1}))</th>
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<tr>
<td>4.0</td>
<td>( 2.7 \times 10^{-3} )</td>
<td>( 3.4 \times 10^{-2} )</td>
</tr>
<tr>
<td>14.5</td>
<td>( 3.5 \times 10^{-3} )</td>
<td>( 5.2 \times 10^{-2} )</td>
</tr>
<tr>
<td>23.0</td>
<td>( 1.6 \times 10^{-2} )</td>
<td>( 0.18 )</td>
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<tr>
<td>28.0</td>
<td>( 4.1 \times 10^{-2} )</td>
<td>( 0.21 )</td>
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**FIG. 4.** Photographs of *Batrachochytrium dendrobatidis* grown for 8 d in culture at 10°C, 23°C, and 26°C (400× magnification). Note (A) the irregular shape of zoosporangia and abundance of zoospores at 10°C, (B) colonial growth at 23°C, and (C) spherical thalli with lack of zoospores at 26°C.
short-term decrease in the population growth rate, and the density of sporangia resulting after 50 d was intermediate between the constant temperature regimes.

**DISCUSSION**

Chytridiomycosis epizootics and high infection prevalence often occur during cold seasons or at high elevations (Berger et al. 1998, 2004, Bradley et al. 2002, Retallick et al. 2004, Woodhams and Alford 2005, Bosch and Martinez-Solano 2006, Lips et al. 2006, Rachowicz et al. 2006). Temperature-induced trade-offs in pathogen life-history characteristics may contribute to these biogeographic patterns of disease. We modeled population trajectories at constant 10°C, 23°C, or 23°C with a short 10°C cold shock and found that the fungus is predicted to have the highest long-term growth rate at the higher temperature. However, the long-term growth rate remains high over a wide range of temperatures. Our results support the chytrid thermal-optimum hypothesis, since they indicate that increases in environmental temperature will increase the rate of growth of *B. dendrobatidis* populations on hosts, until temperatures are above the optimal thermal range for the pathogen. Our results also indicate that the effects of variable environmental temperatures are likely to be complex; while a short period of “cold shock” delayed population growth in our simple model (Fig. 6), our laboratory experiments indicated that short periods of “cold shock” may stimulate the release of zoospores, which could compensate or even overcompensate for temporary decreases in the rate of pathogen population growth. Because each zoosporangium releases many more zoospores at lower temperatures, infected frogs may be more likely to transmit the infection. This effect could be compounded by their longer survival.

Adaptation to a range of thermal conditions is facilitated by trade-offs between survival and encystment and between fecundity and maturation rate. These trade-offs operate in a variety of other organisms and act to maximize reproductive fitness over a range of environmental conditions (Wilbur and Collins 1973,

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>10°C</th>
<th>23°C</th>
</tr>
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<tr>
<td>$F$</td>
<td>no. viable zoospores produced per zoosporangium</td>
<td>160.5</td>
<td>64.8</td>
</tr>
<tr>
<td>$\tau$</td>
<td>mean no. hours to sporulation per hour</td>
<td>222.7</td>
<td>87.7</td>
</tr>
<tr>
<td>$d_Z$</td>
<td>per capita zoospore mortality per hour</td>
<td>0.003</td>
<td>0.0161</td>
</tr>
<tr>
<td>$m_Z$</td>
<td>per capita zoospore encystment per hour</td>
<td>0.04</td>
<td>0.178</td>
</tr>
</tbody>
</table>

**PLATE 1.** Artistic representation of the zoosporangium stage of the fungus *Batrachochytrium dendrobatidis*. Artwork by Robert M. Brucker, James Madison University.
Smith-Gill and Berven 1979, Werner 1986, Kavanagh and Alford 2003, Savage et al. 2004). Thus, we show that an ecological framework often observed in zoological systems applies to a microbial pathogen.

Life-history trade-offs allow the amphibian pathogen *B. dendrobatidis* to maintain its fitness across a broad range of temperatures. If development of fatal chytridiomycosis is directly linked to pathogen population size (Carey et al. 2006), then pathogenicity may also be maintained by these trade-offs, at least in controlled experiments in which host factors such as behavior are constrained. Few experiments have tested the effects of temperature on living hosts infected with *B. dendrobatidis*, but these examples support our model. Carey et al. (2006) exposed boreal toads, *Bufo boreas*, to *B. dendrobatidis* at 12°C and 23°C. Survival time was not statistically significantly different between these temperature treatments. Berger et al. (2004) exposed great barred frogs, *Mixophyes fasciolatus*, to *B. dendrobatidis* at 17°C and 23°C and found no significant difference in survival time. Infected orange-eyed treefrogs, *Litoria chloris*, survived longer under naturally fluctuating temperatures between 13.5°C and 23.2°C with or without a 2-d exposure to 8°C than frogs kept at constant 20°C (Woodhams et al. 2003). Although naturally warm conditions may inhibit disease (Woodhams et al. 2003, Berger et al. 2004), the fungus appears to maintain pathogenicity over a broad range of lower temperatures. We found that life-history trade-offs facilitate the fitness of *B. dendrobatidis* in colder environments.

**Zoospore survival and encystment**

*Batrachochytrium dendrobatidis* zoospores are clearly active for a longer time in cold conditions than in relatively warmer conditions (Fig. 2A, Table 1). This confirms a finding by Berger (2001), who collected *B. dendrobatidis* zoospores into distilled water and used motility as a measure of survival time at two temperatures, 4°C and 23°C. She found that few zoospores were active after 24 h and that more zoospores survived at 4°C than at 23°C. Hyatt et al. (2007) quantified zoospores by flow cytometry and also found that zoospores came out of solution faster at 23°C (~2.26% per hour) than at 4°C (~0.43% per hour). Zoospores may therefore move greater distances before encysting in cold conditions, and they may be infective for a longer period of time than in warmer conditions. Formation of monocentric thalli (single individuals) rather than...
colonies may be favored at low temperatures (Fig. 4). Zoospores encysting quickly may add to the size of colonies, whereas zoospores surviving longer and moving greater distances may start more colonies, perhaps adding to the surface area of skin covered by the fungus.

**Thallus maturation and zoospore production**

These results have important implications for understanding disease virulence and transmission. Transmission efficiency is likely increased at lower temperatures where it appears that larger numbers of zoospores are produced per zoosporangium, and these are likely to remain infectious for a longer period of time than at warmer temperatures. The number of active zoospores counted per inoculated active zoospore after one generation approximates the number of zoospores produced per zoosporangium but may be an underestimate due to factors such as mortality of inoculated zoospores and the effect of density on the continued development of encysted zoospores (Longcore et al. 1999) (Fig. 3). Careful microscopy is needed to confirm the number of zoospores inside fully developed zoosporangia under various conditions in vitro and in vivo. Estimates in the literature range from 4 to 150 zoospores per zoosporangium (Berger et al. 1999, Annis et al. 2004). Here we estimate 65–161 zoospores per zoosporangium (Table 2).

We found that at increasing density, *B. dendrobatidis* grew faster (Fig. 3A) and matured faster (Fig. 3B). However, zoosporangia grew to a smaller size, and this may indicate that fewer zoospores are produced per zoosporangium at higher densities (Fig. 3C). The net effect of these changes could cause the rate of population growth on hosts to increase, decrease, or remain constant; more work is needed to ascertain how density dependence operates in populations of *B. dendrobatidis*.

**Cold-induced zoospore release**

data suggests that a cold shock could trigger disease emergence by temporally increasing the population growth rate of *B. dendrobatidis* and increasing infectivity. Large numbers of infectious zoospores were released when temperatures dropped. After growing to mature-sized thalli at 26.5°C, zoospores may have been quickly produced and released within a few days of the onset of cold conditions (Fig. 2C). This experiment indicates that the course of development and maturation is not set at the time of zoospore encystment, but changes with temperature. Cold shocks may also affect host behavior and immune function.

**Host behavior in the cold**

Although the amphibian chytrid is well adapted to a range of temperatures between 8°C and 25°C, host factors such as behavior and immune function may contribute to the development of fatal chytridiomycosis and disease-associated population declines at low temperatures. *Batrachochytrium dendrobatidis* has a direct life cycle and does not require an intermediate host (Longcore et al. 1999). An infected amphibian may shelter many generations of the pathogen, and pathogen spread over the host is perhaps facilitated in moist microhabitats and by low temperatures in which zoospores survive longer. Temperature strongly affects the activity patterns and microhabitat choices of ectotherms (Hutchison and Dupre 1992). The thermal environment of host amphibians affects the outcome of *B. dendrobatidis* infection (Woodhams et al. 2003, Berger et al. 2004). If high temperatures are available for basking, some amphibians may be able to inhibit fungal infections (Woodhams et al. 2003). Some behaviors of amphibians under cold conditions may increase their susceptibility to infection by *B. dendrobatidis*. At temperatures below 16°C, the tropical frogs *Litoria nannotis*, *L. rheocola*, and *Nyctimystes dayi* in Queensland, Australia, often shelter in streams or streamside refuges (Hodgkinson and Hero 2002, Retallick 2002). In Central America, association with aquatic habitat was a primary factor correlated with amphibian population declines (Lips et al. 2003) and may be linked to the emergence of chytridiomycosis (Lips et al. 2006). Many temperate amphibians and larvae overwinter in aquatic habitats, particularly at high elevations with short seasons for larval development (Pinder et al. 1992). Low temperatures concurrently affect host and pathogen, in this case, perhaps, to the advantage of *B. dendrobatidis*.

**Host immune suppression in the cold**

Research on ectotherms has shown that low temperatures can inhibit immune function (Bennett and Neville 1975, Cooper et al. 1992, Bly et al. 1993, Le Morvan et al. 1998, Carey et al. 1999, Jackson and Tinsley 2002). For some amphibians, short-term cold exposure as well as seasonal variability in temperature suppresses immune function. Maniero and Carey (1997) found that several immune defenses, including complement activity and lymphocyte proliferation, were suppressed in northern leopard frogs, *Rana pipiens*, by low temperatures. In African clawed frogs, *Xenopus laevis*, cold conditions increased the establishment of infection with the helminth parasite *Protopolyctoma xenopodis* (Jackson and Tinsley 2002). Red-spotted newts, *Notophthalmus viridescens*, decreased leukocyte production and lysozyme activity in the cold (Raffel et al. 2006). Wood frogs, *Rana sylvatica*, stopped synthesis of antimicrobial peptides at low temperatures (Matutte et al. 2000). Our data suggest that immune defenses that target the infective zoospore stage may be critical at low temperatures.

Innate epithelial defenses such as regulation of skin sloughing and mucosal barriers including antimicrobial peptides and commensal microbiota may influence the parameters modeled here, particularly zoospore survival and encystment rates. These defenses may inhibit *B. dendrobatidis* infections in some species (Davidson et al. 2003, Rollins-Smith and Conlon 2005, Harris et al. 2006, Woodhams et al. 2006, 2007) and are likely to be affected by temperature. For example, the microbiota of four-toed salamanders, *Hemidactylium scutatum*, include bacteria that inhibit the growth of oophagus fungus and *B. dendrobatidis* (Harris et al. 2006). However, microbiota may vary with environmental conditions, as shown in the large intestine of hibernating and nonhibernating *Rana pipiens* (Carr et al. 1976, Gossling et al. 1982, Banas et al. 1988). The effectiveness of these innate skin defenses also varies with species.

Several host factors may influence disease emergence, including suppression of host defenses and changes in the occurrence of risky host behavior. The biogeographical patterns of disease emergence may not be simply defined by characteristics of the pathogen. Although the amphibian chytrid is well adapted to a range of temperatures up to 25°C, host factors contribute to the disease-associated population declines at low temperatures.

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**Literature Cited**


APPENDIX

A description of the cold shock model (Ecological Archives E089-097-A1).