Delayed Chemical Defense: Timely Expulsion of Herbivores Can Reduce Competition with Neighboring Plants

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Abstract: Time delays in plant responses to insect herbivory are thought to be the principal disadvantage of induced over constitutive defenses, suggesting that there should be strong selection for rapid responses. However, observed time delays between the onset of herbivory and defense induction vary considerably among plants. We postulate that strong competition with conspecifics is an important codeterminant of the cost-benefit balance for induced responses. There may be a benefit to the plant to delay mounting a full defense response until the herbivore larvae are mobile enough to leave and large enough to cause severe damage to neighboring plants. Thus, delayed responses could reduce the competitive pressure on the focal plant. To explore this idea, we developed an individual-based model using data from wild tobacco, Nicotiana attenuata, and its specialized herbivore, Manduca sexta. Chemical defense was assumed to be costly in terms of reduced plant growth. We used a genetic algorithm with the plant’s delay time as a heritable trait. A stationary distribution of delay times emerged, which under high herbivore densities peaked at higher values, which were related to the time larvae need to grow large enough to severely damage neighboring plants. Plants may thus tip the competitive balance by expelling insect herbivores to move to adjacent plants when the herbivores are most damaging. Thus, herbivores become part of a plant’s strategy for reducing competition and increasing fitness.

Keywords: induced defense, intraspecific competition, plant-herbivore interactions, trait diversity, individual-based model, simulation.

Introduction

Optimal defense theories predict that inducible defenses in plants are less costly than constitutive defenses because the defenses are produced only when they are actually needed (Karban and Baldwin 1997; Karban et al. 1999). As long as there are no herbivores, plants with inducible defenses can allocate their resources to primary, fitness-enhancing processes—such as growth and reproduction—and hence outcompete plants with constitutive defenses (Clark and Harvell 1992; Herms and Mattson 1992). The time lag between the environmental cue and optimal adjustment of a plant to its current environment is considered to be one of the greatest disadvantages of inducible defenses and other forms of phenotypic plasticity (Moran and Hamilton 1980; Padilla and Adolph 1996; DeWitt et al. 1998). This time lag is particularly detrimental for induced responses because herbivores may remove substantial amounts of biomass before the defenses are effective. Consequently, a strong selection pressure should exist for short time lags—or delay times—between herbivory and the production of effective chemical defenses. However, observed delay times for inducible defenses vary considerably within plant species (Mathur et al. 2011; Schuman et al. 2012). This suggests that there is little selection pressure for short delay times and even that under certain environmental conditions slower responses might actually increase fitness. To test this hypothesis, we explored the potential for neutral or even positive selection for longer delay times using a model parameterized with new and existing data for an annual plant attacked by a caterpillar.
To generate a better understanding of the concepts underlying the model, imagine two adjacent, conspecific plants that are in severe competition. When one of the plants is attacked by an herbivore (caterpillar), there are three possible scenarios (fig. 1): first, the attacked plant does not respond to feeding damage by increasing its defenses, allowing the caterpillar to complete its life cycle on the plant. In this scenario, the plant will suffer substantial damage or may even be entirely destroyed (fig. 1, left). For an annual species, this constitutes a severe fitness loss.

Second, the attacked plant may increase its defense levels very rapidly. A fast response may deter or kill the herbivore by producing direct defenses, for example, nicotine and protease inhibitors (Steppuhn and Baldwin 2007). Rapid induction may have several benefits: the direct defenses may deter or kill nonspecialized herbivores very quickly, thereby reducing leaf loss. This may not apply to specialist herbivores that can tolerate or detoxify their host plant’s specific defenses (Ali and Agrawal 2012). However, even specialist herbivores show significantly slower growth rates and higher mortality when feeding on well-defended host plants (Harvey et al. 2007; Steppuhn and Baldwin 2007). In addition, indirect defenses—such as volatile organic compounds that attract predators or parasitoids—may be induced (Kessler and Baldwin 2001; fig. 1, middle). A rapid induction of volatiles may be beneficial because parasitoids and insect predators usually prefer the early developmental stages. On the other hand, individual plants may not incur an immediate benefit from parasitoids if parasitized larvae feed more to sustain the endoparasitoids or even go through a supernumeral instar (Beckage et al. 1994). In addition, there are also costs for inducing defenses, as they may reduce the plant’s compet-

**Figure 1:** Conceptual background of the model. Plants that have induced chemical defense are violet; plants that have not are green. When a plant is attacked by a larva, it has three ways to react to the attack: it does not produce defense compounds (left), it reacts immediately by producing defense compounds (middle), or it reacts after a certain delay time \( \tau \) (right). Possible defense actions of a plant include producing toxic substances to affect larval growth and health (violet plants) and producing volatile organic compounds to attract predators, which is most effective when the larva is small because predators or parasitoids do not attack bigger larvae (middle, top two panels).
itive ability by decreasing vegetative growth because of an investment trade-off between chemical defense and vegetative growth (Louda et al. 1990; Augner et al. 1997; Nötzold et al. 1997; Baldwin et al. 1998; Redman et al. 2001). In environments with high plant growth rates, small differences in competitive ability may lead to large differences in plant size and eventually plant fitness (Nicotra and Rodenhouse 1995; Schwinning and Fox 1995; van Dam and Baldwin 2001). Plants treated with the induction hormone methyl-jasmonate (MeJA) or genotypes that overexpressed MeJA produced up to 50% less seed mass than their untreated neighbors (van Dam and Baldwin 1998, 2001; Cipollini 2002; Cipollini 2007). Because the wound signal was applied without damaging the plant, these costs are independent of leaf loss (Baldwin 1996; van Dam and Baldwin 1998; Cipollini 2007). Thus, if herbivore feeding immediately elicits an induced response, the induced plant suffers a fitness decrease compared with its noninduced neighboring competitor, regardless of how little damage the herbivore has done.

As illustrated above, short delay times may not be uniformly beneficial to the plant. This opens the possibility of a third scenario in which the attacked plant delays its defense response by a certain amount of time. If the costs of losing competitive capacity are larger than the costs of leaf loss due to herbivory, this can lead to a net increase of fitness (fig. 1, right). A similar idea (but applied to constitutive defense strategies) was first proposed by Tuomi et al. (1994). They hypothesized that plants with less effective defenses may gain an associational protection when growing close to well-defended plants. If the well-defended plants kill most of the herbivores, they might then easily be outcompeted by less defended plants (Tuomi et al. 1994). Although their model applies to constitutive defense polymorphisms and plants of different species, we propose that a similar strategy may pertain to herbivore-induced plants competing with conspecifics. Under these conditions, we postulate that plants should optimize the timing of their defense production to induce herbivore movement to a neighboring plant when the herbivore is the most damaging. This optimal delay time would be determined by larval developmental rates. In a plant-caterpillar system, later instars consume the majority of the biomass, with 90% consumed during their last instar. Thus, there is an optimal time for sending a larva to a neighboring plant so that the costs of induced defense will be offset by the gains in fitness resulting from reduced competition from the attacked neighbor.

To further explore our hypothesis and test its relevance on the population level, we developed an individual-based model representing fast-growing plants competing in dense cohorts with different densities of their mobile herbivores. We used an existing model of intraspecific plant competition (Lin et al. 2013, 2014) and included plant defenses, fitness costs, and herbivory as well as growth, survival, and movement of individual larvae. Our model is based on observations and data on *Nicotiana attenuata* (Solanaceae) and an important specialized invertebrate herbivore, *Manduca sexta* (Lepidoptera: Sphingidae). We chose this interaction because it is one of the best-studied models for induced defenses (from chemical mechanisms to ecological effects), with more than 20 years of data, providing realistic parameter values for the model.

*Nicotiana attenuata* is a fast-growing annual plant from the Great Basin Desert (southwestern United States) whose seeds germinate synchronously after a fire has destroyed the dominant vegetation (Young and Evans 1978; Koniak 1985; Baldwin and Morse 1994; Preston and Baldwin 1999). The fire pyrolyzes litter, removes negative regulators of germination, and releases smoke cues, which seeds use to synchronize their germination from long-lived seed banks (Baldwin and Morse 1994; Baldwin et al. 1994). The synchronization of seed germination results in dense cohorts of even-aged plants of more or less the same size, so intraspecific competition is a common factor affecting plant performance even under the high nutrient levels after a fire (Lynds and Baldwin 1998). The most limiting resource is water, which is present at the beginning of the season but runs short after some time and thus limits the length of the growth season.

In the first year after a large fire, the herbivore community is dominated by arthropod herbivores, such as *M. sexta*, while later in the succession vertebrate browsers will cause extensive damage (Baldwin 1998). Occasionally, *M. sexta* and its close relative *Manduca quinquemaculata* can reach outbreak densities and completely defoliate *N. attenuata* (A. Kessler, personal communication). *Manduca sexta* moths oviposit single eggs on the leaves of *N. attenuata*, on which the larvae feed for about 3 weeks until they have reached the prepupal stage and leave the plant to pupate in the soil (Gilmore 1938; Madden and Chamberlin 1945). During this time the larvae grow exponentially; while neonate larvae weigh a few milligrams, the prepupal stage may weigh 8–10 g (van Dam et al. 2001a).

*Nicotiana attenuata* employs a variety of inducible chemical defenses in response to damage or herbivore feeding, such as nicotine, proteinase inhibitors, phenolics, and volatile emissions (Keinanen et al. 2001; Kessler and Baldwin 2001; van Dam et al. 2001b). Despite the fact that *M. sexta* is a specialist herbivore and possesses several physiological mechanisms to deal with the nicotine in tobacco plants (Murray et al. 1994; Snyder et al. 1994), both its performance and its behavior are negatively affected by high levels of nicotine (Harvey et al. 2007) and other jasmonate-induced responses (Steppuhn and Baldwin 2007). Larvae that are reared on leaves of induced plants grow significantly slower and suffer a higher mortality than larvae on control leaves (van Dam et al. 2000). Moreover, if larvae are placed on jasmonate-treated plants growing adjacent to an untreated...
plant, the larva will move to the noninduced plant, and the sooner it moves and feeds on it, the larger it grows (van Dam et al. 2000).

Our model explores whether delaying chemical defense of herbivores can improve individual fitness of plants growing in dense cohorts. We tailored our model in terms of parameter values and functional relationships to mimic a specific system but also performed simulation experiments that explore a wider range of settings and discussed implications for general theory.

Methods

The Model

A full and detailed description of our model, dubbed TIMELY (short for time delay), following the overview, design concepts, and details (ODD) protocol (Grimm et al. 2006, 2010) is provided in the supplemental material (available online). Below we present a summary description. Further information on the model is given in the transparent and comprehensive model “evaluation” (TRACE) document (Schmolke et al. 2010; Grimm et al. 2014) in the supplemental material. TRACE documents contain evidence that a model was thoughtfully designed, correctly implemented, thoroughly tested, well understood, and appropriately used for its intended purpose.

We used an existing well-established model of intraspecific plant competition (Lin et al. 2013, 2014) that combines a two-layer influence approach (May et al. 2009) with an ontogenetic growth model for plants. Two layers were used because belowground competition for nutrients and water was found to be size symmetric (Casper and Jackson 1997), while aboveground competition for light is largely asymmetric (Schwinning and Weiner 1998). We adapted the model parameters to match values obtained on wild tobacco (Nicotiana attenuata) growth and competition data. The most important parameters and variables are given in table 1.

We used this model for simulations of two plants and one larva (two-plant scenario) to test the main idea of this publication. The two-plant scenario helped demonstrate and explore the consequences of delayed chemical defense in a highly simplified setting. However, in real plant populations, several factors will strongly affect the fitness consequences of delayed defense, in particular the density of plants and herbivores, the spatial distribution of plants, and the frequency distribution of delay times within the plant population. Therefore, we also extended this model to whole plant populations of several hundred plants and larvae (400-plant scenario).

The model comprises 250 × 250 square grid cells, corresponding to a total area of 15 × 15 m². Grid cells were used to represent zones of influence in a discretized way (Weiner et al. 2001; May et al. 2009). Time proceeded in discrete steps of 4 h; simulations were run for 27 days, thus the full life cycle of the caterpillars. Plants were characterized by their position, their above- and belowground biomass, their defense level, and the delay time $\tau$, defined as the time between the attack of a larva and start of defense production of the plant.

Table 1: Parameters and variables of the individual-based model

| Description                                      | Default | Range tested          |
|--------------------------------------------------|---------|-----------------------|
| Parameters:                                      |         |                       |
| No. plants                                       | Starting no. plants | 400 | 2, 100–1,000          |
| $R_a$                                            | Resource limitation aboveground | .15 | 0–6                   |
| $R_b$                                            | Resource limitation belowground | .45 | 0–6                   |
| $\tau$                                           | Delay time of the plant; $\tau$ = difference in days between first larva feeding and defense production and last larva feeding and defense production stop | Random | 0–10 days            |
| Conversion factor                                | Ratio of eaten biomass converted into larva mass | .19 | 0–1.0                 |
| Movement radius                                  | Maximal distance a larva moves to another plant | 5 m | 2–15 m                |
| Defense tolerance                                | Threshold at which mobile larvae switch plants | .24 | .15–30                |
| Variables:                                       |         |                       |
| Ticks                                            | A tick = 1/6 day | 0 | 0–240                 |
| Biomass plant                                    | Mass of whole plant | 60 g | 10–600 g              |
| Aboveground mass                                 | Mass of shoot | 30 g | 10–300 g              |
| Belowground mass                                 | Mass of root | 30 g | 10–300 g              |
| Biomass larva                                    | Proportion of defense compounds and plant mass | 1 mg | 1 mg–10 g            |
| Defense level                                    | Proportion of defense compounds and plant mass | .0 | .0–.3                  |

Note: Default and initial values of parameters and variables, respectively, are given along with possible ranges and, if applicable, units.
assumed that $\tau$ was also the time between the larva leaving the plant and the stop of defense compound production by the abandoned plant. The plants were assigned to random positions, and their delay time differed within the population but stayed constant for each plant over the simulation. Plants grow, compete for resources, produce defense compounds when being attacked, and die if they are eaten entirely by a caterpillar.

Intraspecific plant-plant competition for resources was represented separately for both roots and shoots by a two-layer zone of influence (ZOI) model. Plants obtain resources only within the area of their ZOI, with the aboveground area representing competition mainly for light and the belowground area for water, nitrogen, and other nutrients. The radii of the ZOIs were allometrically related to the respective above- and belowground biomasses. Competition occurred only where ZOIs overlapped. In simulations including genetic algorithms, plant reproduction was represented as well (see “Genetic Algorithm”).

Larvae were characterized by their biomass, age, and position. Initially they were distributed on randomly chosen plants. They first stayed on their host plant, where they fed and grew. After having reached a certain age and weight (third instar), they became mobile enough to switch host plant if needed, that is, in response to the host plants’ chemical defense (larval movement between host plants has been observed in the field [Kessler and Baldwin 2002] and in the laboratory [Casey 1977; van Dam et al. 2000]). Caterpillars grew exponentially and consumed plant mass proportionally to their weight; this assumption is based on two studies that show that *Manduca sexta* larvae reared on *Nicotiana* feed proportionally to their body weight (Gilmore 1938; Madden and Chamberlin 1945). In reality, the relationship between body mass and consumption might be nonlinear because the efficiency of assimilation might change with body mass, but we do not expect strong nonlinearity and therefore (also to reduce model complexity) used a linear relationship. When the larvae reached their maximum size, they left the plants for pupation and were thus inactive in the model. Caterpillar growth and instar progression were negatively affected by plant defense compounds (fig. 2).

In each time step caterpillars had a certain probability to die. Their mortality rate scaled proportionally with their current host plant’s defense level and inversely with the logarithm of the caterpillar’s body mass. This means that at the beginning of simulations, caterpillar mortality risk was very high but decreased as the caterpillars grew larger.

If the plant defense level reached a certain threshold, caterpillars that were heavy enough moved to another nearby plant. Caterpillars that were too small were not able to move and thus remained on their host plant. The next host plant was chosen randomly among all plants within the movement radius of the larva. The probability of being chosen decreased exponentially with its distance to the larva’s previous host plant. Caterpillars that were in the process of switching plants had a higher probability to die than caterpillars feeding on a plant.

![Figure 2: Growth of *Manduca sexta* larvae. Larvae reared on plants that are unable to produce defense compounds (− def; red line; start: $N = 30$ larvae; end: $N = 16$ larvae) are compared with larvae raised on plants that are maximally defended (+ def; start: $N = 30$ larvae; end $N = 6$ larvae). The photo shows the difference in the size of a larva raised on a defenseless plant (lower larva) relative to one raised on a maximally defended plant (upper larva) after 14 days. Asterisks denote $P$ values: three asterisks indicate $P < .001$, two asterisks indicate $P < .01$, and one asterisk indicates $P < .05$.](image-url)
When a caterpillar encountered a new host plant, it immediately started feeding and thereby inducing defense with the delay time \( \tau \). When a plant is induced, 30\% of the plant’s metabolic resources are invested in defense production; the rest is invested in growth and reproduction.

**Parameterization**

Most of the parameter values were taken from experimental data on *N. attenuata* and *M. sexta*. Model parameters and variables are summarized in table 1. *Manduca sexta* larvae reared on *N. attenuata* leaves grow exponentially in time and feed proportionally to their body weight (Gilmore 1938; Madden and Chamberlin 1945). The average conversion factor of plant to larval biomass is 0.19, and the largest preupal larval mass observed is 10.11 g (van Dam et al. 2001); hence, we set the maximal biomass to 10 g (van Dam et al. 2001a).

Shoot growth rates were derived from noninduced and undamaged, soil-grown, hydroponic *N. attenuata* plants grown for 50 and 20 days in climate rooms, respectively (van Dam and Baldwin 1998; Glawe et al. 2003). Both soil and hydroponically grown plants grew exponentially during those periods and attained relative growth rates of 0.25 and 0.16 g g\(^{-1}\) day\(^{-1}\), respectively. Accordingly, we chose a maximal growth relative growth rate of 0.2 g g\(^{-1}\) day\(^{-1}\). The maximal fresh weight of the aboveground biomass of a plant in the field was estimated to be 500 g (table 1; A. Kessler, personal communication; TRACE document fig. ST6: field data of A. Weinhold; figs. ST1–ST35 are available online).

The parameter value for allocation to defense production is difficult to estimate, since there are only few data to quantify the magnitude of investment on a whole plant basis. Induced *N. attenuata* plants allocate 6\% of their whole plant nitrogen to de novo nicotine production alone (Baldwin 1998). Induced *N. attenuata* plants, however, also produce significantly more protease inhibitors (van Dam et al. 2001b), volatiles (Halitschke et al. 2000; Kahl et al. 2000), phenolics, and sugar esters (Keinänen et al. 2001). Thus, the total allocation to defense production may well be much higher than 6\%, and therefore we tested the model for allocation values ranging from 10\% to 40\% (fig. ST5). In the default case, we assumed that induced plants temporarily allocate 30\% of their growth to the production of defensive compounds when attacked by an herbivore.

We used the commonly seen response times as assessed in the many experiments carried out in the Baldwin lab (Baldwin 1988, 1989; Ziegler et al. 2001) and also response times measured for different plants (Karban and Myers 1989; Underwood 1998; Agrell et al. 2003; Kant et al. 2004) as the potential range for the delay time \( \tau \).

We chose 27 days for the simulation of one generation because we concentrated on the time where interaction between larvae and plants occurred. Because *M. sexta* is both the main herbivore and pollinator of *N. attenuata*, both pollination services and oviposition are motivated by floral scent and nectar (Kessler 2012). Our simulations thus started with plants already in an early flowering stage because at this time oviposition rates started rising.

**Field Experiment**

In a field experiment, we compared the growth trajectories of *M. sexta* larvae feeding on defenseless plants with larvae feeding on well-defended plants. We therefore recorded the growth trajectories of 30 larvae of *M. sexta* per treatment in their native environment in the Great Basin Desert (Utah). We recorded larval growth on two different plant lines: either well-defended wild-type plants or plants of the jasmonate-deficient inverted repeat allene oxide cyclase (irAOC) line. Jasmonic acid accumulation and perception are critical for the activation of most defense responses, and the irAOC line we used in the field study shows a reduction of herbivory-induced jasmonate levels of more than 95\% (Kallenbach et al. 2012; Machado et al. 2013; for further information, see fig. ST1 and “Data Evaluation” in the supplemental material). The results were used to parameterize larval growth.

**Simulation Experiments**

At the start of our simulation, all plants had the same weight of 30 g, and the neonate larvae started with a body mass of 1 mg on randomly selected plants. For the robustness analysis (see the appendix; supplemental material), we also ran simulations where caterpillars started at any time point on randomly selected plants.

We ran simulations with two plants (for test purposes) and with 400 plants. The model was implemented in NetLogo (Wilensky 1999; Tisue and Wilensky 2004). Field data and model output were analyzed with R (R Development Core Team 2016). The NetLogo program is provided in GitHub: https://github.com/PiaBackmann/TIMELY-MODEL. Data are available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.gh2m22t (Backmann et al. 2018).

Two-Plant Scenario. We ran simulations with only two plants and one larva to visualize the scenario discussed in the introduction. Both plants had the same delay time, and one plant (plant A) received a newly hatched caterpillar. We performed simulations with three different delay times: no delay time (\( \tau = 0 \) days), intermediate (\( \tau = 4 \) days), and long (\( \tau = \infty \) days); the latter option equals no defense response.
Main Simulations (400-Plant Scenario). All 400 plants were given either the same delay time for all plants (for the experiments conducted for figs. 4, 5) or random delay times, with \( \tau \in [0, \ldots, 10] \) (for the results shown in fig. 6). Caterpillars were placed randomly on the plants, according to their chosen density.

Genetic Algorithm. The heritable trait for the genetic algorithm was the plant’s delay time \( \tau \). At initialization, the delay times of all plants were drawn from a uniform distribution between 0 and 10 days. The plants that had the largest average shoot biomass at the end of a simulation or generation are considered to have the largest fitness, assuming that larger plants produce more seeds (Aarssen and Taylor 1992; TRACE document fig. ST6). Therefore, in the following generation, the 400 plants were assigned to genotypes (delay times \( \tau \)) proportional to their total shoot biomasses at the end of the preceding generation. This was repeated for 300 generations, with 27 days per simulation. To prevent being caught in local fitness maxima, we represented mutation by adding a random number \( r \) every generation, with \(-3 < r < 3\), to every assigned value of \( \tau \). Simulation experiments were conducted with different larval densities (0–500 larvae). Per larva density, the genetic algorithm was repeated 100 times.

We compared our genetic algorithm with a different one (Stonedahl et al. 2008; see TRACE document) to see whether the results were persistent. We also started the genetic algorithm with all plants having the same delay time to see whether initial setting would affect the final distribution of delay times.

Results

Field Study

Larvae feeding on low-defense plants showed significantly higher growth rates (fig. 2) than larvae feeding on high-defense plants. Furthermore, their instar progression was faster. The fifth instar, for example, was reached 4 days earlier (table ST1; tables ST1–ST7 are available online). Additionally, the maximal weight attained in each instar (table ST1) and survival rates of larvae raised on low-defense plants were higher; 53\% of the larvae survived on low-defense plants and 20\% on well-protected plants, showing that there is a fitness cost for larvae to stay on a well-protected plant.

Two-Plant Scenario

For the scenario with immediate defense production (\( \tau = 0 \) days), the larva died after 7 days, thus before it would have been able to switch plants. The final biomass of plant A was significantly lower than the biomass of its uninfected competitor (fig. 3, left). For the scenario with a long delay time (corresponding to no defense at all), the larva stayed on plant A until pupation and the biomass of the infested plant was reduced by larval feeding, resulting in a lower biomass than the other plant (fig. 3, right). For the scenario with \( \tau = 5 \) days, the larva was driven from the first host plant to the neighboring plant on day 11 of the simulation. The final mass of plant A was higher than the mass of plant B (fig. 3, middle).

Main Simulations (400-Plant Scenario)

We conducted simulations with populations of 400 plants having all the same delay time and 300 larvae to record average larval mortality for different \( \tau \) values. For \( \tau = 0 \) and \( \tau = \infty \) days, on average 21.7\% and 45.8\%, respectively, of all larvae survived until pupation stage. For an intermediate delay time (\( \tau = 4 \) days), on average 50.3\% survived until pupation, and 57.4\% of all caterpillars reached the third instar, which is the instar in which larvae can switch their host plants.

As the next step, we calculated the mean number of interplant movements of larvae per simulation. These switches were either motivated by a high defense level of the host plant or because the host plant was entirely consumed by the larva. For our analysis, we only counted switches motivated by a high defense level because those were a larval reaction to the plant’s defense mechanism. The mean number of switches peaked when the delay time of plants was intermediate (3 days \( \leq \tau \leq 5 \) days; fig. 4). If the plant defended immediately (\( \tau = 0 \) days) or after a very long delay time, the number of switches was significantly smaller (fig. 4).

The higher the defense levels of the host plant, the shallower the growth curves of larvae. As a result, if a larva encountered mostly fast-defending plants during its development, it needed 2–3 days longer to reach its final size and go into the pupation stage (fig. ST2; supplemental material).

Larvae are sufficiently mobile to switch host plants when they have reached a certain size. In our simulations, the larvae moved earliest after 8 days from one plant to another (fig. 5, bottom). Larvae were mobile that quickly only when the delay time of their host plant was intermediate (\( \tau = 4 \) days). In case of feeding on a plant with a delay time of \( \tau = 0 \) days, caterpillars needed on average 2 days longer (10 days) to reach a size where interplant movement was possible because their growth was inhibited by the fast-rising defense levels in their host plants during the early instar phases.

In most of the cases, when switching, larvae moved to one of the other plants in the closer neighborhood, thus direct competitors of the originally infested plant (fig. ST3). A single larva, which survived until pupation, visited on aver-
Figure 3: Plant biomass in one two-plant simulation. Three different \( \tau \) values are shown (left, \( \tau = 0 \) days; middle, \( \tau = 5 \) days; right, \( \tau > 10 \) days). In all cases, plant \( A \) (green line) starts with a freshly hatched larva. In the left panel, it defends immediately and eventually kills the larva at day 7. In the right panel, it reacts so slowly that the larva completes its full life cycle on the plant. Therefore, the plant suffers considerable damage. In the middle panel, the plant reacts after a certain delay. The larva survives, reaches third instar, and thus becomes mobile. It is sent to the competing plant, and plant \( A \) can recover. Plant \( B \) receives more damage in this scenario than the formerly infested plant.
age four to five plants per simulation. Reasons for switching were complete consumption of the host plant (in ~70% of cases) or that the host plant defense level increased over the threshold level (in ~30% of cases). If a plant was infested by a larva and started inducing defense, the critical defense level to repel larvae was reached approximately 4–7 days after induction (fig. 5, top). This time depended on the current growth rate of the plant and of the size of the plant when induction had started.

### Genetic Algorithm

Starting from the random initialization of delay times in generation 1, variance of the delay times decreased and the distribution started to peak around a maximum value. In all cases, a specific dominant delay time $\tau$ emerged. However, it never fully suppressed other delay times. Average delay time depended on initial herbivore density (fig. 6, left); for high densities, the mean of the delay times increased over the generations, and for low densities, the mean delay time decreased. For higher initial herbivore densities, variance of delay times decreased more rapidly, and the final stationary distribution developed more pronounced peaks (fig. 6, right; see also video of the development of the delay time distribution over time in the supplemental material). More or less the same final distributions were obtained when the genetic algorithm started with the same delay time for all plants (TRACE document fig. ST18). All these results were also obtained when using the alternative genetic algorithm (see TRACE document).

To test whether interplant competition was one important driver for longer delay times, we also run the genetic algorithm for plants that were not affected by interplant competition. Here it showed that without competition, optimal delay times should be as short as possible (fig. ST4).

### Discussion

Our results show that in highly competitive environments, plants can realize a fitness benefit by delaying induction of defenses. Expelling an herbivore to one of the neighbors may be the best chance for an infested plant competing with uninduced conspecifics to compensate for the costs of induction, despite the amount of leaf loss. Consequently, there is no strong selection for reducing response times, which may explain the large variety in delay times found in plants. Therefore, the postulation that the delay in deploying antiherbivore defenses is unequivocally costly (Herms and Mattson 1992) can be rejected.

Our results indicate that instead of being as fast as possible, it is more important that defense induction is timely. For *Nicotiana attenuata*, the peak in defense production should co-occur with a window of sensitivity for the herbivores. Behavior of *Manduca sexta* larvae is most sensitive to the induced defenses of *N. attenuata* when they are in their third instar (van Dam et al. 2001a). From this instar onward, the larvae start to cause significant damage to the plant and are large enough to reduce the risks of starvation or predation while searching for a less defended host plant (Gilmore 1938; Haccou and Hemerik 1985; fig. 2). For the host plant, it would be most beneficial if larvae in this instar are driven to the neighbor plants and start feeding on their competitor. We could also show that for plants that are not exposed to interplant competition, the opposite is true and shorter delay times are preferable (fig. ST4).

Not producing defense at all results in a higher plant lethality; larvae would stay on the host plant during their whole life cycle and cause severe damage (fig. 6, right). In our simulations, on average 171 out of 400 plants die because of herbivory and competition if no defense is produced, whereas only 117 plants die if plants immediately produce defense. However, for a plant inducing defense immediately after an attack, the defense level peaks too early and more larvae will die before reaching the third instar. If the plant would be growing in isolation, this would be the best option. However, in natural environments, *N. attenuata* and many other plant species that mass germinate (such as spring annuals) grow in dense cohorts, and defense induction comes with loss in competitive abilities (van Dam and Baldwin 1998;
Glawe et al. 2003). Thus, in competitive environments, producing defense early means a loss in growth rate that can hardly be compensated for in a relatively short growth season. Both extremes—immediate induction or no induced defense at all—seem to be suboptimal strategies. Our model shows that one possible way out of this dilemma is to induce at a time that larvae are causing the most of the damage and are likely to move to neighboring plants.

During the first two instars, when larvae are small and not able to move between plants, the damage they cause is negligible because damage scales with caterpillar size. In this stage, the effect of the 30% growth loss when being induced is more relevant than the loss of biomass due to larval feeding.

For higher initial herbivore densities—for example, when every plant receives at least one larva—slowly reacting plants will have an initial growth advantage because they do not invest in producing defense first. In all plants, the defense maximum peaks 4–7 days after induction started (fig. 5, top), whereas larvae stay at least 8–10 days on the host plant where they hatched. Therefore, if the plant defends without delay, it will be at a maximum defense level before the larva becomes mobile, and the mobility phase will be reached even later (fig. 5, bottom).

Under high herbivore pressure, selection by herbivore feeding as represented by our genetic algorithm leads to higher delay times. Under these conditions it is a better strategy to wait, whereas for low herbivore pressure a broad range of delay times can coexist. This is underscored by the fact that under higher herbivore pressures, variance of delay times in the population decreased more rapidly, and the curve of the stationary distribution of $r$ values showed a sharper peak.

To test whether the resulting stationary distribution of delay times was robust, for each herbivore density we used the resulting peak values of $r$ to start a genetic algorithm.
where all plants were assigned to this same value of $\tau$. After 300 generations, the genetic algorithm leveled out to the same resulting frequencies of the stationary distribution found for the genetic algorithm, starting with the uniform distribution of $\tau$ (fig. ST18). This means that the fitness related to delay times was always frequency dependent, leading to a distribution of delay times rather than resulting in a single dominant time or revealing time intervals that were completely suppressed. In their natural habitat, herbivore densities change from season to season. Therefore, there will be no distinctive and constant selection pressure toward shorter or longer delay times. We therefore hypothesize that the observed variance in delay times of individual plants should be high in natural populations.

In order to test the model predictions, the variance in reaction times (kinetics) of $N. \text{ attenuata}$ in natural populations should be measured by damaging the plants mechanically and simultaneously adding larval saliva to the produced wound. Then, the concentrations of different defense compounds within the leaf tissue can be measured at different time points after elicitation. This outcome can then be compared with the predictions of the TIMELY model (fig. ST8 shows how the experimental corroboration could be done).

Plants can perceive attacks by herbivores as well as the strength of their local competition. Consequently, they may also phenotypically adapt their delay time to the actual level of competition. By performing experiments with different plant densities, we could show that the severity of herbivore load is the most important factor for the resulting delay times. However, without interplant competition, the resulting delay times would be as short as possible (fig. ST4). Therefore, the evolution on optimal delay times depends on both the number of attacking herbivores and the severity of interplant competition.

One more possible mechanism that might interfere with the mechanism represented in our model is the production of volatile organic compounds that are produced once larvae feed on a plant and that might induce chemical defense in neighboring plants. However, such secondary induced defense has not been observed so far for $N. \text{ attenuata}$, and we therefore ignored this effect. Still, this priming of neighbors is included as an option in our model and thus can be explored for situations and species where such priming can be assumed to play a role (see ODD in the supplemental material, p. 28).

We designed and calibrated our model with data from $N. \text{ attenuata}$ as annual plant species and $M. \text{ sexta}$ as its main herbivore. However, it can be easily adapted for other annual plant-insect herbivore systems. In order to do so, we tested our model for a range of settings that differ from the $N. \text{ attenuata}–M. \text{ sexta}$ system (see the appendix; supplemental material). Here we found that the generally held assumption that damage done by herbivores before the defense levels are induced is always costly in terms of fitness cannot be sustained. In that sense, the model addresses a general hypothesis that now is refuted.

Whether a particular plant species may benefit by delaying induction depends strongly on the characteristics of the herbivores. Delaying defenses within the plants’ lifetime may be profitable only when the plant is attacked by an herbivore that is mobile and selective and feeds increasingly more over time (Tuomi et al. 1994; Underwood 1999; van Dam et al. 2001a). Lepidopteran larvae are good examples of such herbivores, but these criteria may also apply to other arthropod herbivores that have two discrete
generations within the plants’ lifetime, of which the second is more abundant (Underwood 1999). Furthermore, our model was designed to analyze the fitness effects of time delays within one growing season; however, it could be adapted to analyze the consequences in slower-growing long-lived plants, such as trees. Trees may have delayed induced defenses that are not expressed until the next growing season after herbivore attack (Haukioja 1980; Rhoades 1983; Karban and Baldwin 1997). Such long-term delays in perennial plants may, depending on the mobility and selectivity of the herbivores, drive fluctuations in insect populations over the years (Underwood 1999); hence, the benefits of such long-term delays can be assessed only over several reproductive cycles of the plant.

To conclude, for induced defense responses in plants, “the faster the better” is not necessarily true: delays in inducible defenses can evolve to be part of the set of successful defense strategies when the plants use a small proportion of their biomass to rear larvae for reducing competition from their neighbors/competitors. Herbivore-mediated competition among plants via the evolution of delays in response timing has not been described previously and provides an intriguing example for the adaptive behavior of plants (Meyer et al. 2014), in particular as it involves life-history traits of other organisms, in this case caterpillars. The behavior of plants is no less adaptive than that of animals, but the potential for adaptation seems to be limited by their sessile nature. We have shown that plants can include insect herbivores into their fitness-maximizing strategy and thereby modify the local forces of competition to their own benefit. This indicates the high potential of merging chemical ecology and behavioral ecology via individual-based models of populations.

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Appendix

Robustness Analysis

We used the rationale of robustness analysis (Levin 1966; Grimm and Berger 2016), using unrealistic settings in terms of parameters in functions to try to break a model and thereby explore robustness of the main findings of a model and identify key components of a model system’s organization. We therefore ran further simulations with different plant densities, mobility types of larvae, and different defense investment proportions of plants to see whether they may lead to similar results, even for systems that differ from the Nicotiana attenuata–Manduca sexta system (all performed robustness experiments are listed in table ST2).

The robustness experiments show that for most settings, the delay time \( \tau \) remains positive (see table ST2; figs. ST5–ST7). This holds true as long as the following prerequisites are fulfilled: herbivores must be mobile, plants must compete with one another (inter- or intraspecific), and plants must share their herbivores. We therefore are convinced that our main conclusion holds: the generally held assumption that damage done by herbivores before the defense levels are induced are always costly in terms of fitness cannot be sustained. In that sense, the model addresses a general hypothesis that now is refuted.

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Above left, *Nicotiana attenuata*, the studied plant species, in its natural habitat. Above right, *Manduca sexta* feeding on a tobacco plant. Photo credits: Pia Backmann. Below, view of the Great Basin Desert, habitat of the study plant, the wild tobacco, *N. attenuata*. Photo taken at the field station of the Max Planck Institute for Chemical Ecology at Brigham Young University’s Lytle Ranch Preserve in southwestern Utah. Photo credit: Danny Kessler.