Integral Projection Models for host–parasite systems with an application to amphibian chytrid fungus

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Summary

1. Host–parasite models are typically constructed under either a microparasite or macroparasite paradigm. However, this has long been recognized as a false dichotomy because many infectious disease agents, including most fungal pathogens, have attributes of both microparasites and macroparasites.

2. We illustrate how Integral Projection Models (IPMs) provide a novel modelling framework to represent both types of pathogens. We build a simple host–parasite IPM that tracks both the number of susceptible and infected hosts and the distribution of parasite burdens in infected hosts.

3. The vital rate functions necessary to build IPMs for disease dynamics share many commonalities with classic micro and macroparasite models and we discuss how these functions can be parameterized to build a host–parasite IPM. We illustrate the utility of this IPM approach by modelling the temperature-dependent epizootic dynamics of amphibian chytrid fungus in Mountain yellow-legged frogs (Rana muscosa).

4. The host–parasite IPM can be applied to other diseases such as facial tumour disease in Tasmanian devils and white-nose syndrome in bats. Moreover, the host–parasite IPM can be easily extended to capture more complex disease dynamics and provides an exciting new frontier in modelling wildlife disease.

Key-words: Batrachochytrium dendrobatidis, devil facial tumour disease, fungal disease, macroparasite models, microparasite models, parasite aggregation, Rana muscosa, white-nose syndrome

Introduction

Following the influential papers by Anderson and May (Anderson & May 1979; May & Anderson 1979), host–parasite models have usually been constructed within one of two model structures. In their simplest form, microparasite models classify individuals as susceptible, infected or recovered, with the implicit assumption that all infected hosts can be considered similar because once a host is infected, microparasites can rapidly multiply within the host. Under this simple structure, prevalence, the proportion of infected individuals, is therefore adequate to characterize the level of infection within a host population. In contrast, macroparasite models generally assume that parasites cannot complete their entire life cycle within an individual host. Therefore, infection levels within a host are strongly influenced by the number of infective stages the host has encountered, and parasite burden influences host survival, reproduction and the transmission of infective stages. As a result, in macroparasite models, the proportion of individuals infected is not adequate to characterize the level of infection within a host population, and therefore, it is necessary to model the frequency distributions of parasites among individuals. In standard macroparasite models, the distribution of parasites among hosts is often modelled using a negative binomial distribution with a fixed aggregation parameter (Dobson & Hudson 1992).

In some pathogens traditionally categorized as microparasites, pathogen within-host reproduction occurs at a slow enough rate that it can be tracked from one time point to the next (Briggs, Knapp & Vredenburg 2010; Langwig et al. 2015a). In these instances, it is useful to take a macroparasite approach and model the distribution of loads across hosts as this measure is both more consistent with the type of data collected on these diseases and allows for the prediction of additional epidemiological patterns such as the patterns and dynamics of parasite aggregation (Scott 1987; Shaw & Dobson 1995). For example, fungal pathogens are increasingly recognized as important threats to biodiversity, agricultural production and human health (Fisher et al. 2012) and may exhibit this relatively slow, measurable on-host reproduction. A modelling framework that accounts for the microparasite and macroparasite characteristics of fungal pathogens is critical for understanding their dynamics.

To this end, Briggs, Knapp & Vredenburg (2010) developed an individual-based model for the fungal pathogen Batrachochytrium dendrobatidis in frog populations and were able to predict the biological criteria necessary for population persistence as well as the efficacy of different treatment strategies.
during epizootics (C.J. Briggs, unpublished data). However, this model required a separate equation for the fungal load on each individual and was difficult to parameterize from field or experimental data. In general, there is a need for an intermediate modelling framework for ‘slow’ microparasites that accounts for heterogeneity in the distribution of parasites across hosts (and how host distributional patterns change over time), while allowing for straightforward parameterization from laboratory or field data.

In this paper, we illustrate the potential for Integral Projection Models (IPMs) to address this need. Several recent papers have provided excellent overviews of the construction and use of IPMs (Rees & Ellner 2009; Coulson 2012; Metcalf et al. 2013; Merow et al. 2014a; Rees, Childs & Ellner 2014; Metcalf et al. 2016). In very general terms, IPMs assume that demographic parameters of individuals are affected by one or more continuous variables that describe some property of those individuals. The models then iterate population dynamics in discrete time with state variables of the form \( N(x, t) \), representing the frequency of individuals with continuous property \( x \) at the time \( t \).

The models can be readily parameterized from data using linear or nonlinear regression-based approaches. For population models, the continuous variable \( x \) is often the size, such as body mass (Coulson 2012), or age of an organism, but in principle any continuous variable or variables could be used. Here, we illustrate their use, using a measure of parasite load. It has been pointed out that this approach be used. Here, we illustrate their use, using a measure of parasite load. It has been pointed out that this approach be used. Here, we illustrate their use, using a measure of parasite load. It has been pointed out that this approach be used. Here, we illustrate their use, using a measure of parasite load. It has been pointed out that this approach be used. Here, we illustrate their use, using a measure of parasite load. It has been pointed out that this approach be used. Here, we illustrate their use, using a measure of parasite load. It has been pointed out that this approach be used. Here, we illustrate their use, using a measure of parasite load. It has been pointed out that this approach be used. Here, we illustrate their use, using a measure of parasite load. It has been pointed out that this approach be used. Here, we illustrate their use, using a measure of parasite load. It has been pointed out that this approach be used. Here, we illustrate their use, using a measure of parasite load.

The basic model we examine is a modification of a susceptible-infected-susceptible model. In our model, individuals that clear the infection immediately re-enter the susceptible class, with no immunity. Including a recovered class simply requires adding an additional discrete stage to the IPM (Metcalf et al. 2016). The model has the following state variables:

\[
S(t): \text{Number of susceptible/uninfected hosts at time } t \\
I(x, t): \text{Frequency of infected hosts with load } x \text{ at time } t \text{ (where } x \neq 0); \\
R(x, t): \text{Frequency of infected hosts with load } x \text{ at time } t \text{ (where } x \neq 0); \\
U(x, t): \text{Frequency of infected hosts with load } x \text{ at time } t \text{ (where } x \neq 0); \\
L(x, t): \text{Frequency of infected hosts with load } x \text{ at time } t \text{ (where } x \neq 0); \\
S(t + 1) = S(t)\phi(L(x, t)) + \int_{L}^{U} f(x)I(x, t)s(x)L(x)dx + \int_{L}^{U} f(x)I(x, t)R(x, t)dx.
\]

Materials and methods

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\]

\[
\text{eqn 1}
\]

Fig. 1. Life history flow chart for the host–parasite Integral Projection Model. The chart shows how an infected host with a parasite load of \( x \) at time \( t \) can transition to an infected host or susceptible/uninfected host with a parasite load of \( x' \) or 0, respectively, at time \( t + 1 \). The chart also shows how a susceptible/uninfected host at time \( t \) can transition to an infected host or susceptible/uninfected host with a parasite load of \( x' \) or 0, respectively, at time \( t + 1 \).
The first term in this equation gives the number of hosts who remain uninfected in a time step. The second term gives the number of infected hosts who lose an infection and enter the uninfected class in a time step. The third and fourth terms give the number of uninfected hosts who are born from uninfected and infected hosts in a time step.

The equation for infected hosts with load \( x' \) at time \( t + 1 \) is given by

\[
I(x', t + 1) = \int_{L}^{U} I(x, t) s(x) (1 - f(x)) G(x', x) dx + S(t) \phi(I(x, t)) G_{0}(x').
\]

The first term in this equation gives the number of infected individuals with load \( x \) that transition to load \( x' \) in a time step. The second term gives the number of uninfected individuals that transition to an infected individual with load \( x' \) in a time step. Below we more thoroughly discuss the terms in eqns (1) and (2), how they relate to classic macroparasite and microparasite models, and how they can be parameterized. When parameterizing the functions below, we assume that each process obeys the Markov property such that only the load at time \( t \) predicts the event at time \( t + 1 \) (e.g. growth of the parasite, host survival, loss of infection, etc.; Easterling, Ellner & Dixon 2000).

**THE GROWTH FUNCTION: \( G(x', x) \)**

For continuous measures of parasite load, the growth function \( G(x', x) \) specifies the probability density of transitioning to an infected load \( x' \) at time \( t + 1 \), dependent on having a load of \( x \) at time \( t \). In comparison with standard macroparasite and microparasite models, this function allows for pathogen growth on a host to be driven by both within-host pathogen birth/rapid self-reinfection (e.g. microparasites and some macroparasites) and from acquiring additional parasites from the environment or other infected hosts. The dependence of \( G(x', x) \) on the free-living stages of the parasites can be made explicit by writing \( G(x', x) \) as dependent on the number of free-living parasites at time \( t \).

This function can be estimated with data on the parasite load of individual hosts at time \( t \) and time \( t + 1 \). Using standard regression techniques, load at time \( t \), the number of free-living parasites at time \( t \) and/or the density and abundance of other infected hosts can be regressed against load at time \( t + 1 \) and the resulting model can be used to parameterize the growth function \( G(x', x) \) (Easterling, Ellner & Dixon 2000). For continuous parasite loads, the load at time \( t + 1 \) could be described by a log-normal or gamma distribution, while discrete disease loads could be fit by a negative binomial distribution (Anderson & May 1978; Shaw, Grenfell & Dobson 1998). The growth of a parasite on a host will often depend on other abiotic variables that can be accounted for as additional fixed or random effects in the regression model (Rees & Ellner 2009).

**THE SURVIVAL FUNCTION: \( s(x) \) and \( s_{0}(x) \)**

\( s_{0} \) specifies the survival probability of uninfected hosts. \( 1 - s_{0} \) gives the probability of a host dying without any infection, which parallels the death rate of uninfected hosts in classic micro and macroparasite models. \( s_{0} \) can be estimated by the proportion of uninfected hosts that survive from \( t \) to \( t + 1 \).

The survival function \( s(x) \) specifies the probability of a host with a parasite load \( x \) surviving from time \( t \) to time \( t + 1 \). In classic macroparasite models, it is assumed that parasite-induced host mortality increases linearly with load at rate \( \alpha \), where \( \alpha \) specifies the pathogenicity of the parasite (Anderson & May 1978). In the IPM framework, a commonly used function to measure survival probability is the logistic function given by

\[
s(x) = \frac{\exp(b_{0} - b_{1}x)}{1 + \exp(b_{0} - b_{1}x)}.
\]

where \( b_{1} \) is similar to the pathogenicity parameter \( \alpha \) (Anderson & May 1978; Wilber, Weinstein & Briggs 2016). When \( b_{0} \) is held constant, \( b_{0} \) dictates the parasite load at which substantial parasite-induced host mortality begins to occur (Wilber, Weinstein & Briggs 2016). The logistic function could be replaced by other functions, as dictated by the data (Dahlgren, García & Ehrlen 2011).

The survival function \( s(x) \) can be estimated with logistic regression using host survival and load data from laboratory or mark–recapture studies conducted at the appropriate time-scale. If other biotic or abiotic factors are also thought to contribute to the survival probability of a host from \( t \) to \( t + 1 \), they can be included as additional predictor variables in the survival function.

**THE LOSS OF INFECTION FUNCTION: \( l(x) \)**

The loss of infection function \( l(x) \) specifies the probability of a host having a parasite load of \( x \) at time \( t \) and losing the infection by time \( t + 1 \). In comparison with classic microparasite susceptible-infected models, this function is analogous to the rate at which infected individuals recover from infection and reenter the susceptible/uninfected class. We similarly assume that individuals that lose an infection immediately reenter the susceptible/uninfected class, though a resistant class could easily be included in this modelling framework (Metcalfe et al. 2016).

A logistic function (eqn 3) could also be used for the loss of infection function and could be parameterized using parasite load data at time \( t \) and \( t + 1 \) and fitting a logistic regression where the response variable is whether or not a host lost an infection by time \( t + 1 \) and the predictor variable is the infection intensity \( x \) at time \( t \). As with the survival function, if other biotic or abiotic factors also contributed to \( l(x) \) they could be included as additional predictor variables in the logistic regression.

**THE TRANSMISSION FUNCTION: \( \phi(I(x, t)) \)**

The transmission function \( \phi(I(x, t)) \) specifies the probability of transitioning from the uninfected class to the infected class. The transmission function is critically important for the dynamics of a disease and can take a variety of different functional forms. Some common examples include the density-dependent or mass action transmission function \( \beta S/N \) and the frequency-dependent transmission function \( \beta S/I/N \) (McCallum, Barlow & Hone 2001).

Over a unit time interval, a density-dependent, mass action transmission function results in the following probability of an individual host not being infected: \( \exp \left( -\beta \int_{L}^{U} h(x, t) dx \right) \), where \( \int_{L}^{U} h(x, t) dx \) gives the total number of infected individuals. Allowing \( \beta \) to be a function of parasite load, \( \phi(I(x, t)) \) can be written as

\[
\phi(I(x, t)) = 1 - \exp \left( -\int_{L}^{U} \beta(x) h(x, t) dx \right),
\]

where \( \int_{L}^{U} \beta(x) h(x, t) dx \) is the force of infection and \( \beta(x) \) specifies the effect of an individual with an infection load of \( x \) on the infection probability of an uninfected individual. This formulation assumes that new infections occur following a Poisson process with rate \( \int_{L}^{U} \beta(x) h(x, t) dx \).

While this functional form may be appropriate for many microparasites in which direct transmission among hosts is the primary mode of acquiring infection, the transmission of some pathogens depends on the number of free-living parasites in a system as well as the number of
infected hosts (Briggs, Knapp & Vredenburg 2010). If we assume that number of free-living parasites is proportional to the total number of parasites in all infected hosts in the system at time \( t \), then we can modify \( \Psi(x, t) \) to \( \Phi(x, \Psi(x, t)) \) to capture this biology.

Finally, some pathogens have an environmental reservoir such that the probability of infection is nonzero even when no infected hosts are present. This could be captured by rewriting eqn (4) as

\[
\Phi(I(x, t)) = \exp \left( -\left( a + \int_{0}^{t} \beta(x)I(x, t)\,dx \right) \right),
\]

where \( 1 - \exp(-a) \) defines the probability of infection when no infected hosts are present (e.g. from an environmental reservoir). This environmental reservoir could be more explicitly accounted for by including an additional state variable in the IPM that tracks how the number of parasites in the environment grows and decays in a time step (Rohani et al. 2009).

Methods for estimating the transmission function and/or its corresponding parameters are well-described in the host–pathogen literature (McCallum 2000; Smith et al. 2009), though choosing between transmission functions is typically a data-intensive procedure (Rachowicz & Briggs 2007; Smith et al. 2009).

THE INITIAL INFECTION BURDEN FUNCTION: \( G_b(x') \)

The function \( G_b(x') \) specifies the probability density of the infection intensity of a host when it first becomes infected and can be a function of the total number of infected hosts in the population, the total number of infectious agents in the population and/or various other host or abiotic covariates. This function can be estimated by fitting a regression model where the response variable is the pathogen load of infected hosts at time \( t + 1 \) that were uninfected at time \( t \). For continuous disease loads, a variety of different distributions such as gamma, log-normal and normal could be explored.

In comparison with standard stochastic macroparasite models, this function is analogous to clumped infection distributions in which a host can acquire a random number of free-living parasites in a small time step (Isham 1995; Pugliese, Rosà & Damaggio 1998). However, depending on the time step used to parameterize the IPM, \( G_b(x') \) will also be influenced by the growth of the parasite on the host as a ‘clump’ of parasites can infect a host and then grow in the time interval \( t \) to \( t + 1 \).

Moreover, the above host–parasite IPM assumes that after acquiring an initial infection burden the growth of the parasite on a host is then driven by \( G(x', x) \) and is independent of the density of infected hosts. If one has reason to believe that transmission and the function \( G_b(x') \) are important drivers of disease dynamics on hosts after the initial infection, the growth function may be redefined as

\[
G'(x', x, I(x, t)) = \left( 1 - \Phi(I(x, t))G(x', x) + \Phi(I(x, t)) \left[ G_b(x' - x) + \text{higher order terms} \right] \right),
\]

where an increase in load from \( x \) to \( x' \) in a time step could be because of (i) no transmission occurring and parasite load increasing due to within-host pathogen growth (first term) or (ii) transmission occurring and a host acquiring a ‘clump’ of infections of size \( y \) such that \( y = x' - x \) (second term) or (iii) some combination of both within-host growth and transmission occurring such that parasite load increases from \( x \) to \( x' \) in a time step. This is given by higher order terms and will depend on the length of the time step \( t \) to \( t + 1 \) relative to the dynamics of the pathogen.

In Appendix S1, Supporting information, we derive \( R_0 \), a canonical epidemiological measure of the ability of a parasite to invade a fully susceptible host population (Diekmann, Heesterbeek & Metz 1990), for the host–parasite IPM and show how it depends on the vital rate functions \( G(x', x), s(x), R(x), \Phi(I(x, t)), \) and \( G_i(x') \).

THE FECUNDITY FUNCTION: \( f_0 \) and \( f(x) \)

The fecundity function \( f(x) \) specifies the mean number of offspring produced by individuals with a parasite load of \( x \) (or by susceptible/uninfected individuals \( f_0 \)) and the host–parasite IPM assumes that all offspring enter the uninfected class. It is easy to relax this assumption and include vertical transmission into the host–parasite IPM by allowing newly born hosts to enter the infected class with a parasite load specified by some probability density function. Standard macroparasite models assume that host reproduction decreases linearly with increasing parasite load (May & Anderson 1978). However, as pointed out by May & Anderson, this is an over simplification as the response of host reproductive effort to parasitism is often nonlinear (e.g. Weatherly 1971) and reproduction itself can never take on a negative value (Roberts, Smith & Grenfell 1995). Alternative formulations of parasite-induced reduction in host fertility that account for this nonlinear relationship have been discussed (Roberts, Smith & Grenfell 1995).

In the IPM framework, the fecundity function can be fit using Poisson or negative binomial regression where the predictor variable is parasite load and the response variable is the number of offspring produced by a host with that parasite load (Easterling, Elnner & Dixon 2000). If the response variable is a non-integer value, a continuous distribution such as gamma or log-normal could be used. Depending on the link function, these regressions can produce nonlinear fecundity functions that are always positive. Similar to the other vital functions discussed above, other factors that affect mean reproductive output can be included in the regression.

For many host–parasite systems, host reproduction occurs on a much longer time-scale than the dynamics of the parasite and it may not be biologically realistic to include host reproduction at each time step in the IPM model as is done in eqn (1). For example, if host reproduction occurs at a particular time during the year it may be useful to break the year into separate IPMs (e.g. an IPM for summer, fall, winter and spring; Caswell 2001) such that load-dependent host reproduction only occurs in a particular season or as a discrete pulse at the beginning of a particular season (e.g. host reproduction is only nonzero in the spring and fall). One may also want to include host age as an additional discrete or continuous host attribute (Childs et al. 2003) to account for reproductive differences among hosts of different ages. On the other hand, if one is particularly interested in the fate of a host population over a single seasonal epizootic where host reproduction does not occur, the fecundity function may be excluded from the host–parasite IPM as it will not affect host population persistence during the epizootic. In this case, appropriately modelling vital rates such as the survival function \( s(x) \) and the growth function \( G(x', x) \) will be critically important for understanding host–parasite dynamics. In general, how to include host reproduction into the host–parasite IPM will depend on the questions that are being addressed.

APPLICATION OF MODEL TO AMPHIBIAN CHYTRID FUNGUS: LABORATORY EXPERIMENT

We use the above IPM framework to examine the population dynamics of amphibian hosts infected with the fungal pathogen *Batrachochytrium dendrobatidis* (*Bd*). *Bd* is a devastating amphibian
pathogen that has led to declines in many amphibian populations around the globe (Skerratt et al. 2007; Kilpatrick, Briggs & Daszak 2010). *Bd* is a cutaneous fungus that disrupts the osmoregulatory ability of amphibian skin, eventually leading to chytridiomycosis and amphibian mortality (Voyles, et al. 2007, 2009). In contrast to traditional macroparasites, *Bd* reinfests an infected host (Rollins-Smith 2009). The generation time of *Bd* is between 4-10 days depending on temperature (Woodhams et al. 2008), such that the on-host *Bd* growth dynamics can be captured via repeated swabbing of an animal every few days, with the fungal load on the frog estimated as the number of copies of *Bd* DNA detected on the skin swabs via quantitative PCR (Boyle et al. 2004). Quantitative PCR provides a continuous measure of infection intensity between 0 (uninfected) and an arbitrarily large *Bd* infection. These characteristics of *Bd* make it an ideal candidate for applying the host–parasite IPM described above.

We use the IPM framework to gain insight into how temperature affects the epizootic dynamics of *Bd* in populations of the Mountain yellow-legged frog complex (*Rana muscosa* and *Rana sierrae*, henceforth *R. muscosa*). *Rana muscosa* are native to the California Sierra Nevada mountains and have suffered severe *Bd*-induced population declines (Briggs, Knapp & Vredenburg 2010; Vredenburg et al. 2010). The severity of *Bd* infection is highly temperature-dependent (Berger et al. 2004; Andre, Parker & Briggs 2008), with optimal *Bd* growth occurring between 17–25 °C in laboratory conditions (Piotrowski, Annis & Longcore 2004), but depending on-host *Bd* interactions (Piotrowski, Annis & Longcore 2004; Raiffel et al. 2012). While these are the temperatures at which amphibians often suffer more severe chytridiomycosis and mortality, this pathology is species-dependent (Kilpatrick, Briggs & Daszak 2010).

We use data from a laboratory experiment in which 20 adult *R. muscosa* were housed separately at three different temperatures (4 °C, 12 °C, 20 °C; 5 frogs per temperature), exposed to *c.* 10^6 zoospores of *Bd* and then monitored for 119 days. Every 3 days starting 8 days after exposing the frogs to *Bd*, the frogs were swabbed and *Bd* zoospore load was estimated using quantitative PCR. Mortality that occurred between swabbing events was recorded at the next swabbing event.

### MODEL DESCRIPTION

To fit the IPM to *Bd* load data from laboratory experiments, we made two simplifying assumptions. First, we excluded reproduction/recruitment because we lack data on the effect of infection on reproduction. As a result, we used this model to address questions regarding epizootic dynamics of *Bd* and *R. muscosa* over the course of a single summer season, rather than to examine long-term population persistence with disease.

Secondly, we assumed the probability of infection *ϕ(T)* was temperature (*T*)-dependent, but independent of the density of infected hosts (i.e. *I(x,t)*) does not affect the probability of infection. In our experiments, individual animals were housed in separate containers and initial infection was solely due to an amphibian acquiring *Bd* zoospores from the environment. We subsequently explored different transmission functions that do include *I(x,t)* to understand their implications on *Bd* epizootic dynamics. With these assumptions, the modified IPM is given by

\[ S(t+1) = S(t)S_0(T)(1 - \phi(T)) + \int_{L}^{U} I(x,t)x I(x,t) f(x,T) dx, \]

**eqn 7**

where the various vital functions are now dependent on temperature *T*. Note that *x* refers to ln(χ) (log zoospore load) when *x* = 0. In this case, susceptible/uninfected represents a discrete state of the frog and is not equivalent to ln(χ)=0. In this model, a single time step represents 3 days, which was the time between swabbing events in the laboratory experiment.

### VITAL RATE FUNCTIONS

We modelled the survival function *s(x)* of a frog with a log zoospore load of *x* as a logistic regression with the link function given by

\[ \logit(s(x)) = b_{0,0} + b_{1,0}x, \]

**eqn 9**

where *b_{0,0}* is the intercept of the link function on the logit scale, and *b_{1,0}* is the effect of log zoospore load on the logit-transformed probability of survival. We could not estimate the effect of temperature on this vital rate function because no individuals died at 4 or 12 °C (Fig. 2a). This result was surprising because individuals at 12 °C had loads as high or higher than individuals at 20 °C and did not experience mortality. Based on previous results in the field which show that *R. muscosa* suffer a roughly consistent *Bd*-induced mortality across a variable lake temperatures in the field (i.e. the ≈10,000 zoospore threshold, Vredenburg et al. 2010), additional results in the laboratory that show that frogs experience significant *Bd*-induced mortality at temperatures below 20 °C (17 °C, Andre, Parker & Briggs 2008), and extensive field observations that decreased temperature does not have a large protective effect on *R. muscosa* in the field (Knapp et al. 2011), we think there is very little evidence that the survival curve of *R. muscosa* and *Bd*-load interacts with temperature. Therefore, we assumed that *Bd*-induced mortality is dependent only on load and not on temperature directly. We parameterized the survival function using only individuals at 20 °C (Fig. 2a, see Appendix S2 for a comparison with a survival function fitted with all of the temperature data) and assumed a temperature-independent survival function. However, temperature influenced fungal growth, as detailed next.

We modelled the growth function *G(x',x)* as a normal distribution

\[ X \sim N(\mu(x,T), \sigma^2(x)) \]

where *T* is temperature. Mean fungal loads were modelled as

\[ \mu(x,T) = b_{0,1} + b_{1,1}x + b_{2,1}T, \]

**eqn 10**

where *b_{0,1}* is the intercept and *b_{1,1}* and *b_{2,1}* give the effect of a unit change in log zoospore load and temperature on the log zoospore load at time *t* + 1, respectively. We also allowed the variance of *G(x',x)* to be an exponential function of log zoospore load at time *t*

\[ \sigma^2(x) = \sigma_{0,1} \exp(2\sigma_{0,1}x), \]

**eqn 11**

where *σ_{0,1}* is a constant and *σ_{0,1}* dictates the effect of log zoospore load on the variance.

We modelled the loss of infection function *l(x)* as a logistic regression with the link function

\[ \logit(l(x,T)) = b_{0,2} + b_{1,2}x + b_{2,2}T, \]

**eqn 12**

where *b_{0,2}* is the intercept and *b_{1,2}* and *b_{2,2}* are the coefficients giving the effect of a unit change in log zoospore load and temperature on the logit-transformed probability of losing an infection in a single time step, respectively.

We modelled the initial infection burden function $G_0(x')$ as a normal distribution $X \sim N(\mu(T), \sigma^2(T))$ because infection burden was modelled on the log scale, which allowed for negative values. We defined the mean of the distribution as $\mu(T) = b_{0,3} + b_{1,3}T$ where $b_{0,3} \text{ and } b_{1,3}$ are defined similarly to the growth function. We modelled the variance as $\sigma^2(T) = v_{0,3} \exp(2c_{0,3}T)$ where $v_{0,3}$ and $c_{0,3}$ are defined similarly as in the growth function.

Finally, we modelled the probability of an individual becoming infected $\phi(T)$ in a time step as a function of temperature $T$ using a logistic model $\logit[\phi(T)] = b_{0,4} + b_{1,4}T$ where $b_{0,4}$ and $b_{1,4}$ are defined similar to the recovery function. We performed model selection and validation for each vital rate function described above and these results are given in Appendix S2. We fit the vital rate functions in R version 3.1.2 (R Foundation for Statistical Computing, Vienna, Austria) and all code used for this analysis can be found at http://github.com/mqwilber/ipm_for_parasites.

**ANALYSING THE IPM**

After fitting the parameters of the vital rate functions, we analysed the resulting IPM (eqns 7 and 8) by discretizing the continuous variable $Bd$ load and using the mid-point rule to evaluate the IPM at each time step (Rees, Childs & Ellner 2014). For the infected portion of the host–parasite IPM, we used 100 discretized bins (i.e. a mesh size of 100) and lower and upper bounds of $-5$ and $18$ log zoospore load, which we chose to minimize the effects of evocation on the IPM predictions (loss of individuals from the model because their predicted future loads are outside the model range, Appendix S3; Williams, Miller & Ellner 2012). To put these bounds in context, the log zoospore range from our experiment was $(-1.14, 13.15)$ and the approximate log zoospore load at which $R. \text{ muscosa}$ begin experiencing substantial die-off in the field is at or above a log zoospore load of 9.21 (Vredenburg et al. 2010). To incorporate the discrete, uninfected stage into the IPM, we appended an extra row giving transitions of various infected individuals to an uninfected state (top-most row) and an extra column specifying the transition of uninfected individuals into various infected states (left-most column) to the $100 \times 100$ parasite load transition matrix described above (Merow et al. 2014a).

We calculated the local elasticity of the population growth rate ($\lambda$) to the lower-level regression parameters $b_{ij}$ of the vital functions defined above by perturbing each regression parameter by $\delta = 0.001$ and calculating the elasticity as $e_{ij} = (\lambda_{\text{perturbed}} - \lambda_{\text{stable}})/\delta \times b_{ij} / \lambda_{\text{stable}}$ (Merow et al. 2014b). To propagate the uncertainty in our estimates of the lower-level vital rate parameters through to our estimates of the population growth rate and lower-level parameter elasticity, we took the following parametric bootstrap approach. Using standard asymptotic likelihood results (McCullagh & Nelder 1989), we assumed that each parameter set from a vital rate function followed a multivariate normal distribution with a mean and variance–covariance matrix equal to the values given by the regression procedure used to fit the vital rate
function. Next, we ran 1000 simulations in which we randomly drew the lower-level regression parameters from their respective multivariate normal distributions and parameterized the IPM using these parameters. We then calculated either the population growth rate or the elasticity of a given lower-level parameter with these randomly drawn values and stored the result. This provided us with estimates of the population growth rate and lower-level parameter elasticity while accounting for the uncertainty in the lower-level parameters used to build the IPM. We note that this approach likely underestimates the uncertainty as it does not account for the uncertainty in the variance estimates, does not account for covariance of parameters between vital rate functions, and assumes multivariate normality.

**Experiencing Density-Dependent Transmission Dynamics**

In eqn (7), in order to match the conditions of our laboratory experiment in which animals were housed singly, we assumed transmission of *Bd* did not depend on the density or frequency of infected hosts. Here, we explored how a mass action, density-dependent transmission function affects the epizootic dynamics of *Bd*. In particular, we assumed the following transmission function

\[ \phi(I(x, t)) = 1 - \exp \left( -\left( a + \beta \int_{0}^{t} x(t, \tau) \, d\tau \right) \right) \]  

\[ = 1 - \exp \left( -\left( a + \beta \int_{0}^{t} x(t, \tau) \, d\tau \right) \right), \quad \text{eqn 13} \]

which specifies that the probability of infection at time *t* is dependent on the total number of zoospores present in the host population \( \int_{0}^{t} x(t, \tau) \, d\tau \) at time *t* as well as a constant probability of infection from an environmental reservoir \( \omega = 1 - \exp(-a) \). We followed the example of previous *Bd* modelling work and assumed that the *Bd* epizootic dynamics depend on the number zoospores in the aquatic environment rather than just the number of infected amphibians in a population (Briggs, Knapp & Vredenburg 2010). The term \( \int_{0}^{t} x(t, \tau) \, d\tau \) reflects this assumption, albeit ignoring potential dynamics of free-living zoospores. Moreover, we assumed that density dependence affects only the probability of transitioning from uninfected to infected, such that once an amphibian is infected the increase in *Bd* is independent of infected host density. This assumption is realistic if the parasite reproduction on the host swamps out the effects of reinfection from other individuals or the environment.

To explore the effects of this transmission function on epizootic dynamics, we first parameterized the density-independent portion of the IPM model using the maximum likelihood estimates of the vital rate function parameters discussed above. Because we could not estimate the density-dependent transmission function from the data we collected, we explored the effect of this function on population dynamics by choosing \( (\omega, \beta) \) pairs on a grid and using these values to parameterize the density-dependent transmission function. The estimated values of the environmental infection probability \( \omega \) used in the density-independent model suggested that \( \omega \) was between \( 0.22 \) and \( 0.6 \) depending on the temperature, so we explored values of \( \omega \) between \( 0.01 \) and \( 0.6 \). We did not have a good \textit{a priori} estimate of \( \beta \), so we explored \( \beta \) within the range \( 0 \) to \( 1 \times 10^{-2} \), where this upper bound was chosen arbitrarily after preliminary simulations showed that larger values of \( \beta \) had little effect on the population dynamics. For every \( (\omega, \beta) \) pair, we iterated the density-dependent IPM for 120 days, which is the approximate length of the summer in the Sierra Nevada during which *Bd* epizootics tend to occur (Briggs et al. 2005; Briggs, Knapp & Vredenburg 2010). We initialized each population with 100 uninfected individuals, and for each combination of \( (\omega, \beta) \), we calculated the proportion of surviving amphibians and the prevalence of *Bd* infection at the end of the epizootic.

**Results**

**VITAL RATE FUNCTIONS**

Increasing log zoospore load \( x \) significantly decreased the survival probability of amphibians (Fig. 2a; \( \chi_{21}^{2} = 12.197, P = 0.0005 \); Table 1).

Both temperature and log zoospore load at time *t* significantly increased log zoospore load at time \( t + 1 \) (Likelihood ratio test (LRT) for load at time *t*: \( \chi_{21}^{2} = 196.36, P < 0.0001 \); LRT for temperature: \( \chi_{21}^{2} = 13.56, P = 0.0002 \). Moreover, log zoospore load at time *t* was important for describing the variance structure of the growth function, as compared to a model with constant variance structure (LRT comparing full model to model with constant variance: \( \chi_{21}^{2} = 14.555, P = 0.0001 \); log zoospore load: \( \chi_{21}^{2} = 23.701, P < 0.0001 \); Fig. 3). Amphibians were more likely to clear infection at lower temperatures and when the load at time *t* was smaller.

Increasing temperature significantly increased the mean and variance of the initial infection load distribution \( G_{0}(x') \) (temperature effect on mean: \( \chi_{21}^{2} = 2.53, P = 0.015 \); temperature effect on variance: LRT comparing model with variance structure to without: \( \chi_{21}^{2} = 6.00, P = 0.0143 \); Fig. 3).

Finally, increasing temperature significantly increased the probability of infection \( \phi \) (\( \chi_{21}^{2} = 6.0361, P = 0.014 \); Table 1).

**LABORATORY DYNAMICS OF AMPHIBIANS AND BD**

The parameterized IPM model predicted that individual amphibians at low temperatures would survive significantly longer than amphibians at high temperatures, with the largest difference being when log zoospore loads were low (Fig. 4a). Over a summer epizootic, amphibian populations at low temperatures experienced a minimal effect of *Bd*-induced population declines (\( \lambda \approx 1 \)), while amphibians at higher temperatures experience substantially more rapid declines, with large uncertainty around these estimates (Fig. 4b). Elasticity analysis on the lower-level parameters used in the vital rate functions showed that overall population growth rate was most sensitive to proportional changes in the growth rate of *Bd* (the parameters of the growth function; Fig. S5) as well as the pathogenicity of *Bd* and the threshold at which *Bd*-induced mortality began to occur (the parameters of the survival function; Fig. S5).

The IPM model also allowed us to examine how the stable log zoospore distribution of *Bd* on surviving hosts changed with temperature. For surviving, infected amphibians, the mean infection intensity increased with temperature, but the variance to mean ratio decreased with temperature (Fig. 5).
consistent with experimental and model results showing that hosts experienced greater Bd-induced mortality at higher temperatures. This is also consistent with previous theoretical results from macroparasite models which predict that increased parasite-induced host mortality generally decreases the variance to mean ratio (Barbour & Pugliese 2000).

### Effects of Density-Dependent Transmission on Epizootic Dynamics

The effect of density-dependent transmission on *Bd*-R. *muscosa* population dynamics varied with temperature, the probability of infection from the environmental reservoir ($\omega$) and the transmission coefficient ($\beta$). In general, over the range of density-dependent transmission values we examined, density-dependent transmission had little effect on prevalence and the proportion of population decline over the course of a summer epizootic (Figs S7 and S8). In contrast, the probability of infection from the environment had a large effect on both prevalence patterns and population decline (Figs S7 and S8). Given a probability of infection from the environment above approximately 0.15, increasing density-dependent transmission had very little effect on *Bd* prevalence or *R. muscosa* population decline. Over the parameter space we examined, the density-dependent transmission model predicted that populations at 12 °C will experience a maximum of a 20% population decline over the course of an epizootic with 70% prevalence, while populations at 20 °C will experience a >80% population decline with close to 100% prevalence (Figs S7 and S8).

### Discussion

Integral Projection Models provide an ideal framework to model diseases that do not fall neatly into the microparasite/macroparasite dichotomy and a way to explicitly model heterogeneity and changes in the pathogen load distribution in the host population. By taking an intermediate approach between individual-based disease models which explicitly track the parasite load on every individual in a population (Briggs, Knapp & Vredenburg 2010) and classic macroparasite/microparasite models which only track the total number of hosts and parasites in a population (Anderson & May 1978), IPMs can elegantly investigate population outcomes of infectious diseases while still incorporating critical information about disease dynamics at the individual-level (Metz et al. 2016). While the IPM approach can theoretically be used to explore the dynamics in any macroparasite or microparasite system, we believe it will be especially useful in host–parasite systems where the growth rate of a parasite is slow enough that measurements of parasite load at time $t$ and $t+1$ are on the same time-scale as the growth rate of the parasite. This allows for empirical estimation of the vital rate functions and an investigation regarding how these vital rate functions vary with environmental factors such as temperature and/or differ between host populations in which a disease is established or invading. This approach could also be used to explore how the distribution of parasite load among individuals changes over time, without the constraints imposed by choosing a fixed distribution and/or aggregation parameter.

We used the host–parasite IPM model to explore the consequences of different temperatures on *R. muscosa–Bd* dynamics over the course of an epizootic. The effect of temperature on *Bd* growth is well-known both in culture and on amphibian hosts (Longcore et al. 1999; Berger et al. 2004; Piotrowski, Annis & Longcore 2004; Andre, Parker & Briggs 2008; Raffel et al. 2012) and previous work has estimated the expected time to death of amphianbs infected with *Bd* over various different temperatures (Berger et al. 2004; Andre, Parker & Briggs 2008). However, the effect of temperature-*Bd* interactions on amphibians at the population level is much less clear (Rohr & Raffel 2010; Knapp et al. 2011). Using an IPM model, we were
able to make specific, quantitative predictions about how temperature and transmission dynamics affect population growth rates of *Rana muscosa*.

The density-independent IPM model predicted that population-level growth rate decreased with increasing temperature and naïve populations at or above about 18 °C had a 50% chance of experiencing an 80% decline or greater over the course of a summer epizootic. This result likely represents a best case scenario for *Rana muscosa* as this density-independent model does not account for *Bd* transmission dynamics (Rachowicz & Briggs 2007) or additional factors leading to increased frog mortality or *Bd*-susceptibility in the field. Our elasticity analysis showed that the population-level growth rate was most sensitive to proportional changes in parameters relating to the *Bd* growth function and the survival function. *In situ* factors slightly reduced the *Bd*-load at which frogs began experiencing disease-induced mortality, for example, *R. muscosa* populations could experience extirpation during a summer epizootic for a wide range of temperatures, which would be consistent with the patterns observed in the field (Knapp et al. 2011). In particular, we assumed a temperature-independent survival function in the IPM model (described in ‘Vital Rate Functions’) and including temperature dependence into this function would have significant impacts on the ability of *R. muscosa* populations to persist through an epizootic.

We extended this density-independent IPM to explore how density-dependent transmission and transmission from an environmental reservoir affected population dynamics. Our results suggest that density-dependent transmission had a small effect on the population dynamics of *Bd* epizootics, particularly when an environmental reservoir was present. While this result is largely due to our assumption that density-dependent transmission does not affect the growth of *Bd* on an
already infected frog, it is consistent with predictions from a fully individual-based model that predicts that density manipulations (i.e. culling infected frogs) will likely have little effect on mitigating population outcomes during \(Bd\) epizootics in this system (C.J. Briggs, unpublished data). A natural next step will be to use this IPM to investigate how varying temperature regimes and \(R.\) muscosa demography affect the persistence of \(R.\) muscosa populations infected with \(Bd\) over longer timescales. In general, the question of how temperature interacts with \(Bd\) and in turn affects amphibian host persistence is a critical question in amphibian conservation (Roehr & Raffel 2010) and IPMs provide a novel means by which this question can be quantitatively addressed.

In addition to these population-level predictions, host–parasite IPMs also allow for explicit predictions about how the distribution of parasites loads over hosts changes with different vital parameters and/or over the course of an epizootic or enzootic. Macroparasite models have long recognized the importance of the distribution of parasite loads over hosts for determining the dynamics of host–parasite interactions (Anderson & May 1978; Tompkins et al. 2002), and classic macroparasite models addressed this by using a statistical distribution (often negative binomial, Shaw, Grenfell & Dobson 1998) and then looking at how different levels of parasite aggregation affected host–parasite dynamics (Anderson & May 1978; Kretzschmar & Alder 1993). These approaches have been extended to include fluctuating aggregation (Rosá & Pugliese 2002; Rosá et al. 2003), but still rely on explicitly defining the shape of the host–parasite distribution. In contrast, IPMs do not assume a host–parasite distribution, rather one emerges as a result of the vital functions specified when parameterizing the model and the transmission and population dynamics over time. Therefore, one can explore how sensitive the aggregation of the host–parasite distribution is to different vital function parameters, providing an intriguing way to parse the contribution of different processes to parasite aggregation. Moreover, as it is straightforward to include fluctuating aggregation (Rosá & Pugliese 2002; Rosá et al. 2003), but still rely on explicitly defining the shape of the host–parasite distribution. In contrast, IPMs do not assume a host–parasite distribution, rather one emerges as a result of the vital functions specified when parameterizing the model and the transmission and population dynamics over time. Therefore, one can explore how sensitive the aggregation of the host–parasite distribution is to different vital function parameters, providing an intriguing way to parse the contribution of different processes to parasite aggregation.

Using the parameterized IPM for \(Bd\)-\(R.\) muscosa, we examined how the distribution of \(Bd\)-loads changed with temperature. The IPM showed that fundamental insight from...
macroparasite distributions also applies to Bd. For example, as predicted by macroparasite models (Barbour & Pugliese 2000), increasing Bd-induced host mortality with increasing temperature decreased the aggregation of Bd across hosts and reduced positive skew as individuals with high Bd loads were removed from the population through mortality. In fact, a sensitivity analysis of the variance to mean ratio of the Bd-load distributions showed that this measure of aggregation became progressively more sensitive to the survival function as temperature increased and more frogs experienced Bd load-dependent mortality (Fig. S6). In addition, the variance to mean ratio was more sensitive to the variance in the growth function \(v_{0,1} = 0.1 \) than the variance in the initial infection burden function \(v_{0,3} = 0.3 \), suggesting that explaining the individual-level heterogeneity in Bd growth rate may be more important for understanding the shape of the Bd-load distribution than explaining the individual-level heterogeneity in the load of Bd at initial infection. The IPM approach highlights the importance of this unexplained variance in the Bd growth function, and future studies could identify whether this heterogeneity is due to biological factors such as differences in immune responses among hosts or methodological factors such as quantitative PCR error when measuring Bd load.

In addition to allowing for a more rigorous analysis of parasite aggregation, an IPM approach can be used to examine a variety of different classic patterns in host–parasite systems. For example, host age can easily be included as an additional host attribute (Childs et al. 2003; Childs et al. 2004), such that IPMs could then be used to examine observed patterns between parasite intensity and host age (i.e. age-intensity profiles, Duerr, Dietz & Eichner 2003). Similarly, host-heterogeneity in susceptibility could be included as an additional host attribute such that IPMs could be used to explore nonlinear dose–response relationships (Dwyer, Elkinton & Buonaccorsi 1997; Gomes et al. 2014). We also discuss in Appendix S1 how \( R_0 \) can be calculated from the host–parasite IPM. While these are just a few examples, the theoretical application of IPMs for exploring observed host–parasite patterns is extensive.

While this study focused on using IPMs to describe epizootic dynamics of amphibian chytrid fungus, there are a variety of other wildlife diseases in which host–parasite IPMs could be applicable to explore the population and evolutionary outcomes of infection. For example, Tasmanian devils Sarcophilus harrisii are threatened with extinction by an infectious cancer, Tasmanian devil facial tumour disease (McCallum et al. 2009). A critical question for management is to predict the impact of the disease as it enters currently uninfected populations and to investigate evidence of selection for increased resistance to infection or reduced tumour growth rates. Intensive mark–recapture data are available, enabling the estimation of survival rates of infected and uninfected animals, together with transition rates from uninfected to infected states (Hamde et al. 2012). In addition, measurements of tumour size are taken from all infected animals at every capture opportunity and repeated tumour measurements are available for a substantial number of individuals, which could be used to estimate the tumour growth function. One could examine whether the death rate of infected devils is related to the size of the tumour and then use the IPMs to examine how differences in tumour growth among populations or over time might alter the dynamics of devil populations. It is highly likely that the death rate of infected devils is related to the size of the tumour. This problem may therefore be well-suited for an IPM approach, permitting more accurate modelling of the impact of the tumour on devil population dynamics.

Similarly, an IPM approach could also be taken to explore various aspects of the ecology and evolution of bats affected by white-nose syndrome, an emerging fungal disease of North American bats (Bleihert et al. 2008). White-nose syndrome is characterized by intense transmission, such that nearly 100% of bats of multiple species often become infected during the first winter after the fungus reaches a site (Langwig et al. 2015b). Mortality, which occurs 70–100 days after initial infection in laboratory studies (Warnecke et al. 2012), usually occurs in mid to late winter when fungal loads are highest (Langwig et al. 2015a). IPMs could be fit to pathogen loads and population dynamics of bats to explore how temperature and humidity influence pathogen growth and disease impacts (Langwig et al. 2012). Through modification of the growth function and survival function, IPMs could be used to determine whether persistence of some stabilizing populations could be explained by resistance or tolerance, or other factors affecting host–parasite interactions. For example, resistance could be described by a reduction or saturation in the growth function \( G(x, x') \), whereas tolerance could be described by a higher probability of survival \( s(x) \) for the same fungal load, \( x \).

In conclusion, IPMs can be used to answer important questions regarding host–pathogen interactions in wildlife and plant disease. Moreover, IPMs can provide new insight into many classic micro- and macroparasite patterns such as the distribution of parasites across hosts, age-intensity profiles and the dynamics of infection prevalence. By bridging the gap between micro- and macroparasites, IPMs provide an exciting new frontier in modelling wildlife disease.

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Data accessibility

The data used in this analysis can be found at http://dx.doi.org/10.5061/dryad.06s9. All the R scripts can be found in the Github repository https://github.com/mqwilber/ipm_for_parasites.

References


