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Distance-dependent shifts in net effects by an unpalatable nettle on a palatable plant species

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ABSTRACT

We examined whether the relative importance of facilitation and competition effects by an unpalatable perennial (*Urtica thunbergiana*) on a palatable annual (*Persicaria longiseta*) change with the spatial distance between them in a long-term deer-grazing habitat. Survivorship, growth, size, and fecundity of *Persicaria* were recorded at 1–2-month intervals during growing seasons in 2 successive years and were compared among individuals located in different positions relative to the canopy of *Urtica*: at the centre, internal edge, external edge, and far from the canopy. Survivorship of *Persicaria* was significantly higher at the centre of the *Urtica* canopy than outside the canopy early in the growing season in both years. No significant differences in *Persicaria* growth were observed among the four positions in most periods, except in one when growth was significantly higher at the centre, internal, and external edges of the canopy compared to outside the canopy. We found spatial shifts in the net effects of *Urtica* on *Persicaria* fecundity, from positive effects under the canopy centre to negative effects under the external edge of the canopy in the first year, and from negative effects under the centre to positive effects under the external edge in the second year. These results demonstrate that the relative importance of positive and negative effects of *Urtica* on *Persicaria* vary temporally within and among years and spatially around a single *Urtica* plant. Spatiotemporal variation in plant interactions may be attributable to annual and seasonal variation in vegetation productivity and grazing pressure.

**Key words:** competition, facilitation, deer-grazing, Nara Park, *Persicaria longiseta*, *Urtica thunbergiana*
1. Introduction

Plant–plant interactions, such as facilitation and competition, play a major role in shaping the structure of plant populations and communities (Callaway et al., 2005; Callaway et al., 2000; Callaway and Walker, 1997; Hacker and Gaines, 1997; Purves and Law, 2002). Facilitation and competition often act simultaneously (Callaway, 1994), and thus the net effect of a particular plant–plant interaction is determined by the relative strengths of the facilitative and competitive effects (Menge, 2000). The intensity of interactions among sessile plants gradually decreases with distance between individuals (Matlack and Harper, 1986; Weiner, 1984). How the interactions decrease with inter-plant distance and whether the intensities of facilitation and competition decrease similarly or differently depends on the species involved and the environmental context. Due to such contingency, it is less predictable whether the net effect of a plant–plant interaction changes with the distance between individuals.

Unpalatable plants with traits that deter animals, including toxicity, spines, and thorns, have potentially strong facilitative effects on palatable plants through indirect protection (Milchunas and Noy-Meir, 2002; Oesterheld and Oyarzábal, 2004; Osem et al., 2007; Rebollo et al., 2002; Smit et al., 2006). However, considerable evidence has demonstrated that such facilitation effects are balanced under some circumstances through competition between the palatable and unpalatable species (Alberti et al., 2008; Graff et al., 2007; Veblen, 2008). The balance between facilitation and competition depends on a variety of factors, e.g., grazing intensity (Graff et al., 2007; Levenbach, 2009), the defensive ability of the unpalatable plant (Rebollo et al., 2005), the grazing tolerance of the palatable species (Vandenberghe et al., 2009), community productivity (Goldberg et al., 1999; Rebollo et al., 2005), soil humidity (Veblen, 2008), and plant performance such as growth, survival, and reproduction (Maestre et al., 2005). Some studies
have demonstrated that temporal variation in the effects of these factors can cause temporal
shifts in the balance between competition and facilitation (Alberti et al., 2008; Veblen, 2008).
However, few studies have examined spatial shifts in the net effects of unpalatable plants,
especially with respect to finer spatial scales around each unpalatable plant (but see Gómez et al.
2001).

We assumed that the strength of facilitative and competitive interactions and the net effect
of an unpalatable plant changes as a function of the distance between palatable and unpalatable
plants (Fig. 1). The net effect of an unpalatable plant on a palatable plant at each distance is
expressed as a combined function of facilitative and competitive effects. We hypothesised four
possible patterns for the net effect of an unpalatable plant changing with spatial distance. First,
when facilitative effects are greater than competitive effects, regardless of spatial distance, and
the two effects decrease similarly with distance, the net effect is generally positive and simply
decreases to zero over space (Case 1, Fig. 1a). The reverse case is also possible (Case 2, Fig.
2b). Most of previous studies have assumed that plant–plant interactions correspond to Case 1
or 2, but more complicated outcomes can occur when the strengths of competition and
facilitation change differently with spatial scale. For example, positive effects might be strong
below the canopy of the unpalatable plant and decrease abruptly from the centre to the edge,
whereas negative effects operate at an intermediate level below the canopy and decrease
gradually outside of the canopy (Case 3, Fig. 1c). In this case, the net effect is positive at the
centre of the canopy and negative near the edge of the canopy. This combination can arise when
competitive suppression that may be due to belowground competition operates at an
intermediate level at a wide range around the unpalatable canopy, whereas positive effects are
strong only at the centre of the unpalatable canopy because herbivores occasionally graze on
plants within the unpalatable canopy by inserting their heads through it. The opposite scenario
is also possible, that is, the net effect may be negative at the centre of the canopy and positive near the edge of the canopy. This pattern can arise when competitive suppression through severe shading is strong, whereas positive effects operate at an intermediate level at a wide range around the unpalatable canopy because herbivores generally avoid grazing plants around unpalatable plants but do occasionally (Case 4, Fig. 1d).

To test these hypotheses, we examined the spatial shift in net effects of unpalatable nettles (Urtica thunbergiana) on palatable plants (Persicaria longiseta) at fine spatial scales around unpalatable individuals in Nara Park, Japan, where a dense population of sika deer (Cervus nippon) has persisted over a long time period. Our previous study demonstrated that Urtica plants have protective effects on neighbouring Persicaria plants from deer grazing by concealing them within the Urtica canopy (i.e., associational resistance), but they also have negative effects on small neighbours, probably due to competition for light (Suzuki and Suzuki 2011). In this study, survivorship, growth, and fecundity of Persicaria specimens near Urtica plants were recorded at 1–2-month intervals in 2 successive years. We classified P. longiseta individuals into four distance classes based on the relative distance from the centre of the nearest Urtica canopy, and compared performance among the distance classes. Our specific question was whether the net effect on palatable plants shifted spatially from positive to negative around an unpalatable plant (Case 3 or 4).

2. Materials and methods

2.1. Study site

The study was conducted at Nara Park (34°41’ N, 135°50’ E), in Nara Prefecture, western Japan
(see Suzuki and Suzuki, 2011 for details). Sika deer (*Cervus nippon* Temminck) populations have been distributed in this park for more than 1200 years, as they are protected for religious reasons. Over the last decade, the population density of deer has reached extremely high levels (ca. 1200 individuals / 5 km² in the park; Foundation for the Protection of Deer in Nara Park, 2011). The study areas were established in a shady-moist environment in the understory of a sparse Japanese cedar forest. Tall, erect species (e.g., *Erechtites hieracifolia* (L.) Raf., *Hypolepis punctata* (Thunb.) Mett. ex Kuhn, and *Urtica thunbergiana*) were particularly abundant in this environments (Suzuki et al., 2009). These tall species are unpalatable to deer because of their physical and chemical defences (Kato et al. 2008).

### 2.2. Study species

We examined the interactions between a palatable species, *Persicaria longiseta* (De Bruyn) Kitag., and an unpalatable species, *Urtica thunbergiana*. *Persicaria longiseta* is an annual herb distributed widely throughout East Asia and is common to roadsides, farms, edges of rice fields, and gardens in Japan. Seed germination occurs primarily from the end of March to April. During the growing season, plants form a number of lateral shoots from axillary buds proximal to the base. Flowering can start from July but occurs mostly from September onwards and peaks in October at the study site. Reproductive plants produce a large number of small seeds (ca. 1.5 mg per seed) but do not regenerate vegetatively. Fruit maturation and seed dispersal occur from July to December. After reproduction, the plant dies by mid-December. Deer graze individuals of *Persicaria* in Nara Park and the grazed plants exhibit inherently dwarf morphology, with shorter shoots and smaller leaves compared with populations with no grazing history (Suzuki, 2008). The small stature of *Persicaria* is likely a grazing response (Suzuki,
The nettle, *Urtica thunbergiana* is a perennial herb distributed within forest understories throughout central and southern Japan. This species possesses stinging hairs containing toxins on its stems and both surfaces of leaves, which can protect them from deer grazing (Kato et al., 2008). During a growing season, plants develop a number of shoots proximal to the base. Most aboveground parts wither during the winter season (December–March). The population of *U. thunbergiana* in Nara Park has evolved an extremely higher density of stinging hairs compared to those in ungrazed areas as a consequence of local adaptation to heavy grazing (Kato et al., 2008). The maximum plant heights of *Persicaria* and *Urtica* in Nara Park are ca. 10 and 50 cm, respectively, and thus *Urtica* plants seemingly suppress *Persicaria* neighbours through competition for light.

### 2.3. Field censuses

In May 2007, we randomly selected 10 *Urtica* individuals and established ten 50 × 50-cm plots, each positioned with an *Urtica* individual at the centre. Some plots included several *Urtica* specimens. In June 2008, we established five 100 × 100 cm plots that contained several *Urtica* individuals. We drove metal stakes into the ground at the corners of all plots. In total, 273 and 512 individuals of *Persicaria* and 17 and 18 individuals of *Urtica* were observed in 2007 and 2008, respectively (Table 1, 2). All plants of the two species in each plot were mapped by recording the $x,y$-coordinates of stems. Plant fates and sizes were recorded in May, July, September, and October 2007 (ten 50 × 50-cm plots), and in June, August, and October 2008 (five 100 × 100-cm plots). In October of both years, the flowers on each *Persicaria* plant were also counted. To express plant size, we
measured the maximum diameter and the perpendicular diameter of the canopy ($D_1$ and $D_2$, respectively) covered by the aboveground parts of each plant of the two species during each census. As a measure of plant size, we calculated the area covered by each plant ($D_1 \times D_2 \times \pi/4$).

2.4. Analysis

To standardize distances between palatable and unpalatable plants, we calculated a relative distance between them by dividing the actual distance (from stem to stem) by the canopy size of the unpalatable plant. The net effect of unpalatable plants on palatable plants was predicted to vary nonlinearly with relative distance. One approach for modelling such a nonlinear relationship is to fit a nonlinear function of the relative distance. However, models with nonlinear functions might be too complex, making the interpretation of estimated parameter values difficult. Therefore, we used a categorical variable for relative distance as an explanatory variable. The positions of Persicaria plants were classified into four categories based on the distance from the centre of the nearest Urtica canopy and the relative size of the Urtica canopy as follows:

- Centre of the canopy (position 1, P1): an area within one-half of the radius of the Urtica canopy (relative distance < 0.5).
- Internal edge of the canopy (position 2, P2): an area under the canopy between the halfway point of the canopy radius and the edge of the canopy (0.5 < relative distance ≤ 1).
- External edge of the canopy (position 3, P3): an area from the limit of the radius of the Urtica canopy to a distance 1.5 times the canopy radius (1 ≤ relative distance < 1.5).
- Far from the canopy (position 4, P4): an area beyond 1.5 times the canopy radius of the *Urtica* plant (1.5 ≤ relative distance). The performance of *Persicaria* at P4 was regarded as a control because we assumed that plants at P4 were rarely affected by *Urtica*.

We also analysed *Urtica* effects using five position categories, including an outer position (position 5: 2 ≤ relative distance), but no significant differences were found between positions 4’ (1.5 ≤ relative distance < 2) and 5, with the exception of two cases for which the general trends were similar to the results of analysis with four positions. Therefore, we concluded that *Urtica* rarely affected *Persicaria* individuals at P4.

By assuming that the shape of an *Urtica* canopy was a circle, the radius of the canopy was calculated as $\sqrt{\text{canopy size}/\pi}$. The position category into which each plant was classified varied between censuses because *Urtica* canopy size changed during the growing season. Thus, the position categories of plants were determined during each census.

Survivorship and changes in plant size during periods between censuses, survivorship during the whole growing season (May–October 2007 and June–October 2008), and final plant size and fecundity (as the number of flowers) in October were analysed. For statistical analysis, we used a generalised linear mixed model inferred by a Bayesian approach (Barker, 2010; McCarthy, 2007). Bayesian inference is one of the most effective and robust methods to infer parameters in a hierarchical model, especially when the number of replicates is limited or unbalanced (Clark 2005; McCarthy 2007). The model formula is

$$y_{ijk} \sim f(\mu_{ij})$$

$$g(\mu_{ij}) = \mu_0 + pos_i + plot_j,$$

where $y_{ijk}$ is the observed performance (i.e., survivorship, growth, plant size, or number of flowers) of *Persicaria* individual $k$ at position $i$ in plot $j$, and it is assumed to follow an
appropriate distribution \( f(.) \) [with a variance parameter if required], \textit{binomial} for survivorship, \( normal \) for growth, \textit{Gamma} for plant size, and \textit{Poisson} for the number of flowers; \( \mu_{ij} \) is the mean at position \( i \) in plot \( j \); \( g(.) \) is an appropriate link function, \textit{logit} for survivorship, \textit{non-transformed} for growth, and \textit{log} for plant size and number of flowers; \( g^{-1}(\mu_0) \) [\( g^{-1}(.) \) is the inverse function of \( g(.) \)] is equal to a mean performance of \textit{Persicaria} that are not affected by unpalatable plants; a \( pos_i \) is an effect of position \( i \) as a fixed factor; and \( plot_j \) is an effect of plot \( j \) as a random factor following a normal distribution with mean of 0 and variance of \( \sigma^2_{plot} \). In this study, \( pos_4 \) (effect of position 4) was set to 0 by assuming no effects of \textit{Urtica} at P4. Then, position effects \( pos_i \) (\( i = 1, 2, 3 \)) indicate the performance of \textit{Persicaria} plants relative to those at position 4, which could be equivalent to the relative neighbour effects (NUE, Markham, J. H., Chanway, 1996). A positive \( pos_i \) indicates that the net effect by \textit{Urtica} is positive at position \( i \).

Posterior means and credible intervals (Bayesian confidence intervals) of parameters were estimated by Gibbs sampling, a Markov chain Monte Carlo method (MCMC). When the lower limit of the credible interval of \( pos_i \) is higher than 0, the performance of \textit{Persicaria} at position \( i \) is significantly greater than that at position 4, which indicates that the net effect exerted by \textit{Urtica} on \textit{Persicaria} at position \( i \) is positive.

Gibbs sampling was implemented using OpenBUGS 2.0 (Thomas, 2005) run through the statistical package R v. 2.6.2 (R development Core Team, 2008) using the contributed package, \textit{R2WinBUGS} (Sturtz et al., 2005). Convergence of three independent chains was assessed via the Gelman–Rubin statistic, and sufficient burn-in periods and thinning rates were used to ensure satisfactory sampling of the posterior distributions.

3. Results
The observed values of the performance of \textit{Persicaria} are shown in Table S1 in the Appendix.
In general, survivorship of *Persicaria* tended to be higher under the *Urtica* canopy than outside of it (Fig. 2). In 2007, *Persicaria* survivorship in May–July was in the rank order P1 > P2 > P4 > P3 (Fig. 2a). The effect at P1 (centre of the *Urtica* canopy) was significantly positive. No significant effects were found in July–September (Fig. 2b). Survivorship fell into the rank order P1 > P3 > P2 > P4 in September–October, and effects at P1, P2 and P3 were significantly positive (Fig. 2c).

In 2008, survivorship fell into the rank order P1 > P2 > P3 > P4 in June–August (Fig. 2d). The effect on survivorship at P1 was significantly positive. No significant effects were found in August–October (Fig. 2e).

Effects on *Persicaria* growth were not obvious during most periods. The growth of *Persicaria* in May–July and July–September in 2007 did not differ among groups (Fig. 3a, b). Growth was highest at P1 and lowest at P4 in September–October in 2007 (Table 1), and effects at P1, P2, and P3 were significantly positive during this period (Fig. 3c). In 2008, growth tended to be low at P1 and high at P3 in June–August and August–October, although these differences were not significant (Fig. 3d, e).

Throughout the growing season (May–October), the effect on survivorship was significantly positive at P1 in 2007 and 2008 (Fig. 4a, d). Final *Persicaria* plant size revealed a rank order of P1 > P2 > P4 > P3 in 2007 (Fig. 4b), whereas a rank order of P4 > P2 > P3 > P1 was observed in 2008 (Fig. 4e). The effects at P1 and P2 on plant size were significantly positive in 2007, but no significant effect was observed in 2008.

A clear spatial shift in the effects on *Persicaria* fecundity was observed, although the directions were quite different between the two years. In 2007, the effects on the number of flowers produced per *Persicaria* individual were significantly positive at P1 and P2 and significantly negative at P3 (Fig. 4c). In 2008, the effect at P1 was significantly negative, and
that at P3 was significantly positive (Fig. 4f).

4. Discussion

This study showed that the net effect of unpalatable *Urtica* plants was basically positive for the growth and survivorship of palatable *Persicaria* individuals located under the *Urtica* canopy. *Persicaria* survivorship throughout the growing season and final plant size were higher under *Urtica* canopies than outside (Case 1, Fig. 1a). However, facilitative effects on survivorship were not observed in several periods (July–Sept. 2007 and Aug.–Oct. 2008), nor were positive effects found on growth over most periods, suggesting that the net effect of *Urtica* fluctuated during the growing seasons. Moreover, we found spatial shifts in the effects of *Urtica* on *Persicaria* fecundity, with the effect shifting from positive to negative (Case 3, Fig. 1c) or from negative to positive (Case 4, Fig. 1d) at fine scales around the unpalatable plants. Based on these results, we suggest that the relative intensity of competition and facilitation from *Urtica* on the performance of *Persicaria* varies temporally within and among years and spatially at fine scales around individual plants.

Our results suggest that *Urtica* had simultaneous competitive and facilitative effects on *Persicaria*. Additionally, we recently conducted a field experiment that combined *Urtica* removal and deer exclusion, and found that *Urtica* enhanced the growth of *Persicaria* outside exclusion areas, whereas *Urtica* reduced *Persicaria* growth, survival, and reproduction inside exclusion areas (Suzuki and Suzuki, 2011). These results confirm that *Urtica* suppressed *Persicaria* competitively in addition to protecting it from deer grazing (associational resistance).

The relative importance of facilitation is often highest during periods when grazing pressure
is heaviest, then decreases with grazing pressure (Graff et al., 2007). Seasonal variation in grazing intensity may be caused by changes in herbivore behaviour and irregular movements of herbivore populations (McNaughton, 1984). Seasonal variation in grazing intensity can also be caused by changes in vegetation productivity (plant biomass) (McNaughton, 1985). Rich plant biomass leads to lower grazing intensity because herbivores can obtain sufficient food material within limited areas, but also leads to an increase in aboveground competition between neighbour plants (Goldberg et al., 1999). The current study and our previous work (Suzuki and Suzuki, 2011) show the lack of facilitation effects during the summer when plant biomass is highest at the study site. Moreover, facilitative effects of Urtica were weaker during a year with relatively higher productivity (i.e., 2008). Although the difference in Urtica effects between years may have been attributable to the difference in plot size between years, because some Persicaria individuals in position 4 could have been farther from Urtica specimens, our generalised linear mixed model analysis that included an additional outer position (position 5) indicated that the effects of Urtica were minimal outside the canopy. Therefore, the difference in plot size had little effect on our results for positions outside the canopy (i.e., 1.5 < relative distance < 2).

Our results demonstrate that the effects of facilitation and competition by Urtica plants on neighbouring Persicaria operated at different spatial scales. Consequently, the balance of positive and negative effects shifted between different distance categories around Urtica plants. In 2007, Urtica effects on Persicaria fecundity were positive near the centre of the canopy but negative at the external edge. A similar pattern was observed for final plant size, although it was not significant. Hence, these results indicate that facilitative effects operated over a small range near the canopy centre, but negative competitive effects operated over a longer range in the area defined by the external edge of the canopy. The spatial shift in Urtica effects on Persicaria
fecundity was also found in 2008, but the direction of the shift was reversed from that in 2007. The shift from a negative effect at the centre of the canopy to a positive effect at the external edge of the canopy suggests that competition exceeded facilitation under the canopy, whereas facilitative effects operated at the external canopy edge where competitive effects became relaxed. As suggested above, we predict that the reverse patterns might be caused by annual variation in vegetation productivity, which is influenced by weather patterns. In 2007, precipitation was lower than average in August (94 vs. 116 mm) and September (95 vs. 170 mm). The unusually dry conditions in late summer might have resulted in reduced plant growth and increased grazing pressure during this season. Facilitative effects on fecundity in October might have increased significantly under the canopy and declined drastically outside the canopy, whereas weak competition for soil water operated over larger ranges. In contrast, the precipitation pattern in 2008 (162 and 163 mm in August and September) was comparable to that in an average year. Consequently, the weather conditions would have been suitable for plant growth, and competitive interactions (especially aboveground) might have increased under the unpalatable canopy, whereas weak facilitative effects might have operated over a spatial range larger than the canopy radius.

Spatial shifts in net effects around a single plant have also been reported in other types of ecosystems. In arid environments, shade effects (which inhibit soil evaporation) provided by a shrub’s canopy facilitate establishment, survivorship, and reproduction of other plants (Tirado and Pugnaire, 2003). The spatial range of this kind of facilitation by a single plant is likely smaller than the canopy radius (Barbier et al., 2008). In contrast, competition for water resources in arid environments can operate over scales larger than the canopy radius because shrubs have widely spreading root systems (Barbier et al., 2008; Meyer et al., 2008). In contrast, van de Koppel et al. (2006) found that Spartina alterniflora provided large-scale facilitation and
small-scale competition on forb species on cobble beaches. This exertion of bidirectional effects by individual plants may contribute to the spatially heterogeneous distributions and performance of recipient species, heterogeneous community structure, and self-organised patterns of vegetation structure (van de Koppel et al., 2006).

Although many studies have documented temporal and spatial variations in the balance between facilitation and competition along gradients of environmental stress (Alberti et al., 2008; Callaway, 1994, 1997; Graff et al., 2007; Menge, 2000; Oesterheld and Oyarzábal, 2004; Russell and Fowler, 2004; Vandenberghe et al., 2009; Veblen, 2008), they have simply concluded that the net effects of nurse plants are positive, negative, or neutral for recipient plants within particular environmental conditions. In contrast, our study revealed that the net effect of an unpalatable plant can vary spatially, even around a single nurse plant.

5. Conclusions

The balance between the positive and negative effects exerted by unpalatable plants on palatable plants can shift both temporally and spatially at fine scales. We demonstrated the complicated nature of plant–plant interactions, which contributes to the heterogeneous structure of plant communities. As plants are sessile after germination, all interactions among them inevitably depend on the distances separating individuals rather than the presence/absence of neighbours. Therefore, to fully understand the ecological and evolutionary processes of plant species, one must consider the spatial scales at which plant interactions operate.

Acknowledgements

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Appendix. Supplementary data:

Table S1 Survivorship, growth, final plant size, and fecundity of *P. longiseta* plants in the four positions.
References


Suzuki, R.O., 2008. Dwarf morphology of the annual plant *Persicaria longiseta* as a local
adaptation to a grazed habitat, Nara Park, Japan. Plant Species Biol. 23, 174-182.


Table 1. Total number of Urtica individuals and mean radius of their canopy (standard deviation between brackets) in each census.

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<th>Number of plants</th>
<th>Radius of canopy (cm)</th>
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<tr>
<td><strong>2007</strong></td>
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<tr>
<td>May</td>
<td>11</td>
<td>15.0 (4.6)</td>
</tr>
<tr>
<td>July</td>
<td>11</td>
<td>21.0 (7.4)</td>
</tr>
<tr>
<td>September</td>
<td>12</td>
<td>9.0 (4.1)</td>
</tr>
<tr>
<td>October</td>
<td>16</td>
<td>10.0 (5.0)</td>
</tr>
<tr>
<td><strong>2008</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>June</td>
<td>17</td>
<td>9.1 (5.5)</td>
</tr>
<tr>
<td>August</td>
<td>16</td>
<td>11.8 (6.6)</td>
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<tr>
<td>October</td>
<td>15</td>
<td>9.7 (7.1)</td>
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Table 2. Total number of *Persicaria longiseta* individuals at each position $i$ in each census.

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<th>Position 2</th>
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<td></td>
<td></td>
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<tr>
<td>May</td>
<td>16</td>
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<td>15</td>
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<tr>
<td>October</td>
<td>11</td>
<td>16</td>
<td>12</td>
<td>20</td>
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<tr>
<td><strong>2008</strong></td>
<td></td>
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<td></td>
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<tr>
<td>June</td>
<td>25</td>
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<td>August</td>
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Figure captions

Fig. 1  Possible shifts in the net balance between positive and negative effects at fine spatial scales. (a) Positive effects are greater than negative effects and the rates of decrease in influence with increasing distance from the unpalatable plant are similar; (b) negative effects are greater than positive effects and the rates of decrease with increasing distance from the unpalatable plant are similar between effects; (c) positive effects are greater than negative effects at the centre of the unpalatable plant canopy but decrease faster than the negative effects with increasing distance from the unpalatable plant; (d) negative effects are greater than positive effects at the centre of the unpalatable plant canopy but decrease faster than the positive effects with increasing distance from the unpalatable plant. Dashed, dotted, and solid lines indicate positive, negative, and cumulative (net) effects, respectively. The bold horizontal bar indicates the area covered by the canopy of an unpalatable plant.

Fig. 2  Effects of *Urtica* on *Persicaria* survivorship at position *i*, *pos*, in the periods May–July, July–September, and September–October 2007 (a–c), and June–August and August–October 2008 (d, e). Black circles are estimated means, thick vertical lines are 50% credible intervals, and error bars are 95% credible intervals. As position effects on the performance at P1-3 were estimated as differences from P4, the position effects for P4 (*pos*4) that were set to 0 (white circles) are shown for comparison with the hypothetical patterns in Fig. 1 (same for following figures). The estimated performance at position *i* (see Table S1) is calculated as $g^{-1}(\mu_0 + pos_i)$ (see Analysis), and then $\mu_0$ is equal to $g([\text{performance at the position 4}])$.

Fig. 3  Effects of *Urtica* on *Persicaria* growth at position *i*, *pos*, in the periods May–July,
July–September and September–October 2007 (a–c), and June–August, and August –October 2008 (d, e). Black circles are estimated means, thick vertical lines are 50% credible intervals, and error bars are 95% credible intervals.

**Fig. 4** Effects of *Urtica* on *Persicaria* at position $i$, $pos_i$, in terms of survivorship during whole growing seasons (May–October) in 2007 and 2008 (a, d), final plant sizes (b, e), and number of flowers per plant (c, f) in 2007 and 2008. Black circles are estimated means, thick vertical lines are 50% credible intervals, and error bars are 95% credible intervals.
Distance from centre of the canopy of unpalatable plants
Estimated position effect on survivorship

(a) May-July
(b) July-Sept.
(c) Sept.-Oct.
(d) June-Aug.
(e) Aug.-Oct.

Fig. 2
Fig. 3

Graph showing estimated value of position effect on growth for different periods:

- 2007:
  - May-July
  - July-Sept.
  - Sept.-Oct.

- 2008:
  - June-Aug.

Axes:
- Y-axis: Estimated value of position effect on growth
- X-axis: Position (P1 to P4)

Data points and error bars represent the estimated values for each period.
Fig. 4

(a) Survivorship
(b) Plant Size
(c) Number of flowers
(d) Survivorship
(e) Plant Size
(f) Number of flowers

Estimated value of position effect on plant performance
Appendix Survivorship, growth, final plant size, and fecundity of *P. longiseta* plants in four positions relative to the *Urtica* canopy. Posterior mean, and lower and upper 95% credible intervals (between brackets) estimated by MCMC are shown. Bold values indicate significant differences from position 4 (i.e., the position effect deviated significantly from zero in Fig. 2-4).

<table>
<thead>
<tr>
<th></th>
<th>P1</th>
<th>P2</th>
<th>P3</th>
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<tr>
<td><strong>Survivorship</strong></td>
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<tr>
<td>2007</td>
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<tr>
<td>May-July</td>
<td><strong>1.00 (0.91, 1.00)</strong></td>
<td>0.86 (0.71, 0.95)</td>
<td>0.74 (0.56, 0.90)</td>
<td>0.75 (0.52, 0.90)</td>
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<tr>
<td>July-Sep</td>
<td>0.43 (0.14, 0.78)</td>
<td>0.54 (0.24, 0.84)</td>
<td>0.60 (0.26, 0.88)</td>
<td>0.50 (0.16, 0.84)</td>
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<tr>
<td>Sep-Oct</td>
<td><strong>0.99 (0.71, 1.00)</strong></td>
<td><strong>0.85 (0.59, 0.97)</strong></td>
<td><strong>0.88 (0.62, 0.98)</strong></td>
<td>0.50 (0.33, 0.68)</td>
</tr>
<tr>
<td>May-Oct</td>
<td><strong>0.60 (0.22, 0.89)</strong></td>
<td>0.35 (0.11, 0.70)</td>
<td>0.15 (0.03, 0.42)</td>
<td>0.18 (0.05, 0.44)</td>
</tr>
<tr>
<td>2008</td>
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<tr>
<td>June-Aug.</td>
<td><strong>0.98 (0.80, 1.00)</strong></td>
<td>0.93 (0.75, 0.99)</td>
<td>0.88 (0.65, 0.97)</td>
<td>0.85 (0.63, 0.96)</td>
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<tr>
<td>Aug.-Oct.</td>
<td>0.67 (0.48, 0.82)</td>
<td>0.61 (0.47, 0.76)</td>
<td>0.57 (0.42, 0.70)</td>
<td>0.58 (0.46, 0.70)</td>
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<tr>
<td>June-Oct.</td>
<td><strong>0.80 (0.61, 0.92)</strong></td>
<td>0.51 (0.37, 0.65)</td>
<td>0.51 (0.38, 0.65)</td>
<td>0.48 (0.39, 0.58)</td>
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<tr>
<td><strong>Growth (cm²)</strong></td>
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<td>2007</td>
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<tr>
<td>May-July</td>
<td>2.22 (0.97, 3.49)</td>
<td>2.20 (1.14, 3.28)</td>
<td>2.32 (1.30, 3.39)</td>
<td>2.12 (1.00, 3.26)</td>
</tr>
<tr>
<td>July-Sep</td>
<td>0.34 (-0.79, 1.46)</td>
<td>0.21 (-0.58, 0.99)</td>
<td>0.69 (-0.27, 1.70)</td>
<td>0.59 (-0.51, 1.61)</td>
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<tr>
<td>Sep-Oct</td>
<td><strong>1.29 (-0.48, 3.14)</strong></td>
<td><strong>0.37 (-0.66, 1.43)</strong></td>
<td><strong>0.50 (-0.65, 1.61)</strong></td>
<td>-1.45 (-2.29, -0.62)</td>
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<td>2008</td>
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<tr>
<td>June-Aug.</td>
<td>1.05 (0.15, 1.77)</td>
<td>1.38 (0.72, 1.97)</td>
<td>1.53 (1.18, 1.95)</td>
<td>1.49 (1.27, 1.74)</td>
</tr>
<tr>
<td>Aug.-Oct.</td>
<td>-0.28 (-0.97, 0.66)</td>
<td>-0.09 (-0.72, 0.43)</td>
<td>0.55 (-0.05, 1.27)</td>
<td>0.21 (-0.38, 0.60)</td>
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<tr>
<td><strong>Final plant size (cm²)</strong></td>
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<tr>
<td>2007</td>
<td><strong>6.07 (5.01, 7.25)</strong></td>
<td><strong>4.67 (3.81, 5.58)</strong></td>
<td>3.06 (2.26, 3.99)</td>
<td>3.66 (2.96, 4.42)</td>
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<tr>
<td>2008</td>
<td>5.27 (4.24, 6.56)</td>
<td>6.14 (5.17, 7.41)</td>
<td>5.47 (4.60, 6.73)</td>
<td>6.22 (5.47, 7.22)</td>
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<tr>
<td><strong>Fecundity (No. of flowers per plant)</strong></td>
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<td>2007</td>
<td><strong>20.10 (8.04, 44.48)</strong></td>
<td><strong>18.03 (7.25, 40.33)</strong></td>
<td><strong>6.51 (2.56, 14.47)</strong></td>
<td>13.46 (5.41, 30.04)</td>
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<tr>
<td>2008</td>
<td><strong>9.74 (4.22, 21.42)</strong></td>
<td>20.87 (9.18, 44.82)</td>
<td><strong>24.15 (10.52, 51.75)</strong></td>
<td>20.92 (9.13, 44.79)</td>
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