Foraging modality and plasticity in foraging traits determine the strength of competitive interactions among carnivorous plants, spiders and toads

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Summary

1. Foraging modalities (e.g. passive, sit-and-wait, active) and traits are plastic in some species, but the extent to which this plasticity affects interspecific competition remains unclear.

2. Using a long-term laboratory mesocosm experiment, we quantified competition strength and the plasticity of foraging traits in a guild of generalist predators of arthropods with a range of foraging modalities.

3. Each mesocosm contained eight passively foraging pink sundews, and we employed an experimental design where treatments were the presence or absence of a sit-and-wait foraging spider and actively foraging toad crossed with five levels of prey abundance. We hypothesized that actively foraging toads would outcompete the other species at low prey abundance, but that spiders and sundews would exhibit plasticity in foraging traits to compensate for strong competition when prey were limited.

4. Results generally supported our hypotheses. Toads had a greater effect on sundews at low prey abundances, and toad presence caused spiders to locate webs higher above the ground. Additionally, the closer large spider webs were to the ground, the greater the trichome densities produced by sundews. Also, spider webs were larger with than without toads and as sundew numbers increased, and these effects were more prominent as resources became limited. Finally, spiders negatively affected toad growth only at low prey abundance.

5. These findings highlight the long-term importance of foraging modality and plasticity of foraging traits in determining the strength of competition within and across taxonomic kingdoms. Future research should assess whether plasticity in foraging traits helps to maintain coexistence within this guild and whether foraging modality can be used as a trait to reliably predict the strength of competitive interactions.

Key-words: Anaxyrus quercicus, Drosera capillaris, interspecific competition, kleptoparasitism, Sosippus floridanus

Introduction

Understanding how diversity in functional traits affects the distribution of species and ecosystem functions has been a major focus of ecological research in recent years (Bolker et al. 2003; Werner & Peacor 2003; McGill et al. 2006). For instance, predator functional diversity (in terms of foraging modality) has been shown to have cascading effects throughout food webs (Finke & Denno 2004; Lazzaro et al. 2009; Schmitz 2009; Steffan & Snyder 2010). However, few studies have empirically examined interspecific interactions among multiple predators with varied foraging modalities (Eklov & VanKooten 2001; Carey & Wahl 2010), even though determining the strength and outcome of these interactions between predators could have important implications for theory in community ecology (Wilson et al. 1999; Abrams 2010; Lee, Miller & Inouye 2011).

For example, theory suggests that predators with active foraging modalities should encounter prey at a higher rate than more passively foraging predators (e.g. sit-and-wait), and as a result, they might be superior competitors,
Consequently, we predicted that sundews would be negatively affected (compared with spiders and toads), but the addition of prey is known to increase seed production and vegetative propagation in some species (Thum 1988). Consequently, we predicted that sundews would be negatively affected by competitors because they cannot relocate to areas of high prey abundance. However, changes in prey and nutrient availability also have been shown to induce trait responses in other carnivorous plants, such as altering the number of trapping structures (Ellison & Gotelli 2002), amount of mucilage produced (Thoren et al. 2003) and size of leaves (Schulze & Schulze 1990). For this reason, we also predicted that sundews would decrease leaf area but increase trichome densities on leaves when prey abundance was low to increase their chances of catching prey. Spiders are capable of adjusting their web size and location in response to prey availability (Sherman 1994; Harwood, Sunderland & Symondson 2001, 2003), and thus, we predicted that spiders would shift their webs closer to the ground and increase web size as competition increases or prey abundance decreases. As the most active foragers and therefore potentially superior competitors, we predicted that toad growth would only be influenced by prey abundance and not the other two species. Additionally, consistent with theory (Werner & Anholt 1993), we predicted that toads would have the strongest adverse effect on competitors when prey were limiting.

**Materials and methods**

**Experimental design**

We conducted a mesocosm experiment for 1 year in the laboratory, enabling us to examine the long-term effects of competition. Mesocosms consisted of terraria (33 × 20 × 13 cm) filled to a depth of 2.5 cm with a mixture of 1/3 peat moss and 2/3 silica-based sand, with each terrarium containing eight equally spaced pink sundews (mean leaf rosette diameter ± SE = 3.10 ± 0.04 cm) transplanted from the field. We used a 2 × 2 × 5 design [spider presence/absence crossed with toad presence/absence crossed with 12, 16, 20, 24 or 28 commercially obtained pinhead (nymphs ~2 mm total length) house crickets (Acheta domestica) added twice per week], with four replicates of each treatment combination (a total of 80 terraria). All terraria were covered with window screening to prevent animals from escaping, and then Plexiglas to maintain humidity (Plexiglas alone did not provide a sufficient balance between airflow and containment of animals). Light fixtures above terraria were equipped with full spectrum bulbs, and light regimes were varied by season (9L:15D November–February, to 14L:10D March–October). Temperature remained at ~23 °C for the duration of the experiment. We added 100 ml of deionized water three times per week to ensure that moist conditions were maintained in the terraria. At the start of the experiment, terraria selected for toad and/or spider treatments received one individual of the appropriate species, and all animals were able to move throughout their particular terrarium. Thus, densities for each species in the mesocosms (sundews = 121 m⁻², toads = 15 m⁻², spiders = 15 m⁻²) were within or as close as possible logistically to densities found in the field (sundews max. = 148 m⁻², toads max. = 2 m⁻², spiders max. = 16 m⁻²; Greenberg & Tanner 2005; D. Jennings, unpublished data). We used immature oak toads (mean initial toad mass ± 1 SE = 0.79 ± 0.04 g) and spiders because of the long-term duration of the experiment, and any toads or spiders...
Data Collection

We collected data on several performance measures and traits of each species. Counts of living sundews (including any produced through vegetative propagation), number of leaves, leaf traits (trichome density and trapping area) and measurements of sundew diameter (cm) were conducted at the beginning and end of the experiment. To measure leaf traits, one randomly selected leaf from each sundew in a terrarium was removed and photographed. We then quantified the number of trichomes per leaf and trapping area (area of the leaf with trichomes) using ImageJ software (Schneider, Rasband & Elicieri 2012). Total trapping area was then estimated by multiplying the mean trapping area per leaf by the mean number of leaves per sundew per terrarium. We quantified spider web height (cm) and area (cm²) at the end of the experiment. Web area was calculated using ImageJ software (Schneider, Rasband & Elicieri 2012) after photographing webs. Toads were weighed (g) at the beginning and end of the experiment, and growth was calculated as a proportion of the initial mass of an individual. We also gathered baseline data on prey consumption. Terraria received crickets in a randomized order each week, and immediately after releasing crickets, we monitored any spider and/or toad for 2 min to record any predation events. Twenty-four hours after cricket releases, we counted the number of crickets captured on all sundew leaves.

Statistical Analyses

All statistical analyses were conducted using R 3.1.1 (R Core Team 2014). We examined the effects of competitors and prey abundance on each of the three species using generalized linear models with appropriate error distributions. To test for effects on toad growth, web area and web height, and differences in trichome density and total trapping area (from the beginning to the end of the experiment), we used a Gaussian error distribution, and for sundew counts, we used a Poisson error distribution (‘GLM’ function, ‘stats’ package). Significance was assessed with log-likelihood ratio tests and type II sums of squares. Initially, we tested for all biologically relevant two-, three- and four-way interactions in each model, but interactions with prey and/or toad for 2 min to record any predation events. We found that sundew numbers and sundew trapping area were negatively associated with web height, but the strength of this effect was dependent on web size (web area × web height interactions: number of sundews: $P = 0.033$, sundew trapping area: $P = 0.030$; Fig. 1b, Table 1, Appendix S1). When webs were small and thus only weakly competitive with sundews, sundew trapping area (and sundew numbers) was generally independent of web height (Fig. 1b, Appendix S1). However, when webs were large and thus posed a substantial competitive threat, the closer the webs were to the ground where the sundews grew, the less sundew trapping area (and total sundews) there was at the end of the experiment (Fig. 1b, Appendix S1).

Results

Effects on Sundews

Across all treatments, there was an increase in total sundew trapping area per terrarium from the beginning (mean ± 1 SE = 89.88 ± 9.18 cm²) to the end (mean ± 1 SE = 156.04 ± 16.87 cm²) of the experiment, and the number of sundews per terrarium at the end of the experiment (inclusive of original and newly germinated/propagated plants) ranged from 1 to 24 (mean ± 1 SE = 10.03 ± 0.58). Sundews captured nearly five times more prey ($F = 42.39, \text{d.f.} = 1, 78, P < 0.001$) when toads were absent (mean total number of crickets captured by sundews per terrarium = 23.15 ± 3.18) than present (4.80 ± 1.13). In fact, relative to the absence of a toad, the presence of a toad essentially negated the benefit of the food gradient for sundews and reduced the number of sundews and total trapping area per terrarium at the end of the experiment (toad × prey interactions: number of sundews: $P = 0.014$, trapping area: $P = 0.028$; Fig. 1a, Table 1, Appendix S1, Supporting information).

We also found that sundew numbers and sundew trapping area were negatively associated with web height, but the strength of this effect was dependent on web size (web area × web height interactions: number of sundews: $P = 0.033$, sundew trapping area: $P = 0.030$; Fig. 1b, Table 1, Appendix S1). When webs were small and thus only weakly competitive with sundews, sundew trapping area (and sundew numbers) was generally independent of web height (Fig. 1b, Appendix S1). However, when webs were large and thus posed a substantial competitive threat, the closer the webs were to the ground where the sundews grew, the less sundew trapping area (and total sundews) there was at the end of the experiment (Fig. 1b, Appendix S1).

Fig. 1. Effects of toads, spiders and prey abundance on the number of sundews. Grey circles/lines indicate absence of toads (a), and black circles/lines indicate presence of toads (a) or spiders (b).
Table 1. Results from generalized linear models testing effects of toad presence/absence, spider web area and height, and prey abundance, on sundew number (Poisson error distribution), sundew trichome density and sundew total trapping area (Gaussian error distributions). Interactions with $P > 0.1$ were dropped from final models (while respecting marginality). Shown are likelihood ratio $\chi^2$ values with $P$ values in parentheses, and all d.f. = 1

<table>
<thead>
<tr>
<th>Effect</th>
<th>Response</th>
<th>Sundew number</th>
<th>Trichome density</th>
<th>Total trapping area</th>
</tr>
</thead>
<tbody>
<tr>
<td>Toad (presence/absence)</td>
<td>16.18 ($&lt;0.001$)</td>
<td>0.19 (0.664)</td>
<td>0.24 (0.625)</td>
<td></td>
</tr>
<tr>
<td>Web area</td>
<td>0.53 (0.465)</td>
<td>2.03 (0.154)</td>
<td>4.46 (0.035)</td>
<td></td>
</tr>
<tr>
<td>Web height</td>
<td>&lt;0.01 (0.991)</td>
<td>0.07 (0.792)</td>
<td>0.19 (0.662)</td>
<td></td>
</tr>
<tr>
<td>Prey abundance</td>
<td>3.06 (0.080)</td>
<td>2.05 (0.152)</td>
<td>1.11 (0.292)</td>
<td></td>
</tr>
<tr>
<td>Toad $\times$ web height</td>
<td>2.78 (0.096)</td>
<td>0.21 (0.649)</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>Toad $\times$ prey</td>
<td>6.07 (0.014)</td>
<td>0.13 (0.717)</td>
<td>4.81 (0.028)</td>
<td></td>
</tr>
<tr>
<td>Web area $\times$ web height</td>
<td>4.57 (0.033)</td>
<td>–</td>
<td>4.71 (0.030)</td>
<td></td>
</tr>
<tr>
<td>Web height $\times$ prey</td>
<td>–</td>
<td>0.13 (0.715)</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>Toad $\times$ web height $\times$ prey</td>
<td>–</td>
<td>5.02 (0.025)</td>
<td>–</td>
<td></td>
</tr>
</tbody>
</table>

Bold type indicates statistically significant results ($P < 0.005$).

Fig. 2. Effects of toads, spiders and prey abundance on the difference in sundew trichome density (quantified at the beginning and end of the experiment). Three-way interaction plots show the interaction between web height and prey abundance (cross-sections represent 10th, 50th and 90th quantiles) on the difference in sundew trichome density, in the absence (a) and presence (b) of toads (toad $\times$ web height $\times$ prey: $P = 0.025$).

In addition to effects on sundew densities, there were also effects of the treatments on sundew traits. We measured leaf traits on a total of 1442 sundews and counted 115 338 trichomes. Mean trichome density was higher at the end (mean $\pm 1$ SE = 133.97 $\pm$ 7.99 trichomes cm$^{-2}$) than at the beginning (mean $\pm 1$ SE = 85.54 $\pm$ 1.87 trichomes cm$^{-2}$) of the experiment. As a reminder, our hypothesis was that trichome densities would be negatively associated with prey abundance and should increase with competition from spiders and toads. Our findings generally supported this hypothesis. When toads were absent (Fig. 2a), trichome density was more negatively associated with prey abundance when webs were close to the ground (i.e. high spider competition) than when webs were located higher above the ground. When toads were present (Fig. 2b), trichome density was generally higher than when they were absent, but trichome densities were negatively associated with prey abundance at all web heights (toad $\times$ web height $\times$ prey: $P = 0.025$, Table 1).

**EFFECTS ON SPIDERS**

Spiders appeared to adjust the height and size of their webs in response to competitors and prey abundance. Webs were located more than two times higher above the ground when toads were present (mean $\pm 1$ SE = 6.00 $\pm$ 0.61 cm) than when they were absent (mean $\pm 1$ SE = 2.70 $\pm$ 0.60 cm). Spiders also captured significantly more prey ($F = 7.13$, d.f. = 1, 38, $P = 0.011$) when toads were absent (mean total crickets captured per spider $\pm 1$ SE = 21.70 $\pm$ 3.29) than when toads were present (11.60 $\pm$ 1.70). Further, when toads were present, web height was positively associated with prey abundance, but without toads there was a negative relationship between web height and prey (toad $\times$ prey: $P = 0.045$; Fig. 3, Table 2).

There was a significant three-way interaction between sundew trapping area, toad presence and prey abundance on spider web area (total trapping area $\times$ toad $\times$ prey:...
Fig. 3. Effect of toads on spider web height. Box plot shows median (horizontal line within box) and interquartile range (box), and whiskers extend to the lower and upper outermost data points.

Table 2. Results from generalized linear models testing effects of sundew total trapping area, toad presence/absence, and prey abundance on spider web height, and spider web area (Gaussian error distributions). Interactions with \( P > 0.1 \) were dropped from final models (while respecting marginality). Shown are likelihood ratio \( \chi^2 \) values with \( P \) values in parentheses, and all d.f. = 1

<table>
<thead>
<tr>
<th>Effect</th>
<th>Response</th>
<th>Web height</th>
<th>Web area</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total sundew trapping area</td>
<td></td>
<td>0.74 (0.390)</td>
<td>4.92 (0.027)</td>
</tr>
<tr>
<td>Toad (presence/absence)</td>
<td></td>
<td>16.71 (&lt;0.001)</td>
<td>4.59 (0.032)</td>
</tr>
<tr>
<td>Prey abundance</td>
<td></td>
<td>0.21 (0.647)</td>
<td>24.25 (&lt;0.001)</td>
</tr>
<tr>
<td>Trapping area ( \times ) toad</td>
<td></td>
<td>–</td>
<td>2.86 (0.091)</td>
</tr>
<tr>
<td>Trapping area ( \times ) prey</td>
<td></td>
<td>–</td>
<td>1.45 (0.229)</td>
</tr>
<tr>
<td>Toad ( \times ) prey</td>
<td></td>
<td>–</td>
<td>0.30 (0.587)</td>
</tr>
<tr>
<td>Trapping area ( \times ) toad ( \times ) prey</td>
<td></td>
<td>–</td>
<td>5.16 (0.023)</td>
</tr>
</tbody>
</table>

Bold type indicates statistically significant results (\( P < 0.005 \)).

\( P = 0.023; \) Fig. 2b,c, Table 2). When total sundew trapping area was low, web area exhibited only a weak negative association with prey abundance. In contrast, when total sundew trapping area was high, web area was strongly negatively associated with prey abundance. In both cases, the relationship between web area, trapping area and prey was stronger when toads were present (Fig. 4a) than when they were absent (Fig. 4b).

**EFFECTS ON TOADS**

Toad growth was negatively associated with web area (web area: \( P = 0.003; \) Fig. 5a, Table 3) and positively associated with prey abundance (prey: \( P < 0.001; \) Fig. 5b, Table 3), although the mean total number of crickets we observed being captured per toad was similar (\( F = 0.48, \) d.f. = 1, 38, \( P = 0.494 \)) whether spiders were present (mean \( \pm 1 \ SE \) = 110.10 \( \pm 7.10 \)) or absent (mean \( \pm 1 \ SE \) = 116.50 \( \pm 6.48 \)). We did not detect any significant effect of sundew densities or trapping area on toad growth (Table 3). While relatively rare overall, on 97 occasions, we observed toads consuming live crickets that had become trapped by sundews and were struggling in mucilage. Assuming that these crickets would otherwise have been consumed by sundews, this would have amounted to 8% of the total number of crickets captured by the plants.

**Discussion**

Results from our long-term experiment demonstrate that foraging modality and plasticity in foraging traits can be important determinants of the strength and outcome of competitive interactions (Fig. 6). Sundew numbers, total trapping area and trichome density were all affected by competitors and prey abundance. For sundew numbers, toads had a negative effect, but the effect of spiders was mediated by the interaction between the height and size of their webs (although with generally more sundews as webs increased in size). Small webs did not appear to have much of a competitive effect on sundews regardless of where they were, but large webs closer to the ground where the sundews reside were more detrimental to sundews than large webs higher above the ground. Interspecific competition has been implicated as a potential cause of changes in total trapping area of sundews (Jennings et al. 2010), and thus, it was not surprising to observe other leaf trait changes, such as increases in trichome density, associated with competition. In other plants, trichome production is also plastic. For instance, when trichomes are thought to serve a defensive function, herbivory can induce an increase in their densities (Holeski 2007; Bjorkman, Dalin & Ahnre 2008).

Spiders generally increased web area in response to reduced availability of prey caused by competition with toads or sundews, a behaviour consistent with findings from other studies (Sherman 1994). Additionally, toads caused \( S.\) *floridanus* to elevate the height of their webs, consistent with this species locating webs anywhere from on the ground to above the ground among shrubs and cacti in the field (Punzo & Haines 2006). Previous work has indicated that spiders relocate or modify their webs to avoid competition or possibly predation (Harwood & Obrzycki 2005; Manicom et al. 2008; Jennings et al. 2010), and web relocation is energetically costly for spiders that produce complex webs, such as sheet and funnel webs (Tanaka 1989). In this system, we believe that competition (exploitation or interference) rather than intraguild predation (i.e. toads, spiders and sundews competing and depredating one another) was the driver of web location behaviour. Spiders might choose to locate webs higher above the ground to avoid interference competition with toads and to prevent these energetically costly webs from being damaged by their competitors. Although it is possible that under field conditions intraguild predation between these three species occurs, we did not observe it...
in the mesocosms and believe it to be relatively rare for several reasons. For example, field surveys indicate that spiders of any species are rarely captured by pink sundews, and those individuals that are caught are generally very small in size (Jennings et al. 2010). Additionally, S. floridanus spiders display maternal care, with spiderlings staying with their mother in the web for up to 5 months (Brach 1976; Punzo & Haines 2006). By the time these spiders leave their maternal webs, they are generally >1 cm in length (Punzo & Haines 2006) and likely too large to be caught in most sundews. At this size, they might also deter small oak toads from considering them as prey.

Oak toads were the strongest competitors, with just one individual influencing both sundews and spiders. As predicted, prey abundance was the main factor affecting oak toad growth; although at low prey abundance, spiders also caused a significant negative effect on toads. The competitive superiority of oak toads could partly be explained by our observations of them stealing crickets caught in sundew leaves, and this behaviour might partially explain why, contrary to our hypothesis, they still exerted a strong negative effect on sundew numbers even at high prey abundance. Kleptoparasitic behaviour has been seen in a number of carnivorous plant–animal interactions (Zamora 1990; Zamora & Gomez 1996; Anderson & Midgley 2002). This kleptoparasitism between toads and sundews provides another example of the often surprising and complex interactions between carnivorous plants and animals (Clarke et al. 2009; Moon et al. 2010; Grafe et al. 2011). Although we showed that the actively foraging oak toads were superior competitors to the more

Table 3. Results from generalized linear model testing effects of spider web area, sundew total trapping area and prey abundance on toad growth (Gaussian error distribution). Interactions with \( P > 0.1 \) were dropped from final models (while respecting marginality). Shown are likelihood ratio \( \chi^2 \) values with \( P \) values in parentheses, and all d.f. = 1

<table>
<thead>
<tr>
<th>Effect</th>
<th>Response</th>
<th>( \chi^2 )</th>
<th>( P )</th>
</tr>
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<tbody>
<tr>
<td>Web area</td>
<td>9.00 (0.003)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Web height</td>
<td>0.82 (0.365)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total sundew trapping area</td>
<td>3.66 (0.056)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prey abundance</td>
<td>31.96 (&lt;0.001)</td>
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Bold type indicates statistically significant results \( (P < 0.005) \).
passively foraging spiders and sundews, active foragers expend more energy in pursuit of food and their activity could increase their risk of predation. Thus, in the field, where space and predation pressures are greater than in our mesocosms, interaction strengths among oak toads, spiders and sundews could be different.

We endeavoured to recreate a natural system using the mesocosms, but obviously there are some limitations with this type of experimental design. For example, the densities of oak toads used were higher than any that have been documented in the field (although to our knowledge, very few data exist on the densities of this particular species in nature), and this could have been responsible in part for the strong effects of toads that were observed. Additionally, although crickets comprise part of the diets of all three species (Punzo 1995; Punzo & Haines 2006; Jennings et al. 2010), clearly in natural habitats there is a much more diverse prey assemblage which could help to mitigate the intensity of competition. Consequently, any future work in this system using mesocosms should increase prey diversity to determine whether the outcomes of competition remain the same. Lastly, it is possible that certain soil characteristics (e.g. depth, composition and drainage) could have affected the condition of the sundews in some cases. For instance, given the duration of the experiment, it is likely that toad defecation in the mesocosms increased the variability of nitrogen and phosphorous in the soil and provided additional nutrients for some sundews that could partly offset any resources lost from not catching prey directly. These limitations are all important considerations when interpreting our findings, and improvements in the design of the mesocosms could be made for future work, but the trends in the results nonetheless appear to be robust.

In conclusion, our results further highlight the importance of considering functional diversity and plasticity in traits when examining species interactions and other community- and ecosystem-level patterns and processes. Predator functional diversity has previously been shown to have effects on trophic interactions that propagate through food webs (Schmitz 2007, 2009; Lazzaro et al. 2009; Rohr et al. 2015), and it seems plausible that the results we observed could have similar effects. For example, if spiders relocate their webs higher above the ground in response to exploitative or interference competition with oak toads, then the prey assemblage caught in their webs would likely be considerably different than webs on the ground (Brown 1981; Knop, Zund & Sanders 2014). Capturing a greater diversity of arthropod prey (including many herbivorous insects) could have broader community-level effects, such as facilitating an increase in plant diversity, and plasticity in foraging traits could help to mediate coexistence among this guild (Kobler et al. 2009). However, further studies in this system are needed to test these hypotheses and improve our understanding of the population dynamics for species in these increasingly threatened taxa (Skerratt et al. 2007; Rohr et al. 2008; Jennings & Rohr 2011; Jennings, Congelosi & Rohr 2012; Rohr & Palmer 2013).

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Data accessibility
Data available from the Dryad Digital Repository: http://dx.doi.org/10.5061/dryad.m23g6 (Jennings, Krupa & Rohr 2016).

References


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Supporting Information

Additional Supporting Information may be found in the online version of this article.

Appendix S1 (Fig. S1). Effects of toads, spiders, and prey abundance, on sundew total trapping area. Grey circles/lines indicate absence of toads (a), and black circles/lines indicate presence of toads (a) or spiders (b).