Biodiversity loss decreases parasite diversity: theory and patterns

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Past models have suggested host–parasite coextinction could lead to linear, or concave down relationships between free-living species richness and parasite richness. I explored several models for the relationship between parasite richness and biodiversity loss. Life cycle complexity, low generality of parasites and sensitivity of hosts reduced the robustness of parasite species to the loss of free-living species diversity. Food-web complexity and the ordering of extinctions altered these relationships in unpredictable ways. Each disassembly of a food web resulted in a unique relationship between parasite richness and the richness of free-living species, because the extinction trajectory of parasites was sensitive to the order of extinctions of free-living species. However, the average of many disassemblies tended to approximate an analytical model. Parasites of specialist hosts and hosts higher on food chains were more likely to go extinct in food-web models. Furthermore, correlated extinctions between hosts and parasites (e.g. if parasites share a host with a specialist predator) led to steeper declines in parasite richness with biodiversity loss. In empirical food webs with random removals of free-living species, the relationship between free-living species richness and parasite richness was, on average, quasi-linear, suggesting biodiversity loss reduces parasite diversity more than previously thought.

Keywords: coextinction; food web; parasite; biodiversity loss

1. INTRODUCTION

How should parasites respond to biodiversity loss? The field of conservation biology often views infectious organisms as a sign of imbalance and emphasizes how stressors such as climate change [1] and loss of biodiversity [2] might promote infectious disease. Indeed, some studies find that decreased diversity of non-competent hosts can increase transmission of a pathogen to species of concern, the most cited examples being West Nile virus [3], and Lyme disease [4]. Such a reduction in disease risk to human populations is heralded as an ecosystem service that can be used to market the value of biodiversity [5].

A broader perspective recognizes that parasites can decline with biodiversity loss, and parasites could make up the unseen majority of species extinctions [6–10]. Many parasites depend on complex and functioning ecosystems [11–17]. For instance, each stage in a parasite life cycle requires at least one host species. Hence, medical geographers consider how distributions of ‘vector’ hosts set the distributions of human infectious diseases. Mosquito distributions limit the reach of malaria, tsetse flies make sleeping sickness possible, black flies transmit river blindness, schistosomiasis requires certain snails, Chagas disease is absent without kissing bugs and leishmaniasis depends on sand flies. For these reasons, vector control (a type of biodiversity reduction) is a key strategy for controlling infectious disease. The link between host and parasite distributions applies to non-medically important parasites as well. For example, the trematode Pleurogonius malaclemys infects snails only in the presence of the endangered diamondback terrapin, the sole final host for the trematode [18]. When a diamondback terrapin population is extirpated, it takes its host-specific parasites with it. In this study, I examined factors that influence how parasite richness declines with biodiversity loss.

Some studies have used parasite species lists from different hosts or locations to make predictions about how free-living species diversity relates to parasite diversity. For instance, the numbers of human parasite species and free-living species at a location decrease in richness with latitude, suggesting that either the same geographical factors affect both groups or that reductions in free-living species diversity reduce parasite diversity [19]. Similarly, countries with a higher diversity of birds and mammals have a higher diversity of human parasites [20], though this could be a spurious correlation driven by increases in sampling effort with country size. Not surprisingly, locations in North America with many carnivore species have longer combined carnivore parasite species lists, leading to a strong positive correlation between carnivore diversity and the estimated diversity
of carnivore parasites [21]. Such ‘list’ studies are instructive, but their patterns have many alternative explanations, including sampling artefacts.

A few field observations have linked the diversity of parasite communities in a single host to the diversity of other hosts in the system. Most notably, the richness of trematode communities in snails increases with the diversity of birds that are final hosts for the worms [22]. Different final hosts have different diets, and this exposes them to different parasites, leading to distinct parasite ‘signatures’ in the snail population [23,24]. The diversity of invertebrates (many of which are second-intermediate hosts for the trematodes) also correlates with the diversity of trematodes in snails; however, spatial patterns can break down for mobile hosts (such as fishes) that tend to homogenize associations in snail parasites [12]. These examples suggest that efforts to protect free-living biodiversity can also increase parasite diversity. For coral reefs [25], the rocky intertidal [26], lake shores [27] and estuaries [14], sites protected from human disturbance can also increase parasite diversity. For coral reefs suggests that efforts to protect free-living biodiversity may increase parasite diversity.

If one were to plot parasite richness against free-living species richness, what would the shape of the relationship be? Imagine that an intact system with 100 per cent of its parasite species and 100 per cent of its free-living species occupies the upper right-hand corner (1,1) of a standardized richness–richness plot such as in figure 1. Clearly, all parasites must go extinct when there are no hosts, sending the system to (0,0). Although the endpoints of the relationship between parasite and free-living species richness are obvious, the path between these endpoints is not easy to predict. A concave down relationship would suggest that parasites are robust to reductions in free-living species diversity (here, ‘robustness’ is an inverse measure of secondary extinction risk of species or groups of species). By contrast, a concave up relationship would show that parasites are sensitive to biodiversity loss. A sigmoid relationship would indicate a threshold in free-living species diversity loss that, once crossed, leads to a rapid collapse of the parasite community. Might the shape of the relationship vary among systems? If so, what factors affect the relationship?

I start by investigating a series of analytical models (table 1) that relate parasite diversity to free-living species diversity. The parasites in these models vary in the number of life stages and host specificity per life stage, whereas the hosts vary in extinction order (random, non-random or fixed). The models can be made for single parasite species, or parametrized for an average parasite species, or averaged across parasite species. This reveals classic modelling trade-offs. Mean-field approximations (modelling the average parasite species) take less data and computation, but require more assumptions, than do models that track every stage of every parasite species. A key assumption of current analytical models is that hosts do not suffer secondary extinctions when they lose their resources. Because hosts are embedded in food webs where secondary extinctions might be important, I adapt food-web disassembly to model how parasite diversity relates to free-living species diversity. This is done for hypothetical and empirical food webs. Finally, the patterns produced by the different models are summarized and compared.

2. METHODS

(a) Background

To better understand the effect of free-living species richness on parasite richness, I applied probability theory and simulation modelling to hypothetical and real communities. As a conceptual framework, I assumed a contained system, such as an island or a lake, where species could be extirpated but could not

![Graph showing relationships between parasite and free-living species richness](image-url)
recolonize. The units tracked were parasite species and free-living species, including non-hosts. To be able to compare different systems in a common currency, I expressed the richness of free-living species as a proportion (0−1) relative to the maximum (initial) richness of free-living species in the system. Parasite species richness was expressed on the same relative scale (e.g. proportion of the maximum number of parasite species). Individuals within species were not tracked, so there was no measure of abundance. However, I did specify life stages within species. For simplification, and owing to the focus of this review, I did not consider life stages of free-living species in hypothetical food webs (though some free-living species of empirical food webs did indeed have discrete life stages [28]). I also assumed that parasites did not affect free-living species diversity.

I assumed that an outside force (biodiversity loss) directly removed free-living species in a (usually) random sequence, otherwise called ‘primary extinction’ (see later text). Parasite species were not directly removed (e.g. I assume no targeted parasite-eradication effort such as the one to eliminate smallpox). Parasites that could no longer complete their life cycles experienced a secondary extinction. This form of secondary extinction is a conservative approach for determining the ‘robustness’ of communities to perturbations such as biodiversity loss [29].

I first developed simple analytical models of the system mentioned earlier to generate predictions about how the generality of parasites, life-stage complexity and differential extinction risk of hosts would affect the relationship between free-living species diversity and parasite diversity. I then used food-web disassembly models to investigate these predictions in hypothetical and empirical food webs.

(b) Analytical models

Here, I step through the analytical models from the least data intensive to the most complex, noting the assumptions and limitations of each. In general, I first build models for individual species and then adapt individual species models to communities of parasites. The more promising of these models are then given names and compared in following sections.

(i) Simple host-specific models

If a parasite present in a system is host-specific (generality g = 1), and has a single life stage, s, then the probability it will remain in the system is equal to the probability its host will remain in the system (assuming the system starts with the parasite being present), or

\[ \text{Prob}(P_i | s = 1, g = 1) = \text{Prob}(H_i), \]  

where \( P_i \) is the presence of a parasite species \( i \) in the system and \( H_i \) is the presence of the required host for \( P_i \) (the symbol ‘|’ specifies the model assumes the statements to the right). Assuming hosts and non-hosts have the same probability of being independently lost from the system, we can generalize (2.1a) to

\[ \text{Prob}(P_i | s = 1, g = 1, \text{Prob}(F) = \text{Prob}(H_i)) \]

= \text{Prob}(F).  

At the community level, the relative species richness (0−1) of parasites \( \bar{P} \) and relative species richness (0−1) of free-living species \( \bar{F} \) are equal to the average probability of occurrence, so \( \bar{P} = \text{Prob}(P_{i,j}) \) and \( \bar{F} = \text{Prob}(F_{i,j}). \) Equation (2.1b), expressed in terms of relative species richness, is therefore

\[ \bar{P} = \bar{F} | s = 1, g = 1, \sigma_F = 0, \]  

(2.1c)

where \( \sigma_F \) is the standard deviation of the probability of each free-living species being present. When plotted as relative richness of parasites versus free-living species, there is a straight line (the 1:1 line) from \( \{1, 1\} \) to \( \{0, 0\} \), consistent with the suggestion that the number of threatened parasite species is a linear function of the number of threatened hosts [9]. This is the simplest of all models and its key assumption of strict host specificity is not realistic for most parasite communities.

(ii) Generality

Because generalists are more robust to secondary extinction than are specialists [30,31], the extent of parasite generality will affect the persistence of parasites in the face of biodiversity loss. As an example, one louse species escaped global coextinction with the passenger pigeon because it also infected other pigeons [32]. A community composed of generalist parasite species should maintain its richness even after the random removal of several free-living species. Koh et al. [8] found the association between parasite richness and host richness to be increasingly concave down as the average number of hosts per parasite increases.

Including generality leads to a sampling without replacement problem. Koh et al. [8] provide an analytical approximation for datasets with single-stage parasites. Binomial probability models are far simpler, but will overestimate the probability of a parasite for concave down trajectories and underestimate it for concave up trajectories. However, for sample sizes of more than 25 free-living species, simple binomial models make a good approximation. All the datasets analysed later had sample sizes above this threshold, so sample size corrections were not used. Another complication of generality is that the parasite can persist even if only one of its hosts remains. The probability that this occurs is the complement of not having a host, or \( 1 − (1 − \bar{F})^g. \) For a single parasite species, again assuming equal extinction probabilities among free-living species, the simplest binomial approximation is

\[ \text{Prob}(P_i | s = 1, \sigma_P = 0, N_P > 25) \approx 1 − (1 − \bar{F})^g \]  

(2.2a)

where \( g \) (generality) is the number of hosts for a parasite with a single stage (or of a single stage of a parasite). I used this model to illustrate the effect of generality for single-stage parasites with two and four hosts (figure 1). A similarly simple model for a parasite community requires the assumption that parasites do not differ in generality.

\[ \bar{P} \approx 1 − (1 − \bar{F})^g | s = 1, \sigma_P = 0, \sigma_F = 0, N_P > 25. \]  

(2.2b)

However, if there is variation in generality among parasite species, estimating the relative richness of
the parasite community requires averaging among parasite species.

\[
\mathcal{P} \approx 1 - (1 - \mathcal{F})^{\frac{g}{1 + F \sigma_f}} | g = 1, \sigma_f = 0, N_F > 25. \tag{2.2c}
\]

Koh et al. [8], however, provide an analytical approximation that uses the distribution of \( g \) among species.

(iii) Life stages

Many parasites require more than one type of host to complete their life cycle (e.g. malaria requires a vertebrate and a mosquito). For this reason, parasites with complex life cycles should be more sensitive to reductions in free-living biodiversity [28,33,34], depending on the generality of each life stage. However, the effect of life stages on the relationship between parasite diversity and free-living species diversity has not yet been investigated.

Now, for a parasite species, we can write

\[
\text{Prob}(\mathcal{P}|\sigma_F = 0, \sigma_g = 0, N_F > 25) \approx (1 - (1 - \mathcal{F})^{\frac{g}{1 + m \sigma_g}}), \tag{2.3a}
\]

where \( g \) now refers to the generality of a parasite stage, not a parasite species.

If there is variation in generality among stages within a parasite \((\sigma_g > 0)\), using the arithmetic mean of the generality per stage for \( g \) can greatly overestimate the robustness of a parasite because, as shown later, stages with low generality have a disproportionate effect of robustness. The geometric mean is a better approximation, but still underestimates the robustness of a parasite if there is variation in generality among stages and this underestimate increases with \( F \). To provide a correction to the underestimate, I considered

\[
\bar{g} = \left[ \frac{\text{GM}_g}{1 + m \sigma_g} \right],
\]

where \(\text{GM}_g\) is the geometric mean of the generality among stages within a species. Here, \( m \) is a slope that indicates how the average standard deviation of generality within a species \((\sigma_g)\) influences robustness as a function of \( F \). For values of \( F \) from 0 to 1, I then solved for the \( m \) that best fit the hypothesis for lists of integer exponents that varied in their standard deviations. This resulted in a list of best-fit slopes for each value of \( F \). The best fit to these data was \( m = 0.2F \), so that

\[
\text{Prob}(\mathcal{P}|\sigma_F = 0, \bar{g} = \text{GM}_g/(1 + 0.2F \sigma_g), N_F > 25) \approx (1 - (1 - \mathcal{F})^{\frac{g}{1 + 0.2F \sigma_g}}). \tag{2.3b}
\]

Assuming no variation in the number of stages per parasite species, for a parasite community, the ‘average parasite model’ would be

\[
\bar{P} \approx (1 - (1 - \mathcal{F})^{\frac{g}{1 + 0.2F \sigma_g}})|\sigma_F = 0, \bar{g} = \text{GM}_g/(1 + 0.2F \sigma_g), \sigma_g = 0, \sigma_f = 0, N_F > 25. \tag{2.3c}
\]

However, variation in the number of stages per species and the average number of hosts per stage per species could lead to considerable error in the average parasite model (2.3c). A more data-intensive version (but with fewer assumptions) is the average of equation (2.3b) across parasite species, leading to the ‘average stage model’

\[
\bar{P} \approx (1 - (1 - \mathcal{F})^{\frac{g}{1 + 0.2F \sigma_g}})|\sigma_F = 0, \bar{g} = \text{GM}_g/(1 + 0.2F \sigma_g), N_F > 25. \tag{2.3d}
\]

For a single parasite species, an even more precise model, with even fewer assumptions would explicitly account for variation in generality among parasite stages, resulting in

\[
\text{Prob}(\mathcal{P}|\sigma_F = 0, N_F > 25) \approx \prod_{i=1}^{3} [1 - (1 - \mathcal{F})^{\frac{g}{1 + 0.2F \sigma_g}}]. \tag{2.3e}
\]

To illustrate the combined effect of life stages and generality, I calculated the trajectories of multi-stage parasites and single-stage parasites for equation (2.3e) (figure 1). The parasites differed in the distribution of generality among stages. For example, the first parasite used seven host species in one stage and 20 host species in a second stage. The coding for the first parasite was \( \{7,20\} \) and the other multistage parasites were \( \{4,4,4\}, \{1,10\}, \{1,1\}, \{1,2\} \) and \( \{1,2,2\} \).

Averaging across parasite species gives the most precise (and data intensive) measure of robustness for a parasite community, leading to the ‘variable parasite model’

\[
\bar{P} \approx \prod_{i=1}^{3} [1 - (1 - \mathcal{F})^{\frac{g}{1 + 0.2F \sigma_g}}]|\sigma_F = 0, N_F > 25. \tag{2.3f}
\]

Note that this model has the fewest assumptions of the models so far but requires information about the number of hosts used by each stage of each parasite.

(iv) Variable extinction risk among free-living species

Earlier studies have assumed that the probability of host extinction is uniform and independent of the number of parasite species per host species. This might not be the case. The rare and endangered species most likely to suffer primary extinctions might have few parasites because some parasite species will have been lost when the host became rare [35]. Under this scenario, if populations had already been depleted, parasite richness might thereafter appear to decline relatively slowly with biodiversity loss. Alternatively, large species might be more likely to suffer primary extinctions [36] and tend to host more parasite species [37]. This would decrease the robustness of parasites to biodiversity loss. Free-living species also vary in their risk of secondary extinction. In particular, specialists and top predators are more likely to go extinct owing to lack of resources during biodiversity loss. Because basal taxa have no risk of secondary extinction, plant parasites should be more robust to biodiversity loss than parasites of top predators. This is important because parasite diversity can increase with host trophic level [38], potentially leading to a negative association between parasitism and host robustness to secondary extinction. Pushing the pattern in the opposite direction is that parasites are more diverse in hosts with broad diets [37,39], which are also less sensitive to secondary extinction [36]. Finally, if parasites make hosts more extinction-prone [40], hosts with lots of parasites may be more likely
to go extinct. Although there is no consensus on how sensitive hosts should be to extinction relative to non-hosts, it seems useful to be able to accommodate variation in extinction rates among free-living species. As a starting point, equation (2.1c) can be modified to

$$\text{Prob}(P_i | s = 1, g = 1) = F', \quad (2.4a)$$

where $r$ is the risk of a host being absent from the system relative to all other free-living species. If $r$ is high, the host is sensitive to extinction, and the parasite will be more likely to suffer a secondary extinction. Obviously, the extent parasites decline with free-living biodiversity loss depends on how extinction-prone their hosts are relative to non-hosts. As an example, I used model (2.4a) to calculate trajectories for situations where the single host of a parasite had equal, twice or half the chance of primary extinction of other free-living species (figure 2). A more general model for a single parasite with stages and multiple hosts is

$$\text{Prob}(P | N_F > 25) \approx \prod_{i=1}^{25} \left[1 - \prod_{j=1}^{5} (1 - F^i)\right]. \quad (2.4b)$$

In addition, for a parasite community, the ‘variable host and parasite model’ is

$$P \approx \prod_{i=1}^{25} \left[1 - \prod_{j=1}^{5} (1 - F^i)\right] \mid N_F > 25. \quad (2.4c)$$

This model has the fewest assumptions of all (and requires the most detailed input). For illustration purposes, the variable host and parasite model was run for an empirical food web with hypothesized variation in extinction risk (see later text).

(v) Non-probabilistic models

An alternative non-probabilistic model can be used for cases where an extinction order, $O_p$, of the hosts and non-hosts can be hypothesized. In other words, suppose free-living species will go extinct in order of their rarity or threat level (e.g. host $q$ is the 14th free-living species to go extinct, and host $k$ is the 44th free-living species to go extinct) [41]. To simplify the calculations, simultaneous extinctions of free-living species do not occur (though simultaneous extinctions of parasite species are possible). With this information, we should be able to discern the extinction order of the parasites in the system. Extinction order relates to the previous probability models, because the expected order of extinction of a species is inversely related to its probability of extinction. Estimating extinction order for a parasite species requires the same information as model (2.4c), but uses actual extinction orders of hosts instead of relative extinction risk. For each parasite, one first estimates the extinction order of the hosts for each stage. Because the stage can persist until its last host goes extinct, the extinction order of a parasite, $O_p$, is the minimum of the maximum host extinction orders for each parasite stage

$$O_p \approx X_{(1)}[X_{(2)}(O_a, O_b, \ldots, O_i), X_{(3)}(O_a, O_b, \ldots, O_i)] \ldots X_{(g)}(O_a, O_b, \ldots, O_i)], \quad (2.5a)$$

where $O_p$ is the extinction order of parasite, $X_{(i)}$ is the maximum of the extinction orders of hosts ($a$ through $i$) for a parasite stage $i$, and $X_{(1)}$ is the minimum of the maximum host extinctions across all stages of the parasite. Equation (2.5a) is repeated for each of the $N_p$ parasite species in the system. From the set of $N_p$ parasite extinction orders, it is possible to calculate how relative parasite richness declines with biodiversity loss. After the $n$th free-living extinction, the proportion of parasites that have an extinction order $> n$ will still be present in the system. The set of points representing the relative parasite richness corresponding to a value of relative free-living species richness is the ‘variable host and parasite extinction order model’

$$\{1, 1, \ldots, \{(\sum O_p > 1)/N_p, (N_F - 1)/N_F\}, \ldots, \{(\sum O_p > 2)/N_p, (N_F - 2)/N_F\}, \ldots, \{0, 0\}\}. \quad (2.5b)$$

This model is not particularly useful when there is a lot of uncertainty in the extinction order of hosts (making a probabilistic model such as (2.4c) more appropriate).

(c) Food-web models

The analytical models mentioned above do not account for connections among hosts or between hosts and non-hosts. These connections create network structures that can lead to variation in the risk of secondary extinction among free-living species. Furthermore, food webs can lead to correlated patterns of extinction in hosts that, as I will illustrate, can alter the trajectory of extinction for parasites. For this reason, I used a modified form of robustness analysis to incorporate food-web structure into the relationship between parasite and free-living species richness. Robustness analysis considers the presence–absence of species in a food web, focusing on secondary extinctions that result from resource loss [30, 31]. It takes the concept of secondary extinctions of parasites that have been the key to the analytical models mentioned above and extends them to free-living species as well. Instead of using binomial
analytical models, however, robustness of food webs is performed by computation. This topological approach to simulating disassembly of ecological communities requires few assumptions about how species interact, and thus allows analyses of complex species dependencies not amenable to dynamical modelling [29]. However, food-web disassembly requires a large amount of information about a system and it takes many disassemblies to arrive at an expected result if extinction order is not set.

I created a program in Mathematica to disassemble large matrices multiple times. For each iteration, the program randomly removed one free-living species from a topological food web as a way to simulate biodiversity loss. Any species (parasitic or free-living) left without resources went secondarily extinct. I then calculated the number of species remaining in the food web separately for parasites and free-living species. This process continued until no species remained. The trajectory of parasite species loss during a single iteration depended on the order in which free-living species were removed during a disassembly, and there are up to $N!$ sequences of species removals (where $N$ is the number of free-living species). Unless otherwise indicated, extinction order was randomized, and the disassembly was repeated 500 times (known from past work to give good average estimates of disassemblies [42]). Averaging the 500 simulations gave a prediction for the relationship between average relative parasite richness and relative host richness.

(i) Effects of food-web topology
To illustrate how food-web structure can affect the relationship between parasite and free-living species diversity, I constructed simple food webs with five free-living species and two parasite species. In all cases, the parasites had a single host, but the structure of their food webs varied remarkably. In the first food web, parasites were independent, and the free-living species were all basal, so no free-living species suffered a secondary extinction. This was the same assumption of the analytical models mentioned above. I then changed these assumptions by making a second simple food web in which the three non-hosts formed a food chain, increasing the risk of secondary extinction of non-hosts compared with hosts. In a third system, the two hosts were consumers; each specialized on a separate basal species. A fourth food web consisted of two hosts sharing the same basal resource so that their fates were no longer independent. I then used the disassembly approach described earlier to calculate the average extinction sequence for the parasite community. To illustrate the effect of correlated extinction between parasites and consumers, I constructed two simple food webs with 10 free-living species and one parasite species. In all cases, the parasites had a single host, but the structure of their food webs varied only in that the parasite shared or did not share its host with other consumers.

(ii) Empirical food webs
I applied the disassembly approach to nine empirical food webs with parasites. These were a New Jersey stream, Muskinham Brook [43], the pelagic web of an Arctic lake, Takvatn [44], and seven estuaries: the Ythan River Estuary [45], Otago Harbour [46], Flensburg Estuary [47], Sylt Estuary [48], Carpinteria Salt Marsh, Estero de Punta Banda and Bahia San Quintin [49]. I examined and explained the variation in the shapes of the relationship between free-living and parasite diversity across the nine systems and compared these with predictions from analytical models. I considered that detritus was the last resource to go ‘extinct’. Otherwise, initially, all species were assumed to have the same risk of secondary extinction.

To explain variation in the robustness of single parasite species to secondary extinction, I calculated the relative order of extinction during disassembly for each species. Parasites that went extinct later, on average, than other species in the food web were assumed to be more robust to secondary extinction [33]. For this measure, relative order $= 1$ roughly corresponds to a linear, less than 1 to a concave up, and more than 1 to a concave down trajectory for parasite richness with biodiversity loss. I also tracked the following statistics for each parasite: number of stages, mean and standard deviation of the number of hosts per stage and the average relative order of host extinction, nested within parasite stage. I predicted that the robustness of a parasite to secondary extinction would increase with increasing generality per life stage and with the robustness of its hosts. I also predicted a parasite’s robustness would decrease with the average number (and variation) of life stages. To determine which factors explained parasite robustness, I used a generalized linear model (GLM), considering food web as a random effect.

I asked similar questions about the parasite community as I asked for parasite species. For each food web, I measured the robustness of the parasite community to biodiversity loss as the average relative order of extinction for the parasite species in comparison with free-living species. As before, I predicted that parasite robustness would increase with average parasite generality, decrease with the average number of parasite stages, and increase with the robustness of hosts to secondary extinctions. I again used a GLM, considering each food web as a replicate.

Finally, I considered non-random extinction probabilities for the Carpinteria Salt Marsh food web, because this was a system where I had enough information to propose hypotheses for the relative extinction risk among free-living species. The first hypothesis was that extinction probability decreased with biomass density under the assumption that rare species (controlling for the effects of body size on density) were more likely to be extirpated from a system. The second hypothesis was that the risk of extinction decreased with the frequency that a species was present in three similar estuaries [49]. For instance, species that occurred only in one of the three sites were assumed to be three times more likely to go extinct than species found in all three sites. Some nodes were aggregated taxa such as phytoplankton. I assumed these had a low (one-fifth) rate of extinction controlling for the effects of body size on density. The free-living species C. californica, was known to have been extirpated from several sites in other parts of its range [50], and it
was assumed to have a 10-fold rate of extinction risk. The variable host and parasite model (2.4c) and the variable host and parasite extinction order model (2.5b) was run for each list and the results compared with uniform extinction risk.

(iii) Evaluating model performance

Ideally, an analytical model should be simple, but fit the data well. To compare the performance of the various analytical models, I used the nine food-web models mentioned earlier as a benchmark. For a particular set of inputs, I calculated $P_{\text{fit}}$ and average $P$ for a food-web model and the analytical models. For a range of biodiversity loss from 0 to 100 per cent, I calculated the average absolute deviation of each analytical model from the food web model. I also noted if there was a bias in a particular direction. Deviations were calculated at the level of individual parasites and for the parasite community. In addition, to illustrate model fit, I plotted the predictions of the three analytical models and the food-web model for a lake and stream food web.

3. RESULTS

(a) Analytical models

(i) Generality of parasites

As expected from basic probability theory and past work, generality increased the robustness of parasites to secondary extinction in the analytical model. Although the trajectory of a host-specific parasite was linear (+), generalist parasites (two and four hosts) were increasingly robust to secondary extinction, creating a concave down trajectory with biodiversity loss (figure 1).

(ii) Parasite life stages

As shown by other studies [28,33], the presence of multiple life stages in a parasite species greatly reduced the robustness of parasites to secondary extinction (figure 1). For instance, the concave up trajectory for parasite $\{1,1\}$ is far below the diagonal for a single stage host-specific parasite (+), and opposite to the shape of the generalist parasite $\{2\}$ with the same number of hosts. Adding generality for parasites with complex life cycles resulted in a diversity of curves (figure 1) that show how the opposing effects of generality and life stages affect the trajectory of a parasite species. If stages varied in generality, the stage with the minimum number of hosts tended to dominate the trajectory. For instance, parasite $\{1,10\}$ had nearly the same trajectory as a single-stage host-specific parasite (+).

(iii) Differential extinction risks for hosts

Not surprisingly, a change in the probability of extinction for hosts relative to non-hosts altered the relationship between free-living and parasite richness (figure 2). If host and non-hosts had the same probability of being removed, the relationship was linear for a specialist parasite. If the host was more likely than non-hosts to be removed, the relationship was concave up. If the host was less likely to be removed than non-hosts, the relationship was concave down. Fits of models (2.4) and (2.5) are discussed in §3b.

Figure 3. A food-web disassembly takes multiple paths. Plotted are four randomly selected trajectories for the disassembly of a system with five free-living and two parasite species, indicating how extinction order can lead to different relationships between free-living and parasite richness. Up to 5! trajectories are possible, though many trajectories would be redundant. Figure 4 shows the average trajectory (food web A) is a straight line. Feeding links connect parasite (filled circles) and free-living (open circles) species.

(b) Food-web models

(i) Effects of food-web topology

Figure 3 shows four of the 120 possible disassembly trajectories for a single food web of five free-living basal species and two parasite species that each uses a different host. In nature, it will be difficult to predict which trajectory will be taken unless extinction order is known. Averaging all trajectories would result in the straight line shown in figure 4, food web A, and this gives a general prediction for the trajectory of the parasite.

The topology of the food web altered the relationship between parasite and free-living species richness in ways not predictable from analytical models. In the illustrative example (figure 4), the two parasite species were always specialists and had a single stage, yet they had different average disassembly trajectories owing to simple differences in food-web topology. For food web A, all free-living species were independent and, because they were basal, all had no risk of secondary extinction. This fulfilled the assumptions of analytical models and led to
the linear relationship between free-living and parasitic richness seen for a specialist parasite in figures 1 and 2. For food web B, non-hosts were more likely to suffer secondary extinctions than hosts. This increased the robustness of the parasites relative to free-living species. This was similar to the results of figure 2, except here the difference in extinction risk among free-living species was due solely to topological properties of the web. For food web C, hosts had a higher risk of extinction than non-hosts, leading to a concave up association between free-living and parasite richness (again consistent with figure 2). For food web D, the hosts shared a resource, so their fates were partly dependent. Their joint fate led to a conditional probability in the risk of secondary extinction for the parasite community, slightly changing the trajectory in comparison with food web C.

In the second illustrative example (figure 5), the parasite species were always specialists on a basal species, had a single stage, and were equally robust to secondary extinction, yet they had different disassembly trajectories owing to a change in a single link. For the network in the upper left of the figure, the parasite infected a host that supported many other specialist consumers. If the host went extinct, there were several simultaneous secondary extinctions, leading to a correlation between parasite extinction and biodiversity loss. This led to a strongly concave down relationship between parasite richness and free-living species richness. For the network in the lower right of figure 5, the parasite infected a host that supported no other consumers. When its host went extinct, no additional biodiversity was lost from the system and this led to an initially concave up relationship between host richness and parasite richness. The solid line is the analytical prediction from the binomial distribution (corrected for small sample size), which roughly splits the difference between the two networks. This illustrates how the analytical models cannot easily account for correlations in secondary extinctions between parasites and free-living species.

Table 2. Relative frequency distributions of the shape of the decline in the probability of a parasite species with declines in biodiversity for nine empirical food webs as calculated with a food-web disassembly model. Superscript letters indicate: E, estuary; S, stream; L, lake. BSQ, Bahía de San Quintín; CSM, Carpinteria Salt Marsh; EPB, Estero de Punta Banda and Bahia San Quintin.

| Empirical food webs | % distribution of parasite species trajectories |
|---------------------|-----------------------------------------------|
| web/shape           | concave down | concave up | linear | sigmoid |
| BSQE                | 35           | 40         | 17     | 8       |
| CSM                 | 32           | 26         | 39     | 4       |
| EPB                 | 21           | 28         | 43     | 8       |
| FlensburgE          | 24           | 42         | 27     | 7       |
| MuskinghamSE        | 69           | 8          | 23     | 0       |
| OtagoE              | 61           | 22         | 11     | 6       |
| SyltE               | 24           | 12         | 65     | 0       |
| TakvatnE            | 18           | 73         | 9      | 0       |
| YthanSE             | 62           | 19         | 19     | 0       |
| average             | 38           | 30         | 28     | 3       |

(ii) Empirical food webs
The empirical webs had 11–80 parasite species, which correlated positively with the 39–133 free-living species, although Otago Harbour had relatively few parasites (19) given its 126 free-living species. The average number of stages per parasite ranged from 1.5 to 2.3 and the average number of hosts per stage ranged from 1.9 to 7.7 (there was no association between these two variables).

Individual parasite species’ trajectories with biodiversity loss were split among concave down (38%), concave up (30%) and quasi-linear (28%) shapes (table 2). Only 3 per cent of parasite species had a sigmoid trajectory. However, the relative frequency of trajectory shapes varied among the food webs. For instance, most species in Muskingham Brook food web had concave down trajectories, whereas most of the parasites in the Lake Takvatn food web had concave down.
extinction, as measured by the average parasite relative loss order, could be explained by just two variables. Average parasite robustness to biodiversity loss increased with the average number of hosts per parasite stage and decreased with the average number of stages per parasite (table 4). On average, an increase in one parasite stage per parasite cancelled the robustness obtained by adding more than a dozen hosts.

(iii) Differential extinction

As suggested by the analytical results (figure 2), adding variation to the primary extinction probabilities of the free-living species changed the relationship between parasite richness and free-living species richness (figure 7). When density was used as a measure of the resilience of free-living species to primary extinction, the trajectory was sigmoid, with parasite richness being robust to initial levels of biodiversity loss (relative loss order of 1.24), followed by a sharp decline in parasite richness as free-living species diversity dipped below 50 per cent. When species frequency was used as measure of resilience to secondary extinction (in my opinion, this is the more reasonable hypothesis), the parasite richness trajectory was concave up, indicating that parasites were more sensitive to biodiversity loss (relative loss order of 0.87) than would be expected if all hosts had the same risks of extinction (relative loss order of 1.08). This is consistent with the observation that several parasite species in this system depend on the extinction-prone snail C. californica [51].

(iv) Model performance

The analytical models did not perfectly correspond to the food webs models (table 5; figures 7 and 8). Average absolute deviations ranged from 4 to 17 per cent. The ‘average parasite model’ (2.3c) fit relatively poorly. The simpler ‘average stage model’ (2.3d) fit

Table 3. Results from a general linear model of parasite robustness to biodiversity loss for nine empirical food webs. Parasite robustness is measured as the average relative loss order of a parasite (during a series of food-web disassemblies) compared with the average free-living species in that food web. The statistic for hosts per stage is the geometric mean \((R^2 = 0.79)\). Site was a random factor (variance component = 0.0013). \(N = 347\) parasite species.

| factor                      | estimate | s.e. | d.f. | d.f.den | F-ratio | p          |
|-----------------------------|----------|------|------|---------|---------|------------|
| intercept                   | 1.178    | 0.030| 1    | 68.3    | 3.6466  | <0.0001    |
| stages                      | -0.253   | 0.013| 1    | 341.6   | 31.54   | <0.0001    |
| s.d. hosts per stage        | -0.086   | 0.015| 1    | 336.2   | 14.36   | 0.0002     |
| host robustness             | 0.068    | 0.018| 1    | 341.8   | 811.55  | <0.0001    |
| log(hosts per stage)        | 0.342    | 0.012| 1    | 341.6   | 811.55  | <0.0001    |

Table 4. Results from a general linear model of average parasite robustness to biodiversity loss for nine empirical food webs. Average parasite robustness is measured as the average of relative loss orders (during a series of food-web disassemblies) of all parasites in a food webs compared with free-living species in that food web. The statistic for hosts per stage and stages per parasite is the geometric mean \((R^2 = 0.77; N = 9\) food webs).

| source, factor              | estimate | s.e. | d.f. | SS    | F-ratio | p        |
|-----------------------------|----------|------|------|-------|---------|----------|
| model                       |          |      | 2    | 0.1   | 9.97    | 0.0124   |
| error                       |          |      | 6    | 0.0   |         |          |
| intercept                   | 1.835    | 0.234| 1    | 1.00  | 6.96    | 0.0002   |
| hosts per stage             | 0.065    | 0.025| 1    | 0.0   | 17.34   | 0.0059   |
| stages per parasite         | -0.524   | 0.126| 1    | 1.01  |         |          |
Table 5. Average absolute deviations between food-web models and analytical models for trajectories of parasite species and the parasite community in response to biodiversity loss. Smaller values indicate better fits to the food-web model. The ‘variable parasite model’ (models (2.3e,f)) is the most complex, followed by the ‘average stage model’ (model (2.3d)) and the ‘average parasite model’ (models (2.3b,c)). Superscripts indicate: E, estuary; S, stream; L, lake; BSQ, Bahía de San Quintín; CSM, Carpinteria Salt Marsh; EPB, Estero de Punta Banda. Note, for 500 iterations, the inherent absolute deviation among replicates of the food-web disassembly model was 0.02, suggesting a deviation near 0.02 was a good fit for an analytical model.

| web/model | parasite species | parasite community |
|-----------|------------------|-------------------|
|           | variable parasite | average stage     | variable parasite | average stage | average parasite |
| BSQ_E     | 0.07             | 0.08              | 0.02             | 0.02          | 0.22             |
| CSM_E     | 0.05             | 0.06              | 0.01             | 0.02          | 0.18             |
| EPB_E     | 0.06             | 0.07              | 0.01             | 0.02          | 0.14             |
| Flensburg_E | 0.09            | 0.09              | 0.01             | 0.01          | 0.04             |
| Muskingham_S | 0.04          | 0.04              | 0.01             | 0.02          | 0.05             |
| Otago_F   | 0.20             | 0.13              | 0.07             | 0.07          | 0.13             |
| Sylt_E    | 0.18             | 0.17              | 0.07             | 0.07          | 0.24             |
| Takvatn_L | 0.09             | 0.09              | 0.05             | 0.05          | 0.06             |
| Ythan_E   | 0.09             | 0.09              | 0.01             | 0.01          | 0.05             |
| average   | 0.10             | 0.09              | 0.03             | 0.03          | 0.12             |

Figure 7. The relative extinction risk of hosts affects how parasite richness declines with free-living species richness in an empirical food web. Plotted are associations between free-living species richness and parasite richness obtained from the disassembly of the Carpinteria Salt Marsh food web under two different hypotheses about variation in the relative risk of extinction of free-living species: declining risk with biomass density (solid sigmoid line), and declining risk with the frequency a species was present in three estuaries (solid concave up line). Associated with each disassembly are estimates from the ‘variable host and parasite model’ (dashed lines, model (2.4c)) and the ‘variable host and parasite extinction order model’ (dotted lines, model (2.5b)).

Figure 8. Relative fits of analytical models to trajectories created by food-web disassemblies. The (upper) concave down dashed line shows the disassembly for Lake Takvatn. The ‘variable parasite model’ (model (2.3f), square symbols) and ‘average stage model’ (model (2.3d), dot symbols) had similar trajectories that underestimated the disassembly model whereas the ‘average parasite model’ (model (2.3c), × symbols) initially overestimated and then underestimated. The lower concave up solid line shows the disassembly for Muskimming Brook. The ‘variable parasite model’ and ‘average stage model’ had similar trajectories that fit the disassembly model well. In comparison, the ‘average parasite model’ overestimated the disassembly trajectory.

as well as the more complex ‘variable parasite model’ (2.3f), indicating that the average stage model was a short cut worth taking. The performance of analytical models improved considerably at the community level, indicating that errors cancelled instead of magnified.

The ‘variable host and parasite model’ (2.4c) was a good match for the food-web disassembly data in the case that extinction order was frequency-dependent. However, this model did not fit the density-dependent extinction order results well, primarily because of the high variance in density among the species (leading to several parasite species that could seemingly persist at low free-living biodiversity). Conversely, the ‘variable host and parasite extinction order model’ (2.5b) fit best when extinction order was density-dependent. The fit to the frequency-dependent extinction order was low because the extinction orders were not well defined, meaning that many alternative trajectories were equally likely.

4. DISCUSSION

The average relationship between relative parasite richness and host richness could be linear, sigmoid, concave up or concave down and all empirical webs had a mix of parasites that varied in their robustness to biodiversity loss. This does not imply the lack of a general pattern. The overall average trajectory of
these species was usually quasi-linear, suggesting a stronger effect of biodiversity loss on parasite richness than the concave down relationship previously estimated from models that considered only the number of hosts per parasite [8]. Moreover, the factors responsible for the shape were measurable.

Perhaps the most obvious factor affecting the risk to parasites of biodiversity loss was differential susceptibility of hosts to primary extinction. Assuming all free-living species have the same probability of primary extinction makes it easier to build analytical models. Changing this assumption (as I did for the analysis of Carpinteria Salt Marsh) greatly altered how parasite richness declined with biodiversity loss. Certainly, free-living species are not all equally susceptible to primary extinctions [41]. Humans target some species for fishing, hunting and other types of commercial exploitation. Demographic stochasticity and environmental variability should be more likely to affect large species with small population sizes [52]. In particular, species with small ranges or limited dispersal ability are less likely to be able to recolonize an area if they are locally extirpated [53]. Because parasites are not distributed randomly throughout food webs with respect to topology [39], they might not be randomly distributed with respect to the risk of hosts to primary extinction. Host species with small ranges should have fewer parasite species than species with large ranges, making parasite richness less sensitive to biodiversity loss [54]. By contrast, large free-living species tend to host more parasite species than do small free-living species [37,55,56]. Therefore, host susceptibility to primary extinction in a real system should be given a careful consideration when making specific predictions.

Generality decreased the sensitivity of parasites to losses in free-living species diversity because generalist parasites have multiple ways to complete their life cycles when their hosts go extinct. Unfortunately, knowing the average number of hosts per parasite in a community was not enough to estimate the relationship between parasite richness and host richness. Having multiple life stages increased the sensitivity of parasites to losses in free-living species diversity, because a parasite with a complex-life cycle needs more than one host to be present to persist, increasing the likelihood that some critical host will go extinct under scenarios of biodiversity loss. Life stages are an additional layer of host specificity that makes parasites extra-sensitive to secondary extinction [28]. As indicated in the statistical models, the average number of hosts per life stage is meaningful only in the context of the variation in the number of hosts per life stage. A host-specific life stage can make a parasite susceptible to secondary extinction even if other life stages can use many hosts. The s.d. of hosts per stage was, therefore, a useful predictive variable for parasite robustness in analytical models.

Food-web topology was the hardest factor to account for with analytical models because food webs lead to interactions among species, resulting in conditional probabilities that are difficult to estimate analytically. The removal of one host can lead to the loss of many parasites, and the removal of an important basal species could lead to the loss of many hosts simultaneously. Although it is difficult to predict the net effect of food-web topology on parasites, the results of this study suggest that analytical models without information on the topology of free-living species can often adequately predict the general shape of the empirical relationship between free-living and parasitic richness. Although host robustness to secondary extinction was useful in explaining the robustness of a particular parasite to secondary extinction, the average robustness of the host community (weighted by parasite richness) did not help predict the average robustness of the parasite community. It is difficult to explain why the effect of host robustness did not scale up to the community level. Consistent with past findings [33], hosts were not, on average, more or less likely to suffer secondary extinctions than were non-hosts based on their position in a food-web topology. Averages might obscure important, but variable, contributions at the species level.

Although I focused on biodiversity loss, one could also consider how introduced species or biodiversity restoration could affect the relationship between parasite and free-living species diversity. If, as is common, introduced species leave their parasites behind, parasite diversity will not respond strongly to increases in free-living species diversity due to invasion. If the parasite-poor invaders outcompete native species with many parasites, the result could be a decrease in the parasite richness of the system even as new free-living species are added [34,51]. Otherwise, new free-living species can bring parasites into a system of resident hosts that lack prior exposure. In these cases, the new parasite diversity could reduce free-living species richness through disease-driven extinctions, though such events are rare [57]. Therefore, species additions could lead to a negative association between free-living and parasite richness. There can be a positive relationship between invasive parasites and hosts if invaders do not escape natural enemies. The human-mediated addition of two regionally common fish to a species-poor lake appears to have facilitated the addition of five parasite species [58]. Similarly, if restoration of native biodiversity occurs, we should see an increase in both parasite and free-living biodiversity. For instance, the diversity of trematodes in estuarine snails increased steadily over 6 years after a habitat restoration project, presumably because the restoration succeeded in creating habitat for the various invertebrates, fishes and birds that the trematodes required to complete their life cycles [14]. However, not all restoration efforts will succeed in attracting parasites. In a heathland restoration, the lack of parasites to fully recolonize the region suggested that the effort had failed to recreate the complex trophic interactions found in natural habitats [59].

The alternative perspective that biodiversity loss will increase infectious disease comes from cases of the dilution effect and host compensation. Dilution effects occur when some host species interfere with parasite transmission. The removal of interfering hosts (but not those that are required for transmission) can lead to an increase in the prevalence of certain infectious diseases such as Lyme [4]. The net outcome in prevalence for a particular parasite depends on whether the required hosts or the interfering free-living species are more likely to suffer extinctions. Host compensation can result if some hosts become
abundant due to release from predators or competition, leading to more efficient transmission of infectious diseases [60]. The importance of host compensation depends on the strength of trophic cascades and the relative abundances of impacted versus released hosts. Although dilution and compensation are possible results of biodiversity loss, there is no logical reason to expect that they will be the rule or overshadow secondary extinctions.

The strong association between parasite richness and host richness suggests that parasites can be positive indicators of free-living species richness [13,15,61]. Specifically, a high diversity of parasites indicates a complex and functioning set of interacting free-living species. For example, the abundance of parasitoid insects increases remarkably with native plant diversity in the Azores [62]. Some parasites will be more suitable indicators of free-living species diversity than others. The best source of indicative parasites would be an easy to sample, abundant host species that has a high diversity of parasites with complex life cycles. In aquatic systems, ideal parasite communities for monitoring are found in fishes [17,63–65] and snails [22,24,66,67].

Dynamical models [42,68] are a tool that could help answer some remaining questions. Such models could allow parasites to impact their hosts, and non-competent hosts to impact parasites, leading to more complex relationships between parasite and free-living species diversity. Four possibilities might emerge from dynamical models. If parasites have density-dependent effects on a guild of competitors, they might prevent competitive exclusion and promote diversity [69]. Alternatively, if parasites are generalists and vary in their impact on hosts, they could drive intolerant host species extinct [70]. If some hosts interfere with parasite transmission, their addition to a system could reduce parasite diversity via the dilution effect [4]. Finally, because parasites might go extinct before their hosts [71], parasites might respond even more strongly to biodiversity loss than seen in topological models. Unfortunately, there is not sufficient empirical data to build dynamical models with parasites, so this approach is limited to hypothetical systems for now.

In conclusion, parasite richness declines as free-living biodiversity is lost, resulting in a positive association between parasite richness and free-living species richness. As is the case for free-living species, parasite generality buffers parasites to host losses, whereas complex life cycles add an extra set of resource (host) requirements that reduce parasite robustness to secondary extinctions. Food-web topology leads to conditional probabilities that can complicate how parasite richness relates to free-living species richness. On average, empirical webs showed a quasi-linear decline in relative parasite richness with relative free-living species richness. As a result, parasites are sensitive to free-living species diversity and some can be useful as bioindicators of ecosystem degradation and recovery. Although I focus on parasites, these results should be applicable to most affiliate species (mutuals and commensals).

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