Host phenology can drive the evolution of intermediate virulence strategies in some <u>obligate-killer</u> parasites

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1 Abstract

Mechanistic The traditional mechanistic trade-offs resulting in a negative correlation between transmission and virulence are the foundation of nearly all current theory on the evolution of parasite virulence. Although several Several ecological factors have been shown to modulate the optimal virulence strategies predicted from mechanistic trade-off models, none have but these ecological factors have not yet been shown to be sufficient to explain the intermediate virulence strategies observed in any natural system. The timing of seasonal activity, or phenology, is a common factor that influences the types and impact of many ecological interactions but is rarely considered in difficult to incorporate into virulence evolution studies. We develop a mathematical model of a disease system with seasonal host activity to study the evolutionary consequences of host phenology on parasite virulence the virulence of obligate-killer parasite. Results from this model demonstrated that seasonal host activity is sufficient to drive the evolution of intermediate parasite virulence , in the absence of traditional mechanistic trade-offs, in in some types of natural disease systems, even when a traditional mechanistic trade-off between transmission and virulence is not assumed in the modeling framework. The optimal virulence strategy in these systems can be determined by both the duration of the host activity period as well as the variation in the host emergence timing. Parasites with low virulence strategies are favored in environments with long host activity periods and in environments in which hosts emerge synchronously. The results demonstrate that host phenology can be sufficient to select for intermediate optimal virulence strategies, providing an alternative mechanism to account for virulence evolution in some natural systems.

1 2 Introduction

The evolutionary causes and consequences of parasite virulence remain enigmatic despite decades of research. 2 It was once thought that parasites continue to evolve ever lower levels of virulence to preserve their primary 3 resource for future parasite generations.¹ However, the idea that parasites would limit damaging their 4 host for the benefit of future generations violates multiple core principles of our modern understanding of 5 evolutionary biology.^{2,3} Natural selection, as framed in the modern evolutionary synthesis, favors traits that 6 improve short-term evolutionary fitness even if those traits negatively impact the environment for future 7 generations.^{2,3} Thus, the level of virulence that maximizes parasite fitness is favored by natural selection despite its impact on the host population.⁴ Identifying environmental conditions and mechanistic constraints q that drive the evolution of the vast diversity of parasite virulence strategies observed in nature has been an 10 important research focus for decades. 11

Establishing that mechanistic trade-offs constrain parasite virulence strategies was a key breakthrough that 12 propels virulence evolution research to this day. In the now classic paper, Anderson and May demonstrated 13 that within-host parasite densities increase the probability of transmission to a new host - a component of 14 parasite fitness - but also shorten infection duration resulting in fewer opportunities for transmission to naive 15 hosts.⁴ That is, parasites can only produce more infectious progeny if they cause host damage by utilizing 16 more host resources. This mechanistic trade-off that Mechanistic trade-offs results in a negative correlation 17 between virulence and transmission remains and remain the foundational theoretical framework used to 18 account for the evolution of all-parasite virulence strategies observed in nature.^{5,6} 19 Many ecological factors and environmental conditions have been shown to alter the optimal virulence 20 strategies driven by mechanistic trade-offs within models. For example, it is well-established that varying 21 environmental conditions, such as the extrinsic host death rate, often shift the optimal virulence strategy 22 governed by a mechanistic trade-off.^{4,7-9} However, no environmental condition, in the absence of mechanistic 23 trade-offsan explicitly modeled negative correlation between virulence and transmission, has been shown 24 to select for intermediate virulence. 25 The timing of seasonal activity, or phenology, is an environmental condition affecting all aspects of life 26 cycles, including reproduction, migration, and diapause, in most species.^{10–16} The phenology of host species 27 also impacts the timing and prevalence of transmission opportunities for parasites which could alter optimal 28

virulence strategies.¹⁷⁻²⁴ For example, host phenological patterns that extend the time between infection and
 transmission are expected to select for lower virulence, as observed in some malaria parasites (*Plasmodium*)

³¹ *vivax*). In this system, high virulence strains persist in regions where mosquitoes are present year-round

³² while low virulence strains are more common in regions where mosquitoes are nearly absent during the

³³ dry season.²⁵ While host phenology likely impacts virulence evolution in parasites,^{26–29} it remains unclear

³⁴ whether this environmental condition can have a sufficiently large impact to select for an intermediate

³⁵ virulence phenotype in the absence of a mechanistic trade-off.

Here we investigate the impact of host phenology on the virulence evolution of an obligate-killer parasite. We demonstrate that intermediate virulence is adaptive when host activity patterns are highly seasonalin the absence of any explicit mechanistic trade-off, establishing that environmental context alone is sufficient to drive the evolution of intermediate virulence in disease systems that conform to the assumptions of the model. Further, multiple features of host seasonal activity, including season length and the synchronicity at which hosts first become active during the season, impact the optimal virulence level of parasites. These results provide an alternative framework that can account for virulence evolution in some patural systems

results provide an alternative framework that can account for virulence evolution in some natural systems.

3 Model description

⁴⁴ The model describes the transmission dynamics of a free-living, obligate-killer parasite that infects a seasonally

available host (Figure 1). The host cohort, \hat{s} , enters the system at the beginning of the season over a period

 $_{46}$ given by the function $g(t, t_l)$. Hosts, *s*, have non-overlapping generations and are alive for one season. The

⁴⁷ parasite, *v*, infects hosts while they are briefly susceptible early in their development (*e.g.* baculoviruses of ⁴⁸ forest *Lepidoptera*^{30–34} and univoltine insects parasitized by ichneumonids^{35–37}). The parasite must kill the

⁴⁸ forest *Lepidoptera*³⁰⁻³⁴ and univoltine insects parasitized by ichneumonids³⁰⁻³⁷). The parasite must kill the
 ⁴⁹ host to release new infectious progeny. The parasite completes one round of infection per season because

⁴⁹ host to release new infectious progeny. The parasite completes one round of infection per season because
 ⁵⁰ the incubation period of the parasite is longer than the duration of time the host spends in the susceptible

⁵¹ developmental stage. This transmission scenario occurs in nature if all susceptible host stages emerge over

⁵² a short period of time each season so that there are no susceptible host stages available when the parasite

⁵³ eventually kills its host. Parasites may also effectively complete only one round of infection per season if

the second generation of parasites do not have enough time in the season to complete their life cycle in the

⁵⁵ short-lived host.

We ignore the progression of the susceptible stage, s, to later life stages as it does not impact transmission dynamics. To keep track of these dynamics, we refer to the generation of parasites that infect hosts in the beginning of the season as v_1 and the generation of parasites released from infected hosts upon parasite-induced death as v_2 . τ is the delay between host infection by v_1 and host death when v_2 are released. τ is equivalent to virulence where low virulence parasites have long τ and high virulence parasites have short τ . The initial conditions in the beginning of the season are thus $s(0) = 0, v_1(0^+) = v_2(0^-), v_2(\tau) = 0$. The transmission dynamics in season n are given by the following system of delay differential equations (all parameters are described in Table 1):

$$\frac{ds}{dt} = \hat{s}g(t, t_l) - \frac{ds}{\delta m} - \frac{\mu s}{\delta m}(t) - \alpha s(t)v_1(t),$$
(1a)

$$\frac{dv_1}{dt} = -\delta v_1(t),\tag{1b}$$

$$\frac{dv_2}{dt} = \alpha\beta e^{-d\tau - \mu\tau} s(t - \tau)v_1(t - \tau) - \delta v_2(t).$$
(1c)

⁵⁶ where $\frac{d}{d\mu}\mu$ is the host death rate, δ is the decay rate of parasites in the environment, α is the transmission rate,

 $_{57}$ β is the number of parasites produced upon host deathand au is the delay between host infection and host

⁵⁸ death. au is equivalent to virulence where low virulence parasites have long au and high virulence parasites

⁵⁹ have short τ . We make the common assumption for free-living parasites that the removal of parasites through

transmission (α) is negligible,^{34,38,39} *i.e.* (1b) ignores the term $-\alpha s(t)v_1(t)$.

⁶¹ The function $g(t, t_l)$ is a probability density function that captures the per-capita host emergence rate by

⁶² specifying the timing and length of host emergence. We use a uniform distribution $(U(\bullet))$ for analytical

⁶³ tractability, but other distributions can be used.

$$g(t, t_l) = \begin{cases} \frac{1}{t_l} & 0 \le t \le t_l \\ 0 & t_l < t \le T \end{cases}$$

 t_l denotes the length of the host emergence period and T denotes the season length. The season begins

($t_0 = 0$) with the emergence of the susceptible host cohort, \hat{s} . The host cohort emerges from $0 \le t \le t_l$. v_2

₆₆ parasites remaining in the system at t = T give rise to next season's initial parasite population ($\hat{v} = v_1(0)$).

⁶⁷ Parasites that have not killed their host by the end of the season do not release progeny. Background mortality

Parameter Description		Value
s	susceptible hosts	state variable
v_1	parasites that infect hosts in current season	state variable
v_2	parasites released in current season	state variable
t_l	length of host emergence period	time (varies)
T	season length	time (varies)
\hat{s}	emerging host cohort size	10^8 hosts
α	transmission rate	$10^{-8}/(\text{parasite} \times \text{time})$
β	number of parasites produced upon host death	parasites (varies)
δ	parasite decay rate in the environment	2 parasites/parasite/time
d-µ	host death rate	0.5 hosts/host/time
τ	time between host infection and host death (1/virulence)	time (evolves)

Table 1: Model parameters and their respective values.

arises from predation or some other natural cause. We assume that infected hosts that die from background

⁶⁹ mortality do not release parasites because the parasites are either consumed or the latency period corresponds

⁷⁰ to the time necessary to develop viable progeny.^{40,41} We ignore the impact of infection for host demography

and assume \hat{s} is constant each year (*e.g.* a system where host regulation by parasites is negligible). We solve

⁷² equations 1a-c analytically Appendix A.



Figure 1: **Infection diagram** The host cohort, \hat{s} , emerges from time t = 0 to $t = t_l$, all v_1 parasites emerge at t = 0. Hosts do not reproduce during the season. Infections generally occur early in the season when host density is high. Parasite-induced host death occurs after time τ , at which point new parasites, v_2 are released. v_2 decays in the environment from exposure. Parasites only have time to complete one round of infection per season. v_2 parasites in the environment at t = T will carryover and emerge at the beginning of the next season.

73 3.0.1 Parasite fitness

A parasite introduced into a naive host population persists or goes extinct depending on the length of the host emergence period and season length. The stability of the parasite-free equilibrium is determined by the production of v_2 resulting from infection of s given by

$$v_2(T) = e^{-\delta(T-t_l-\tau)} (v_2(t_l) + \alpha\beta e \underbrace{-d\tau - \mu\tau}{\delta} \hat{v}s(t_l) \int_0^{T-t_l-\tau} e \underbrace{-\frac{\alpha\hat{v}e^{-\delta(s+t_l)}(-1+e^{\delta s})}{\delta} - \delta t_l + ds}_{\delta} \underbrace{-\frac{\alpha\hat{v}e^{-\delta(u+t_l)}(-1+e^{\delta u})}{\delta} - \delta t_l - \mu u}_{\delta} \underbrace{-\frac{\alpha\hat{v}e^{-\delta(u+t_l)}(-1+e^{\delta u})}{\delta} - \mu u}_{\delta} - \mu u}_{\delta} \underbrace{-\frac{\alpha\hat{v}e$$

when $\tau < T - t_l$ and by

$$v_2(T) = \frac{\alpha\beta e^{-\mu\tau}\hat{v}\hat{s}}{t_l}e^{-\delta(T-\tau)} \int_0^{T-\tau} e^{(-\mu u + \frac{\alpha\hat{v}e^{-\delta u}}{\delta})} \int_0^u e^{(\mu x - \frac{\alpha\hat{v}e^{-\delta x}}{\delta})} dxdu$$

74 when $\tau > T - t_l$.

The parasite-free equilibrium is unstable and a single parasite introduced into the system at the beginning of the season will persist if the density of v_2 produced by time T is greater than or equal to $\hat{v} = v_1(0) = 1$ (*i.e.* $v_2(T) \ge 1$, modulus is greater than unity). This expression is a measure of a parasite's fitness when rare given different host phenological patterns. See Appendix A for details of the analytical solution.

79 3.0.2 Parasite evolution

⁸⁰ To study how parasite traits adapt given different seasonal host activity patterns, we use evolutionary invasion

analysis.^{42,43} We first extend system (1) to follow the invasion dynamics a rare mutant parasite

$$\frac{ds}{dt} = \hat{s}g(t, t_l) - \underbrace{d\mu}_{\sim} s(t) - \alpha s(t)v_1(t) - \alpha_m s(t)v_{1m}(t),$$
(2a)

$$\frac{dv_1}{dt} = -\delta v_1(t),\tag{2b}$$

$$\frac{dv_{1m}}{dt} = -\delta_m v_{1m}(t),\tag{2c}$$

$$\frac{dv_2}{dt} = \alpha \beta e^{-d\tau - \mu\tau} s(t - \tau) v_1(t - \tau) - \delta v_2(t),$$
(2d)

$$\frac{dv_{2m}}{dt} = \alpha_m \beta_m e \underbrace{-d\tau_m - \mu \tau_m}_{\sim \sim \sim \sim} s(t - \tau_m) v_{1m}(t - \tau_m) - \delta_m v_{2m}(t).$$
(2e)

where *m* subscripts refer to the invading mutant parasite and its corresponding traits. See Appendix B for
details of the time-dependent solutions for equations (2a-2e).

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The invasion fitness of a rare mutant parasite depends on the density of v_{2m} produced by the end of the season $(v_{2m}(T))$ in the environment set by the resident parasite at equilibrium density \hat{v}^* . The mutant parasite invades in a given host phenological scenario if the density of v_{2m} produced by time T is greater than or equal to the initial $v_{1m}(0) = 1$ introduced at the start of the season $(v_{2m}(T) \ge 1)$. When $\tau < T - t_{l}$, mutant invasion fitness can be found using

$$v_{2m}(T) = e^{-\delta_m (T - t_l - \tau_m)} (v_{2m}(t_l) + \alpha_m \beta_m e_{\underline{-d\tau_m - \mu\tau_m}}^{-d\tau_m - \mu\tau_m} v_{1m}(0) s(t_l) \\ \int_0^{T - t_l - \tau_m} e_{\underline{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (s+t_l)}(-1 + e^{\delta_m s})}{\delta_m} - \frac{\alpha \hat{v}^* e^{-\delta(s+t_l)}(-1 + e^{\delta_s})}{\delta} - \delta_m t_l - ds} ds_{\underline{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{\delta_m u})}{\delta_m} - \frac{\alpha \hat{v}^* e^{-\delta(u+t_l)}(-1 + e^{\delta_s s})}{\delta_m}}}{(3a)}$$

⁹⁰ When $\tau > T - t_l$, mutant invasion fitness can be found using

$$v_{2m}(T) = \frac{\alpha_m \beta_m e^{-\mu \tau_m} v_{1m}(0)\hat{s}}{t_l} e^{-\delta_m (T-\tau_m)} \int_0^{T-\tau_m} e^{(-\mu u + \frac{\alpha \delta^* e^{-\delta u}}{\delta} + \frac{\alpha_m v_{1m}(0)e^{-\delta_m u}}{\delta_m})} \int_0^u e^{(\mu x - \frac{\alpha \delta^* e^{-\delta s}}{\delta} - \frac{\alpha_m v_{1m}(0)e^{-\delta_m x}}{\delta_m})} dx du$$
(3b)

⁹¹ To study the evolution of virulence traits, we first assume all other resident and mutant traits are identical

⁹² (*e.g.* $\alpha = \alpha_m$). Note that when there is no trade-off between β and τ , the parasite growth rate in the host is

essentially the trait under selection. That is, β is constant regardless of τ thus, thus the trait that is effectively

evolving is the time between infection and when the parasite kills the host and releases new parasites is the

⁹⁵ rate that β -new parasites are assembled --in between infection and host death (*e.g.* long τ corresponds to

⁹⁶ slow assembly of new parasites.) To find optimal virulence for a given host phenological scenario, we find

⁹⁷ the uninvadable trait value that maximizes (3). That is, the virulence trait, τ^* , that satisfies

$$\frac{\partial v_{2m}(T)}{\partial \tau_m}\Big|_{\tau_m = \tau_r} = 0 \tag{4a}$$

$$\frac{\partial^2 v_{2m}(T)}{\partial \tau_m^2}\big|_{\tau_m = \tau_r} < 0 \tag{4b}$$

⁹⁸ Note that the measure in equation (3) incorporates the effect of the resident on the population state (number

of susceptibles over one season), which means that it is not a measure of R_0 (which by definition assumes

a non-disease environment). Thus, we can use $v_{2m}(T)$ as defined in (3) as a maximand in evolutionary

¹⁰¹ dynamics.⁴⁴

¹⁰² To study the impact of mechanistic trade-offs between transmission and virulence on virulence evolution, we ¹⁰³ assume that the number of parasites produced at host death is a function of the time between infection and

host death ($\beta(\tau)$). For example, mutant invasion fitness for $\tau < T - t_l$ can be found using

$$v_{2m}(T) = e^{-\delta_m (T - t_l - \tau_m)} (v_{2m}(t_l) + \alpha_m \beta(\tau_m) e^{-\frac{d\tau_m - \mu\tau_m}{2}} v_{1m}(0) s_{\underline{1}}(t_l) \\ \int_0^{T - t_l - \tau_m} e^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (s+t_l)}(-1+e^{-\delta_m s})}{\delta_m} - \frac{\alpha_0^* e^{-\delta(s+t_l)}(-1+e^{-\delta s})}{\delta} - \delta_m t_l - ds} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1+e^{\delta_m u})}{\delta_m} - \frac{\alpha_0^* e^{-\delta(u+t_l)}(-1+e^{-\delta_m s})}{\delta_m}}{(5)}$$

¹⁰⁵ We then find τ^* that satisfies (4a) and (4b) using equation (5).

106 4 Results

Host phenology , in the absence of a mechanistic trade-off, is sufficient to drive the evolution of intermediate 107 virulence in systems that conform to the assumptions of the model. Host phenology is composed of the 108 duration of the activity period and the distribution of initial emergence times, both of which impact the 109 optimal parasite virulence level. Temporally constrained host activity periods within each season can select 110 against both extremely high and extremely low virulence levels resulting in an intermediate optimal level 111 of virulence. Low virulence is selected against as parasites that do not kill the infected host prior to the 112 end of the host activity period fail to produce progeny and thus have no evolutionary fitness. By contrast, 113 highly-virulent parasites kill their hosts quickly and the released progeny decay in the environment for the 114 remainder of the activity period. Thus, progeny released early in the host activity period are more likely to 115 die in the environment prior to contacting a naive host in the following season. An intermediate virulence 116 level that allows parasites to kill their host prior to the end of the activity period, but not so quickly that the 117 progeny produced are likely to decay in the environment, result in the greatest evolutionary fitness. 118

The optimal virulence level increases linearly with decreases in the duration of host activity (Figure 2). 119 Virulent parasites in environments where host activity periods are short minimize the cost of not producing 120 progeny from infected hosts and do not incur the costs of progeny decaying in the environment. By contrast, 121 environments where host activity periods are long favor parasites with a long incubation period to limit the 122 cost of progeny decay due to environmental exposure while still killing hosts prior to the end of the season. 123 The optimal level of virulence in all environmental scenarios results in parasite-induced host death just prior 124 to the end of the seasonal activity period. The linear increase in optimal virulence as season length decreases 125 suggests that parasite fitness is optimized when host death occurs at a fixed time before the end of the season. 126 Variation in the time at which each host first becomes active during the activity period also impacts the 127 virulence levels that maximize parasite fitness (Figure 3). Synchronous host emergence results in a rapid and 128 early spike in infection incidence due to the simultaneous availability of susceptible hosts and the abundance 129 of free parasites. The long duration between host infection and the end of the activity period favors low 130 virulence parasites that kill their host near the end of the season (Figure 3A, i). Variability in the time at 131 which each susceptible host initially becomes active decreases the average time between infection and the 132 end of the season, thus favoring more virulent parasites (Figure 3A, *ii*). That is, the large proportion of 133 infections that occur later in the season require higher virulence to be able to release progeny before the 134 activity period ends. This higher virulence level comes at the cost of progeny from hosts infected early in the 135 season decaying in the environment. Thus, the number of progeny that survive to the next season decreases 136 with increasing variation in host emergence times (Figure 3B). 137

High variability in host emergence timing results in an optimal virulence strategy that is much greater 138 than in environments with synchronous host emergence, but lower than in environments with a moderate 139 distribution (Figure 3). That is, increasing variation in host emergence timing favors parasites with higher 140 virulence, but only when variation in host emergence timing is highmoderate. In environments where the 141 variation in host emergence timing is high, increasing variation in host emergence timing favors parasites 142 with slightly lower virulence (Figure 3A, *iii*). Lower virulence is favored in high emergence variability 143 environments because the number of new infections occurring late in the season, where high virulence would 144 be advantageous, are relatively rare due to small parasite population sizes at the beginning of the season and 145 parasite decay during the season. Initial parasite population sizes are smaller in environments with broadly 146 distributed host emergence timing as fewer total hosts are infected because infection probability is density 147

dependent, and thus fewer progeny are produced. Most parasites that find a susceptible host do so early 148 in the season resulting in additional decreases to the already small parasite population size. The optimal 149 virulence strategy allows parasites that infect hosts around the peak of new infections - occurring mid-season 150 when susceptible host densities are greatest and parasite populations have not decayed substantially - to 151 release progeny while limiting decay of these progeny. Parasites in environments where the distribution in 152 host emergence times is very broad suffer the costs of both decay of the progeny released by early-infected 153 hosts and the cost of late infected hosts not releasing progeny, collectively causing these environments to 154 maintain low densities of moderately virulent pathogens (Figure 3B, *iii*). 155 Mechanistic virulence-transmission trade-offs can modify the optimal virulence strategy in seasonal

156 environments but are not necessary for natural selection to favor intermediate virulence phenotypes. The 157 optimal virulence strategy is slightly lower in models that include a trade-off where duration of infection is 158 positively correlated with progeny production than in models with the same phenological parameters that do 159 not include the trade-off (Figure 4). Including this trade-off increases the fitness benefit of longer-duration 160 infections to a greater extent than the costs associated with infected host mortality not caused by the parasite. 161 By contrast, the optimal virulence strategy is greater in models that include a trade-off where duration of 162 infection is negatively correlated with progeny production than in similar models without the trade-off 163 (Figure 4). Including this trade-off increases the fitness benefit of shorter-duration infections despite the 164 added costs of greater parasite decay due to environmental exposure. Including mechanistic trade-offs 165 modifies the selection pressures on virulence strategies but are not essential for an intermediate virulence 166

¹⁶⁷ strategy to be optimal in seasonal environments.



Figure 2: Host seasonality is sufficient to select an intermediate virulence strategy. A. The temporal duration between infection and host death (τ^*) always evolves to a value that is greater than 0 (extreme virulence) and less than the season length (extremely low virulence); the intermediate virulence strategy maximizes parasite fitness in environments where host activity is seasonal. The optimal parasite-induced host death rate results in host death and progeny release shortly before the end of the season (t = T). The release of progeny just prior to the end of the season limits the decay of progeny due to environmental exposure while avoiding progeny dying within their host at the end of the season. *i* and *ii* are representative examples of optimal virulence strategies in environments with shorter (T = 3.2) or longer (T = 4) host activity periods, respectively. τ^* is found using equation (4a) when there is no trade-off between transmission and virulence. **B.** Higher parasite virulence is favored in environments with limited host activity periods. Parasites with greater virulence produce more progeny that survive to the end of the season when seasons are short. That is, the density of the more virulent progeny (i) at T = 3.2 is greater than the density of the less virulent progeny (ii). The more virulent parasite kill their hosts quickly such that few infected hosts survive to the end of the season and the progeny released spend little time in the environment. By contrast, less virulent parasites (ii) often fail to kill their hosts and release progeny prior to the end of short activity periods (T = 3.2). Longer seasons (T = 4) favor less virulent parasites (*ii*) as they kill their hosts closer to the end of the season such that fewer of their released progeny decay in the environment (ii) than the progeny of the more virulent parasites that are released earlier in the season (i). The blue line represents the incidence rate of new infections; $t_l = 1$; all other parameters found in Table 1.

168 5 Discussion

Nearly all theory developed to explain parasite virulence evolution has utilized mechanistic trade-offs between 169 virulence and other traits important to parasite fitness.^{5,6} The results of this study show that seasonal host 170 activity, in the absence of a mechanistic trade-offan assumed negative correlation between virulence and 171 transmission, can account for the evolution of intermediate virulence in some specific situations. Both aspects 172 of phenology, the duration of the host activity period and host emergence synchronicity, impact the virulence 173 strategy that maximizes the evolutionary fitness of parasites. Although mechanistic trade-offs between 174 virulence and transmission can shift the optimal virulence level as predicted by prior theory, these trade-offs 175 are not essential for intermediate virulence to evolve in this system. The current demonstration that an 176 ecological context is sufficient to select for intermediate virulence broadens the scope of factors that can 177 explain the diversity of parasite virulence strategies. Thus, the evolution of intermediate virulence in natural 178 systems may be governed by a mechanistic trade-off or by ecological factors in some systems. 179

Seasonal host activity can select for intermediate virulence by generating conflicting costs for releasing progeny too early or too late in the season. Low virulence is maladaptive for parasites in this system as they do not kill their host before the end of the season and create no progeny. High virulence is also maladaptive as progeny released early are more likely to die due to environmental exposure. The conflicting costs of not releasing progeny before the end of the season and releasing progeny too early in the season selects for



Figure 3: The variation in host emergence timing impacts the optimal virulence strategy. A. Parasites with lower virulence are favored in environments where nearly all hosts emerge simultaneously (*i*). Progeny from the low virulence parasites are released nearly simultaneously just prior to the end of the season. High virulence parasites are favored in environments where host emergence period length is moderate (*ii*). Moderate variation in host emergence decreases the average time between infection and the end of the season and favors parasites with a high virulence strategy such that few infected hosts survive to the end of the season. Parasites in environments where host emergence variation is very high maximize the number of progeny that survive to the next season by using a moderate virulence strategy (*iii*). Parasites in these environments suffer the costs of hosts that are infected later in the season not releasing progeny as well as progeny decay in the environment when released from early-infected hosts. A moderate virulence strategy allows hosts infected around the mid-season peak in incidence to release progeny while limiting the decay of these progeny. τ^* is found using equation (4a) when there is no trade-off between transmission and virulence. **B.** Equilibrium density of parasites with the optimal virulence results in peak equilibrium in new parasites density, indicated by the vertical lines. T = 3; other parameters found in Table 1.

intermediate virulence levels. Optimal virulence results in parasite-induced host death and the release of
 progeny slightly before the end of the host activity period.

The result predicting adaptive evolution towards intermediate virulence stands in contrast to many 187 prior theoretical investigations of obligate-killer parasites. Prior models of obligate-killer parasites predict 188 ever-increasing virulence in the absence of mechanistic trade-offs.⁴⁵⁻⁴⁸ In simple obligate-killer models, 189 killing infected hosts as quickly as possible is expected to maximize fitness as the early release of progeny 190 permits infection of additional susceptible hosts resulting in a rapid exponential increase of parasites in 191 the system. To date, only mechanistic trade-offs between virulence and transmission-associated factors as 192 well as development time constraints have been demonstrated to constrain maximal virulence in obligate-193 killer parasite models.^{40,46,48–50} In contrast, our results indicate that host phenology - in the absence of an 194 explicit mechanistic trade-off and development time constraints - can create the can create conditions that 195 favor intermediate virulence in obligate-killer parasites even if a negative correlation between virulence and 196 transmission is not included in the model. In the current model, intermediate virulence is favored as seasonal 197



Figure 4: Mechanistic transmission-virulence trade-offs shift the optimal virulence strategy but are not necessary to favor intermediate virulence in environments with seasonal host activity. The optimal virulence level for parasites in which longer durations of infection result in *more* progeny is slightly lower than for parasites that are not constrained with this mechanistic trade-off in the same environment (*i*). This mechanistic trade-off elevates the fitness benefit of longer duration infections by compensating for the cost of infected hosts dying without releasing progeny. The optimal virulence level for parasites in which longer infection durations result in *fewer* progeny is greater than for parasites without this trade-off in the same seasonal environments (*ii*). This mechanistic trade-off elevates the fitness benefit of shorter duration infections despite the cost of greater progeny decay in the environment. τ^* was found using equation (4a) when there is no trade-off between transmission and virulence and then compared to τ^* constrained by a trade-off with transmission. Trade-off for $i : \beta(\tau) = 99(\tau + 0.5)$, trade-off for $ii : \beta(\tau) = 99(-\tau + 4)$. All other parameters found in Table 1.

¹⁹⁸ host absence increases the evolutionary benefit of remaining within hosts in order to reduce deaths in the

¹⁹⁹ free-living stage caused by environmental exposure.⁵¹

Variation in host emergence synchronicity impacts the optimal virulence strategy of parasites in this 200 system. High parasite virulence is favored at low host emergence synchronicity. Low emergence synchronicity 201 slows incidence by decreasing both the rate hosts emerge and parasite equilibrium density. When more 202 infections occur later in the season, parasites have less time to release new parasites before the end of the 203 season. High parasite virulence is adaptive because hosts have a low expected life span at the time of infection. 204 This result is analogous to the prediction that high host mortality drives the evolution of high virulence.^{4,7–9} 205 The timing of host activity can thus lead to the evolution of high virulence in a similar manner to how host 206 demography impacts virulence. 207

The seasonal activity patterns of species with non-overlapping generations may have large impacts on the 208 virulence strategies of the parasites they host. For example, parasites and parasitoids of univoltine insects 209 that complete one round of infection per host generation may maximize their fitness by releasing progeny just 210 prior to the end of the season.^{30–37} The theoretical expectations presented here can be tested empirically by 211 measuring the virulence strategies of parasites across the natural diversity of phenological patterns observed 212 over the geographical range of many insect species. Similarly, experiments could rigorously assess the impact 213 of both season length and host emergence variability on the fitness of parasites with different levels of 214 virulence. 215

The prediction that shorter host activity periods can drive greater virulence is comparable to how the 216 virulence of different Theileria parva strains varies between regions. High within-host densities permit a 217 virulent T. parva strain to be reliably transmitted to feeding nymphal tick vectors shortly after being infected 218 by the adult stage in regions where the activity patterns of the two tick life stages overlap.^{52–54} In contrast, the 219 virulent strain is absent in regions where nymphal and adult activity is asynchronous while a less virulent 220 strain that persists in hosts longer is maintained.^{52,53} Thus, the prediction that the length of the host activity 221 period is inversely correlated with virulence coincides with empirical observations of the distribution of T. 222 parva strains. 223

Several features of the current model can be altered to investigate more complex impacts of phenology on virulence evolution. For example, relaxing the assumption of a constant host population size may result in a feedback between parasite fitness and host demography with consequences for population dynamics⁵⁵. Additionally, parasite virulence evolution may select for alternative host phenological patterns that in turn select for parasite traits with lower impacts on host fitness. We will extend the current model to address these questions in future studies.

The model presented applies to obligate-killer parasites that complete one round of infection per season 230 (monocyclic) in hosts that have non-overlapping generations. Currently, there is no evidence that disease 231 systems that do not conform to these assumptions will not require a mechanistic trade-off to-violate these 232 assumptions can select for intermediate virulence without including a mechanistic trade-off. Nevertheless, 233 several prior models that included both host seasonality and mechanistic trade-offs found qualitatively similar 234 results as those presented here despite relaxing one or more of the strict assumptions in this model,²⁶⁻²⁸ 235 suggesting that phenology can have a large impact on virulence outcomes. For example, longer seasons 236 or longer periods between seasons have been shown to select for lower virulence in polycyclic parasites 237 in seasonal environments,^{27,28} similar to the results presented here. Similarly, explicitly modeling parasite 238 growth rates within hosts, which underlie the correlation between virulence and instantaneous transmission 230 rates, selects for intermediate virulence levels that maximize transmission rates during host activity periods.²⁶ ²⁴¹ By contrast, assuming that virulence levels are mechanistically associated with host density results in selection
 ²⁴² for higher virulence in seasonal environments.²⁹ Future studies incorporating one or more of these competing
 ²⁴³ forces with environmental decay of progeny could be sufficient to select for intermediate virulence in the
 ²⁴⁴ absence of an explicit assumed mechanistic trade-off.

Some of the strict model assumptions can likely be relaxed without altering the result that phenology can 245 be sufficient to select for intermediate virulence strategies. Relaxing the obligate-killer assumption may result 246 in the same qualitative result that intermediate virulence is adaptive in some cases. For example, longer 247 latency periods that result in progeny release near the end of the season would still be adaptive for parasites 248 that reduce host fecundity or increase host death rate, even if there is no correlation between the virulence 249 level and instantaneous or life-time transmission. Longer latency periods are equivalent to lower virulence in 250 this type of system as infected hosts have more time to reproduce and thus higher fitness. This extension 251 is not expected to qualitatively alter the results if the parasite transmission period is short relative to the 252 season length. Many parasite-host systems conform to the assumptions of this model extension such as 253 monocyclic plant pathogens (e.g. soil-borne plant pathogens, demicyclic rusts, post-harvest diseases), and 254 many diseases systems infecting univoltine insects $\frac{56-59}{56-60}$. 255

The importance of parasite virulence to both host-parasite interactions and public health policy has 256 resulted in a concentrated research effort on virulence evolution. Nearly all theoretical research to date 257 has incorporated a mechanistic trade-off between virulence and transmission rates or infection duration, a 258 hypothesis which is still essential to explain the evolution of intermediate virulence in most disease systems. 259 However, ecological factors such as seasonal host activity or spatial structuring provide alternative theoretical 260 frameworks that may account for virulence strategies in some natural systems.^{61,62} Future work that identifies 261 and empirically validates ecological factors that influence virulence evolution would be useful for predicting 262 outbreaks of highly virulent parasites. 263

²⁶⁴ 6 Code and data availability

²⁶⁵ Code is available on the Github repository: https://github.com/hanneloremac/Host-phenology-drives-the ²⁶⁶ evolution-of-intermediate-parasite-virulence

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404 Appendix A

⁴⁰⁵ In Appendix A we find analytical solutions for equations (1a-c) from the main text to study parasite fitness ⁴⁰⁶ given different host phenological patterns.

$$\frac{ds}{dt} = \hat{s}g(t, t_l) \underline{-ds} \underbrace{-\mu s}_{\sim}(t) - \alpha s(t)v_1(t), \tag{A.1a}$$

$$\frac{dv_1}{dt} = -\delta v_1(t),\tag{A.1b}$$

$$\frac{dv_2}{dt} = \alpha\beta e^{-d\tau - \mu\tau} s(t - \tau)v_1(t - \tau) - \delta v_2(t).$$
(A.1c)

with initial conditions: $s(0) = 0, v_1(0^+) = v_2(0^-), v_2(\tau) = 0.$

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⁴⁰⁹ (A.1a-c) is solved analytically by describing host emergence using a uniform distribution

410
$$g(t, t_l) = \begin{cases} \frac{1}{t_l} & 0 \le t \le t_l \\ 0 & t_l < t \end{cases}$$

⁴¹¹ To solve the dynamics during the host's activity period, we first find the analytical solution for $v_1(t)$:

$$v_1(t) = \hat{v}e^{-\delta t}$$

We then use $v_1(t)$ to find the time-dependent solutions for both solution for s(t) and. We can then plug the time-dependent solution for s(t) to find the time-dependent solution for $v_2(t)$:-. Only parasites that infect hosts from $0 < t < T - \tau$ have enough time to kill hosts and release progeny before the end of the season. For $\tau < T - t_l$, parasites that infect hosts during host emergence $(0 < t < t_l)$ have time to kill hosts and release progeny before the end of the season. For $\tau < T - t_l$, parasites that infect hosts during host emergence $(0 < t < t_l)$ have time to kill hosts and release progeny before the end of the season as well as some parasites who infect hosts after host emergence has ended $(t > t_l)$. For $\tau > T - t_l$, only some parasites that infect hosts during host emergence $(0 < t < t_l)$ have time to kill hosts and release progeny before the end of the season. Thus, two separate solutions are required depending on whether τ is greater or less than $T - t_l$. We first consider the case where $\tau < T - t_l$:

$$s(t) = \begin{cases} \frac{\hat{s}}{t_{l}} e^{(-\mu t + \frac{\alpha \hat{v}e^{-\delta t}}{\delta})} \int_{0}^{t} e^{(\mu u - \frac{\alpha \hat{v}e^{-\delta u}}{\delta})} du & 0 < t < t_{l} \\ s(t_{l}) e^{(-\mu (t - tl) - \frac{\alpha \hat{v}e^{-\delta (t + t_{l})}(-1 + e^{\delta t})}{\delta})} & t_{l} \le t < T \end{cases}$$

$$v_{2}(t) = \begin{cases} \frac{\alpha \beta e^{-\mu \tau} \hat{v} \hat{s}}{t_{l}} e^{-\delta (t - \tau)} \int_{0}^{t - \tau} e^{(-\mu u + \frac{\alpha \hat{v}e^{-\delta u}}{\delta})} \int_{0}^{u} e^{(\mu x - \frac{\alpha \hat{v}e^{-\delta x}}{\delta})} dx du & \tau < t < t_{l} \\ e^{-\delta (t - t_{l} - \tau)} (v_{2}(t_{l}) + \alpha \beta e^{-\mu \tau} \hat{v} \hat{s}(t_{l}) \int_{0}^{t - t_{l} - \tau} e^{-\frac{\alpha \hat{v}e^{-\delta (u + t_{l})}(-1 + e^{\delta u})}{\delta} - \delta t_{l} - \mu u} du) & t_{l} \le t < T \end{cases}$$

where $s(t_l)$ and $v_2(t_l)$ are the densities of s and v_2 when the emergence period of s ends.

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For $\tau > T - t_{l}$, only some of the parasites that infect hosts from $0 < t < t_{l}$ have enough time to kill hosts and release progeny before the end of the season. $v_{2}(t)$ are thus only produced from infections that occurred

from $0 < t < t_l$. The solution for $v_2(t)$ in this case is

$$v_{2}(t) = \frac{\alpha\beta e^{-\mu\tau}\hat{v}\hat{s}}{t_{l}}e^{-\delta(t-\tau)}\int_{0}^{t-\tau}e^{(-\mu u + \frac{\alpha\hat{v}e^{-\delta u}}{\delta})}\int_{0}^{u}e^{(\mu x - \frac{\alpha\hat{v}e^{-\delta x}}{\delta})}dxdu \qquad \tau \leq t \leq T$$

A parasite introduced into a naive host population persists or goes extinct depending on the host emergence period length and season length. The stability of the parasite-free equilibrium when $\tau < T - t_L$ is determined by the production of v_2 resulting from infection of s given by

$$v_2(T) = e^{-\delta(T-t_l-\tau)} (v_2(t_l) + \alpha\beta e^{-d\tau - \mu\tau} \hat{v}s(t_l) \int_0^{T-t_l-\tau} e^{-\frac{\alpha\hat{v}e^{-\delta(s+t_l)}(-1+e^{\delta s})}{\delta} - \delta t_l - ds} ds^{-\frac{\alpha\hat{v}e^{-\delta(u+t_l)}(-1+e^{\delta u})}{\delta} - \delta t_l - \delta t_l - ds} ds^{-\frac{\alpha\hat{v}e^{-\delta(u+t_l)}(-1+e^{\delta u})}{\delta} - \delta t_l - \delta t$$

When $\tau > T - t_l$, the stability of the parasite-free equilibrium is determined by

$$v_2(T) = \frac{\alpha\beta e^{-\mu\tau}\hat{v}\hat{s}}{t_l}e^{-\delta(T-\tau)}\int_0^{T-\tau} e^{(-\mu u + \frac{\alpha\hat{v}e^{-\delta u}}{\delta})}\int_0^u e^{(\mu x - \frac{\alpha\hat{v}e^{-\delta x}}{\delta})}dxdu$$

 $_{415}$ The parasite-free equilibrium is unstable and the parasite will persist in the system if the density of v_2

⁴¹⁶ produced by time *T* is greater than or equal to $\hat{v} = v_1(0) = 1$ introduced at the beginning of the activity ⁴¹⁷ period of *s* (*i.e.* $v_2(T) \ge 1$, modulus is greater than unity). This expression is a measure of a parasite's fitness ⁴¹⁸ when rare given for different host phenological patterns - given $\tau \ge T - t_l$.

419

420 Appendix B

In Appendix B we find analytical solutions for equations 2a-e from the main text to study the evolution of
 parasite virulence given different host phenological patterns.

$$\frac{ds}{dt} = \hat{s}g(t, t_l) - \underline{d}\mu s(t) - \alpha s(t)v_1(t) - \alpha_m s(t)v_{1m}(t),$$
(B.1.a)

$$\frac{dv_{1m}}{dt} = -\delta_m v_{1m}(t),\tag{B.1.b}$$

$$\frac{dv_{2m}}{dt} = \alpha_m \beta_m e^{-\mu \tau_m} s(t - \tau_m) v_{1m}(t - \tau_m) - \delta_m v_{2m}(t).$$
(B.1.c)

$$\frac{dv_1}{dt} = -\delta v_1(t),\tag{B.1.d}$$

$$\frac{dv_2}{dt} = \alpha\beta e^{-\mu\tau}s(t-\tau)v_1(t-\tau) - \delta v_2(t),$$
(B.1.e)

with initial conditions: $s(0) = 0, v_{1m}(0^+) = v_{2m}(0^-), v_{2m}(\tau) = 0, v_1(0^+) = v_2(0^-), v_2(\tau) = 0$. *m* subscripts refer to the invading mutant parasite and its corresponding traits.

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Again, separate solutions for (B.1.a-c) has the following time-dependent solution : are required depending on whether τ and τ_m are greater or less than $T - t_l$. The length of τ relative to T and t_l determines \hat{v}^* while the length of τ_m relative to T and t_l determines the within-season dynamics of the mutant parasite. The solutions to all cases can be found in the code on Github (see "Code and data availability" in the main text for the link). We first show the solution to the case when $\tau_m < T - t_l$:

$$\begin{split} v_{1m}(t) &= v_{1m}(0)e^{-\delta_m t} & 0 < t < T \\ s_{\underline{1}}(t) &= \begin{cases} \frac{\hat{s}}{t_l}e^{(-\mu t + \frac{\alpha\hat{v}^* e^{-\delta t}}{\delta} + \frac{\alpha_m v_{1m}(0)e^{-\delta_m t}}{\delta_m})} \int_0^t e^{(\mu u - (\frac{\alpha\hat{v}^* e^{-\delta u}}{\delta} + \frac{\alpha_m v_{1m}(0)e^{-\delta_m u}}{\delta_m}))} du & 0 < t < t_l \\ s(t_l)e^{(-\mu(t-t_l) - (\frac{(\alpha\hat{v}^* e^{-\delta(t-t_l)}(-1+e^{\delta t})}{\delta} + \frac{\alpha_m v_{1m}(0)e^{-\delta_m (t-t_l)}(-1+e^{\delta m t})}{\delta_m})}{\delta_m}) & t_l \le t < T \end{cases} \\ v_{2m}(t) &= \begin{cases} \frac{\alpha_m \beta_m e^{-\mu \tau_m} v_{1m}(0)\hat{s}}{t_l} e^{-\delta_m (t-\tau_m)} \int_0^{t-\tau_m} e^{(-\mu u + \frac{\alpha\hat{v}^* e^{-\delta u}}{\delta} + \frac{\alpha_m v_{1m}(0)e^{-\delta_m u}}{\delta_m})} \\ \int_0^u e^{(\mu x - (\frac{\alpha\hat{v}^* e^{-\delta x}}{\delta} + \frac{\alpha_m v_{1m}(0)e^{-\delta_m x}}{\delta_m}))} dx du & \tau_m < t < t_l \\ e^{-\delta_m (t-t_l-\tau_m)} (v_2(t_l) + \alpha_m \beta_m e^{-\mu \tau_m} v_{1m}(0)s(t_l) \\ \int_0^{t-t_l-\tau_m} e^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1+e^{\delta m u})}{\delta_m} - \frac{\alpha\hat{v}^* e^{-\delta(u+t_l)}(-1+e^{\delta u})}{\delta} - \delta_m t_l - \mu u} du) & t_l \le t \le T \end{cases} \end{cases}$$

When $\tau_m > T - t_l$, the solution for $v_{2m}(t)$ is

The invasion fitness of a rare mutant parasite is given by the density of v_{2m} produced by the end of the season.

The When $\tau_m < T - t_l$ the mutant parasite invades in a given host phenological scenario if the density of

 v_{2m} produced by time T is greater than or equal to the initial $v_{1m}(0) = 1$ introduced at the start of the season

429 $(v_{2m}(T) \ge 1)$ -, following

$$v_{2m}(T) = e^{-\delta_m (T - t_l - \tau_m)} (v_2(t_l) + \alpha_m \beta_m e^{-\frac{d\tau_m - \mu\tau_m}{2}} v_{1m}(0) s_{\underline{1}}(t_l) \\ \int_0^{T - t_l - \tau_m} e^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (s+t_l)}(-1 + e^{-\delta_m s})}{\delta_m} - \frac{\alpha \delta^* e^{-\delta(s+t_l)}(-1 + e^{-\delta_s})}{\delta_m} - \delta_m t_l - ds} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \frac{\alpha \delta^* e^{-\delta(u+t_l)}}{\delta_m} - \delta_m t_l - ds} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \delta_m t_l - ds} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \delta_m t_l - ds}} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \delta_m t_l - ds}} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \delta_m t_l - ds}} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \delta_m t_l - ds}} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \delta_m t_l - ds}} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \delta_m t_l - ds}} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \delta_m t_l - ds}} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \delta_m t_l - ds}} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \delta_m t_l - ds}} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \delta_m t_l - ds}} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \delta_m t_l - ds}} ds^{-\frac{\alpha_m v_{1m}(0)e^{-\delta_m (u+t_l)}(-1 + e^{-\delta_m u})}{\delta_m} - \delta_m t_l - \delta_m t_l$$

)(-:

When $\tau_m > T - t_l$, the mutant parasite invades in a given host phenological scenario if the density of v_{2m} produced by time T is greater than or equal to the initial $v_{1m}(0) = 1$ introduced at the start of the season $(v_{2m}(T) \ge 1)$, following

$$\underbrace{v_{2m}(T)}_{0} = \underbrace{\frac{\alpha_m \beta_m e^{-\mu \tau_m} v_{1m}(0)\hat{s}}{t_l} e^{-\delta_m (T-\tau_m)} \int_0^{T-\tau_m} e^{(-\mu u + \frac{\alpha \hat{v}^* e^{-\delta u}}{\delta} + \frac{\alpha_m v_{1m}(0)e^{-\delta_m u}}{\delta_m})}}_{\int_0^u e^{(\mu x - (\frac{\alpha \hat{v}^* e^{-\delta s}}{\delta} + \frac{\alpha_m v_{1m}(0)e^{-\delta_m x}}{\delta_m}))} dx du$$

We use $v_{2m}(T)$ to find optimal virulence for a given host phenological scenario by finding the trait value that maximizes $v_{2m}(T)$. That is, the virulence trait, τ^* , that satisfies

$$\frac{dv_{2m}(T)}{d\tau_m}\Big|_{\tau_m=\tau_r} = 0 \tag{B.2}$$

$$\frac{d^2 v_{2m}(T)}{d\tau_m^2}\Big|_{\tau_m=\tau_r} < 0 \tag{B.3}$$

⁴³² We use (B.2) to find τ^* in Figures 1A, 2A, 3A and 3B in the main text. For all phenological patterns, we ⁴³³ found that τ^* is uninvadable *i.e.* condition (B.3) is satisfied.

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- For certain phenological patterns, τ_m switches from $T t_l < \tau_m$ to $T t_l > \tau_m$ as it evolves. When t_l is
- small, optimal virulence, τ^* , is short relative to T and t_l . Thus when t_l is short, τ^* times parasite-induced
- ⁴³⁷ death to begin *after* all hosts have finished emerging (*i.e.* the solution where $T t_l > \tau_m$ is required to find
- 438 τ^*). For large $t_{l_{\ell}}$ the value of τ^* that optimizes parasite fitness initiates parasite-induced host death before
- all hosts have finished emerging (*i.e.* the solution where $T t_l < \tau_m$ is required to find τ^*). We found the
- value of t_l that requires a switch from the solution for $T t_l > \tau_m$ to the solution for $T t_l < \tau_m$ to find τ^*
- numerically using Mathematica. We switched which solution we used to find τ^* when the value of τ^* that
- satisfied (B.2) no longer met the inequality. For example, for long t_l , the solution for $T t_l > \tau_m$ returned
- 443 $T t_l < \tau^*$. When this occurred we switched to using the solution for $T t_l < \tau_m$ to find τ^* .
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⁴⁴⁵ To study the impact of mechanistic trade-offs between transmission and virulence on virulence evolution,

- we assume that the number of parasites produced at host death is a function of the time between infection
- and host death ($\beta(\tau)$). This is done in Figures 3A and 3B where a mechanistic trade-off is assumed to
- exist between τ and β in (B.2). The same approach as described above is used to determine which solution
- 449 correctly specifies τ^* .