Field spatial scale and predator colonization behavior mediates pest suppression in diversified agroecosystems

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ABSTRACT

Ecological field studies and theory have demonstrated over the past several decades that the spatial scale at which heterogeneous habitats are deployed can have a profound effect on the growth, movement, reproduction, and mortality of resident plants and animals. Increasing evidence, however, suggests that landscape complexity affects different organisms in different ways. Very little is known about the underlying mechanisms driving such differences in species responses, creating challenges for determining how best to manage landscapes in order to maximize environmental services such as biological control. Diversifying agroecosystems by establishing or retaining natural vegetation in and around crop areas has long been recognized as a potentially effective means of bolstering pest control by attracting more numerous and diverse natural enemies, though outcomes are inconsistent. We address here gaps in our understanding of the link between non-crop vegetation in field margins and pest suppression by using a system of partial differential equations to model population-level predator-prey interactions as well as spatial processes, in order to capture the dynamics of crop plants, herbivores, and two generalist predators. We focus on differences in how these two predators (a carabid and a ladybird beetle) colonize crop fields where they forage for prey, examining differences in how they move into the fields from adjacent vegetation as a potential driver of differences in overall pest suppression. Our results demonstrate that predator colonization behavior and spatial scale are important factors in determining pest suppression.
INTRODUCTION

The role that landscape spatial scale and heterogeneity plays in species’ interactions and distributions is a central theme of modern ecology (Levin, 1992; Wiens, 1989; Andow, 1991; Englund and Hambäck, 2004; Caballero-López et al., 2012). Theoretical and field studies over the past several decades have demonstrated that the spatial scale at which heterogeneous habitats are deployed has a profound effect on growth, movement, reproduction, and mortality of resident plants and animals (Marino and Landis, 1996; Jonsen and Fahrig, 1997; Roland and Taylor, 1997; Turchin, 1998; Banks, 1998; Bommarco and Banks, 2003; Thies et al., 2003; Holland et al. 2004; Hambäck and Englund, 2005). Recent studies in agroecosystems, however, have highlighted the fact that landscape complexity affects different organisms in different ways, prohibiting facile generalizations about the effects of habitat scale and complexity (Caballero-López et al., 2012; Chaplin-Kramer et al., 2011, 2012). Even less is known about the underlying mechanisms driving differences in species responses, creating challenges for determining how best to manage landscapes in order to maximize environmental services such as biological control/pest suppression.

Diversifying agroecosystems by establishing or retaining natural vegetation in and around crop areas has long been recognized as a potentially effective means of bolstering pest control (Landis et al., 2000; Östman et al., 2001; MacLeod et al. 2004; Heimpel and Jervis, 2005; Gardiner et al., 2009; Chaplin-Kramer and Kremen, 2012; Schellhorn et al., 2014, Rusch et al. 2006). The natural enemies hypothesis developed by Root (1973) posits that more diversified agroecosystems tend to attract and harbor larger and more diverse numbers of predators, thereby increasing predation pressure and lowering pest abundance. The addition of non-crop vegetation in and around crop fields often translates to increased resources such as nectar and pollen, which are critical components of many predator/parasitoid diets (Banks et al., 2008; Lee et al., 2006). Precisely how weedy/natural vegetation is incorporated into agricultural fields may greatly influence the extent and reliability of pest suppression by natural enemies. One strategy, which is common on smaller organic farms, is to incorporate weeds in intercropped rows within a field. This type of habitat manipulation effectively fragments crop areas, which can disrupt biological control in some cases (Kareiva, 1987; Prasad and Snyder, 2006) and enhance it in others (Risch et al., 1983; Andow, 1991; Tonhasca and Byrne, 1994). Differences in these outcomes often depend on the spatial scale of crop fields, and also on the movement of herbivores and natural enemies (Banks 1998, 1999; Turchin 1998; Banks and Yasenak 2003, Bommarco and Banks 2003, Thies et al., 2003; Grez et al. 2008). Another common strategy for incorporating natural vegetation into agroecosystems, which is more amenable to use in larger-scale agriculture, is to retain weeds, woody plants, and other non-crop vegetation adjacent to or in the margins of crop production areas (Banks 2000, Banks and Stark 2004, Bianchi et al, 2006; Gardiner et al., 2009).
The adjacent vegetation may act as a source for predators and parasitoids; though increased pest suppression is not always guaranteed (Jonsson et al. 2008; Rusch et al., 2013, Bianchi et al., 2017; Perez-Alvarez et al., 2018). This highlights the difficulty in establishing a general theory relating non-crop vegetation adjacent to crop fields to pest control. We attempt here to address gaps in our understanding of the link between non-crop vegetation in field margins and pest suppression by using a simple mathematical model that captures the dynamics of crop plants, herbivores, and two generalist predators. We focus on differences in how these two predators (a carabid and a ladybird beetle) colonize crop fields where they forage for prey, examining differences in how they move into the fields from adjacent vegetation as a potential driver of differences in overall pest suppression. Furthermore, we examine how differences in colonization behavior may interact with spatial scale in determining the ability of predators to suppress prey in diversified agroecosystems.

METHODS:

Simulation/scenario

We modeled the dynamics of a pest-herbivore-predator agroecosystem, using as an example aphids as the principal herbivore feeding on an annual crop, subject to predation by a ladybird beetle (Coleoptera: Coccinellidae) or a carabid beetle (Coleoptera: Carabidae). This scenario, in which a principle herbivore in an annual cropping system is attacked by generalist predators, is quite common throughout temperate regions, and has been the setting for numerous field studies exploring habitat heterogeneity and predator-prey interactions (Root 1972, Risch et al. 1983, Kareiva 1987, Bommarco and Ekbom 1996, Banks 1998, 1999; Elliott et al. 2002, Grez and Prado 2000, Bommarco et al. 2007, Grez et al. 2008, Caballero-López et al.2012). We used a series of coupled differential equations to describe plant population growth, aphid consumption and population growth, and carabid or ladybird beetle colonization and migration from adjacent non-crop vegetation.

Carabid beetles (Coleoptera: Carabidae) are important predators commonly found in agroecosystems (Lovei and Sunderland, 1996). Primarily ground-dwelling, carabids typically overwinter in annual cropping systems (especially in no-till schemes) or nearby vegetation, moving into crop fields as prey densities increase during the springtime. The adults prey on aphids and other soft-bodied insects throughout the growing season, consuming aphids that fall to the ground or occasionally climbing up on plants to chase down prey.

Ladybird beetles are also common and important agricultural predators; overall they are classified as generalist predators, though many species restrict their diets to one or few aphid species (Hodek et al., 2012). They typically overwinter in non-crop vegetation that may be adjacent or farther away from the annual crop fields, flying in to colonize crop fields early in the growing season when prey densities begin to increase. They feed regularly on nectar, so may
take advantage of nearby non-crop vegetation to bolster their populations during the growing season.

Model and parameterization

We modeled the dynamics of tri-trophic interactions in a simulated agroecosystem over the course of a 90-day growing season. Aphids are the principal herbivore; they feed on crop plants and are subject to predation by one of two predators: (i) a ladybird beetle, or (ii) a carabid beetle. At the start of the simulation, there is an initial population of crop plants, aphids, and either species of predators in the crop field. Crop plants grow throughout the season as aphids feed on them, and additional predators enter the field by walking and flying dispersal behavior.

We describe these dynamics with a system of partial differential equations, which include population-level predator-prey interactions as well as spatial processes. For simplicity’s sake, we assume that carabid beetles migrate into the field by dispersing via the ground from nearby/adjacent vegetation (e.g., weedy margins). In contrast, we assume that ladybird beetles colonize the crop field via the air column, flying in from adjacent/nearby vegetation. We highlight these differences in colonization behavior by varying values of predator population colonization in the model. We then solve for aphid dynamics over 90 days in square fields of varying sizes. In keeping with the mean farm size in the United States in 2017 (approximately 180 ha), we report on results of simulations for farms sizes ranging from 1 to 200 hectares (USDA, 2018), in order to better understand how the way in which the predators colonize crop fields interacts with spatial scale in the suppression of the aphid pest population.

We denote the populations of the crop, aphid, and predator populations by $A$, $B$, and $C$ respectively in the following system of partial differential equations:

$$\frac{dA}{dt} = r_A \left(1 - \frac{A}{K_A}\right)A - \mu_A AB$$

$$\frac{dB}{dt} = r_B AB + \nabla \cdot \left[(d_B + d_{BC} C) \nabla B\right] - \mu_B BC$$

$$\frac{dC}{dt} = \nabla \cdot d_C \nabla C + \nabla \cdot (VC) + M_F$$

$$\frac{dV}{dt} = d_{vB} \nabla B + d_v \Delta V$$

Here the crop grows logistically at rate $r_A$ until reaching the maximum value $K_A$ and is consumed by aphids at rate $\mu_A$. Over a single growing season, we assume that the crop has negligible spatial dynamics. Aphids grow in the presence of crop resources at rate $r_B$ and are consumed by predators at rate $\mu_B$. We model the movement of aphids within the field due to the production of winged alates (Dixon, 1977) by simple diffusion at rate $d_B$, and additionally
assume that predator-induced disturbance increases this diffusion at rate $d_{BC}$. (Weisser 1999). Carabid predators migrate in solely through the field edges at rate $M_E$, and ladybirds migrate directly into the field interior via the air column at rate $M_F$ (this latter term is set to zero for carabids; see Table 3). Once in the field, both predator species diffuse at rate $d_C$ while hunting within the field, corresponding to a random walk (Holmes et al., 1994; Turchin, 1998). We also assume that in close proximity to aphids both predator species exhibit “prey-taxis” (Kareiva and Odell, 1987) and experience directed movement towards prey items with the same velocity, denoted by $V$. We follow (Arditi et al., 2001) in the formulation of this velocity, assuming movement towards aphid resources at rate $d_{VB}$ and smoothing of the movement due to intraspecific competition at rate $d_V$.

At the start of each simulation, crops are uniformly distributed in the field at density $A_0$. Aphids are randomly distributed in the field, but normalized to a constant average density $B_0$. The preliminary predator populations are also randomly distributed in the field at an average density $C_0$, assuming some form of no-till farming. The velocity due to directed motion is initially zero across the field. Aphid populations are unable to pass through the boundary of the spatial domain (the edge of the agricultural field) for the duration of the simulation (we assume no loss of aphids via dispersal out of the system). Velocity is also limited at the edge of the field, as described in (Arditi et al., 2001). We summarize these rates and dynamics in Table 1.

<table>
<thead>
<tr>
<th>Definition</th>
<th>Interior Dynamics</th>
<th>Edge Dynamics</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A$</td>
<td>Crop maturity</td>
<td>N/A - Crops do not move</td>
</tr>
<tr>
<td></td>
<td>Grows logistically, consumed by aphids</td>
<td></td>
</tr>
<tr>
<td>$B$</td>
<td>Aphid population density</td>
<td>No migration in or out</td>
</tr>
<tr>
<td></td>
<td>Grows when consuming crops, consumed by predators,</td>
<td></td>
</tr>
<tr>
<td></td>
<td>spreads due to aerial migration, migration increases</td>
<td></td>
</tr>
<tr>
<td></td>
<td>with predators</td>
<td></td>
</tr>
<tr>
<td>$C$</td>
<td>Predator population density</td>
<td>(Carabids) migration (walking) into field;</td>
</tr>
<tr>
<td></td>
<td>Both species move via random walks, “prey-taxis” (</td>
<td>(Ladybirds) migration (flying) into field</td>
</tr>
<tr>
<td></td>
<td>directed motion at velocity $V$)</td>
<td></td>
</tr>
<tr>
<td>$V$</td>
<td>Velocity of directed motion</td>
<td>No velocity along edges,</td>
</tr>
<tr>
<td></td>
<td>Increases with aphid density, smooths due to</td>
<td>no increase or decrease out edges</td>
</tr>
<tr>
<td></td>
<td>competition</td>
<td></td>
</tr>
</tbody>
</table>

Table 1: Summary of model dynamics in the field and at field edges.
In our simulations, we varied only parameters related to predator migration to the field before or during the season. We fixed other model parameters to values from the literature when possible; the remaining values were selected to produce reasonable model solutions. In Table 2, we list the parameters and values, along with sources for the values where appropriate. We note that most values have been converted to the scale of our simulations.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A_0$</td>
<td>0.2</td>
<td>Fixed</td>
</tr>
<tr>
<td>$B_0$</td>
<td>1</td>
<td>Fixed</td>
</tr>
<tr>
<td>$r_A$</td>
<td>$6.42 \times 10^{-2}$</td>
<td>Calculated (based on $A_0$) such that crops reach 95% maturity in 90 days</td>
</tr>
<tr>
<td>$K_A$</td>
<td>1</td>
<td>Fixed (100% maturity)</td>
</tr>
<tr>
<td>$\mu_A$</td>
<td>$9.37 \times 10^{-7}$</td>
<td>(Larrson 2005, Wang 1994), conversion: (Roubinet 2015)</td>
</tr>
<tr>
<td>$r_B$</td>
<td>0.3</td>
<td>(Jonsson et al., 2018; Laubmeier et al. 2018)</td>
</tr>
<tr>
<td>$d_B$</td>
<td>$8.87 \times 10^{-6}$</td>
<td>(Bommarco et al. 2007)</td>
</tr>
<tr>
<td>$d_{BC}$</td>
<td>$6.81 \times 10^{-6}$</td>
<td>(Weisser 1999)</td>
</tr>
<tr>
<td>$\mu_B$</td>
<td>1</td>
<td>Selected such that aphid populations are comparable to results in (Curtsdotter 2018)</td>
</tr>
<tr>
<td>$d_C$</td>
<td>$3.00 \times 10^{-3}$</td>
<td>(Allema 2014)</td>
</tr>
<tr>
<td>$d_{VB}$</td>
<td>$1.00 \times 10^{-7}$</td>
<td>Selected such that predator speeds are comparable to results in (Wallin 1994)</td>
</tr>
<tr>
<td>$d_V$</td>
<td>$1.00 \times 10^{-4}$</td>
<td>Scaled to $d_{VB}$</td>
</tr>
</tbody>
</table>

Table 2: Parameters fixed in our simulations, with sources for values fixed from literature.

We established baseline parameter values for predator migration such that population densities were comparable to those observed in Thomas (2006); these values are given in Table 3. We additionally considered the effect of lower or higher initial and subsequent rates of migration,
with values of $C_0$ and $M_F$ (for carabids) falling between 0.80 – 0.15. The corresponding values of $M_F$ (for ladybirds) fall between 0.00097 – 0.018. We used the method of lines (Scheisser 2012) to numerically evaluate the PDE for our simulations, in MATLAB version 8.3.0.532 (R2014a).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>$C_0$</th>
<th>$M_E$</th>
<th>$M_F$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>carabids: 0.1 0.1 0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ladybirds:</td>
<td>0.1 0 0.0012</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3: “Baseline” parameter values for ladybird and carabid simulations. The rate $M_F$ was calculated to match daily field-level population growth due to field edge migration $M_E$.

RESULTS:

Predator migration/colonization rates had a marked effect on aphid suppression, with the lowest levels of aphids corresponding to simulations with the highest carabid and ladybird migration levels (Figs. 1-3). Spatial scale was an important mediating factor for aphid suppression for carabids, but not for ladybird beetles; for carabid simulations, aphid densities were substantially higher at larger spatial scales than at lower scales (Figs. 1a, 2a, 3a), whereas aphid densities were invariant across scales for ladybird beetle simulations (Figs. 1b, 2b, 3b). At 90 days, aphids in ladybird simulations were reduced on average by 91-92% of their initial population across all spatial scales. The effects of carabid predation were heavily influenced by spatial scale, with aphid reductions ranging from 88% (1ha) to 3% (200ha) for the same time scale. The relationship between carabid densities and field spatial scale was non-linear, with proportional predation effects diminishing at larger scales (Fig. 4).

Overall, ladybird beetles flying in to colonize crop plants markedly reduced aphids more than carabids walking in from weedy margins, for all spatial scales and migration rates (Figs. 5-11). Differences between carabid and ladybird pest suppression was the highest at high flux (migration) levels, at the largest spatial scales (e.g., Fig. 11a vs. 11c). Finally, there were differences in aphid distributions depending on which predator species was present. Aphids subject to carabid predation were more centrally distributed than aphids subject to ladybird predation, throughout the simulations (Figures 12-15).
DISCUSSION:

Pest suppression in our simulations varied widely as a function of both predator colonization behavior and agroecosystem spatial scale. In all cases and across all spatial scales, ladybird beetles more effectively controlled aphids than carabid beetles (Figs. 1-11). Aphid densities subject to carabid predation demonstrated a non-linear relationship between pest suppression and spatial scale. At baseline migration levels, aphid densities were 6 times higher in the 5ha field simulations than in the 1ha simulations, 35 times higher in the 50ha field simulations than in the 1ha simulations, and 43 times higher in the 100ha field simulations than in the 1ha simulations (Fig. 4). This attenuation of spatial scale effects was even more evident at larger spatial scales, with a similar ten-fold increase in spatial scale (from 10ha to 100ha) yielding only 3.3-fold difference in aphid densities.

The spatial details of predator-prey distributions also reveal striking differences in pest suppression between the two predators. At both small and large scales carabids were able to corral aphids into the center of the field by the end of 90 days (e.g., Figs.12,14), whereas ladybird predation resulted in aphids being more uniformly distributed across the field at the end of the simulation (Figs. 13, 15). However, aphid densities were much higher in simulations with carabid predation at the larger scales (Figs. 1a,2a,3a); despite the fact that carabids were able to concentrate aphids in the center of the field, aphid reproduction was able to overwhelm them by the end of the simulations. Our results highlight the importance of considering both spatial scale as well as predator movement behavior in assessing the effectiveness of habitat manipulations for pest suppression in agroecosystems.

Previous studies exploring predator behavior in response to habitat heterogeneity and spatial scale have focused primarily on the effects of supplemental (e.g., floral) resources on parasitoid physiology and behavior (Heimpel and Jervis, 2005; Lee et al., 2006). Other studies have explored the effects of patch size on predator/parasitoid colonization, exploring how differences in within-field foraging behavior (e.g., directed vs. diffusive) can interact with habitat scale and patterning (Karevia, 1985; Olson et al., 2000; Arditi et al., 2001; Banks and Yasenak, 2003). Bianchi et al. (2017) highlighted the importance of landscape spatial scale, using a mass-action action perspective to illustrate the importance of habitat management for natural enemies while being mindful of predator dispersal abilities. Hambäck and Englund (2005) urged careful attention to scaling migration rates to patch size when attempting to account for the wide range of correlations (both positive and negative) between patch size and herbivore colonization; . We suggest a similar approach might be useful in the context of predator colonization in diverse agroecosystems. In the current study, in addition to spatial-scale effects, we detected differences in aphid suppression stemming solely from variation in colonization type, keeping all other factors – including migration rates and local within-field movement behavior – constant. Taken together, these results have potentially important implications for habitat management aimed at...
pest suppression. In particular, careful consideration of the colonization modality of important predators in any agroecosystem along with the habitat spatial scale is critical when planning or implementing landscape habitat manipulations for pest control.

Species interactions and community dynamics likely play an important additional role in determining the effectiveness of pest suppression in diversified agroecosystems. Predator-predator interactions, along with multi-trophic interactions (including intra-guild predation, hyperparasitism, etc.) have been shown to alter population dynamics and interfere with prey suppression (Snyder and Ives, 2001, 2003; Rand et al., 2012). Furthermore, landscape complexity in agroecosystems may simultaneously increase natural enemy pressure while bolstering pest populations (Perez-Alvarez et al., 2018). Understanding the intersection of predator movement behavior and habitat complexity at the landscape scale in the context of species assemblages and interactions is therefore critical to prescribing useful habitat manipulations for biological control in agroecosystems.

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Figure 1: Final aphid population density for the duration of the season at baseline migration levels for (a) carabid beetle and (b) ladybird beetle predation. Populations are plotted at log scale.
Figure 2: Final aphid population density for the duration of the season at low migration levels for (a) carabid beetle and (b) ladybird beetle predation. Populations are plotted at log scale.

Figure 3: Final aphid population density for the duration of the season at high migration levels for (a) carabid beetle and (b) ladybird beetle predation. Populations are plotted at log scale.
Figure 4: Relationship between final aphid densities and field spatial scale.

Figure 5: Aphid population densities over the duration of the season at (a) low, (b) baseline, and (c) high migration levels in a 1 hectare field. Populations are plotted on a log scale.
Figure 6: Aphid population densities over the duration of the season at (a) low, (b) baseline, and (c) high migration levels in a 3 hectare field. Populations are plotted on a log scale.

Figure 7: Aphid population densities over the duration of the season at (a) low, (b) baseline, and (c) high migration levels in a 5 hectare field. Populations are plotted on a log scale.
Figure 8: Aphid population densities over the duration of the season (a) low, (b) baseline, and (c) high migration levels in a 10 hectare field. Populations are plotted on a log scale.

Figure 9: Aphid population densities over the duration of the season at (a) low, (b) baseline, and (c) high migration levels in a 50 hectare field. Populations are plotted on a log scale.
Figure 10: Aphid population densities over the duration of the season at (a) low, (b) baseline, and (c) high migration levels in a 100 hectare field. Populations are plotted on a log scale.

Figure 11: Aphid population densities over the duration of the season at (a) low, (b) baseline, and (c) high migration levels in a 200 hectare field. Populations are plotted on a log scale.
Figure 12: Heatmaps of the aphid population density over a 1 hectare field in the presence of a carabid predator population on days (a) 0, (b) 30, (c) 60, and (d) 90 of the season.
Figure 13: Heatmaps of the aphid population density over a 1 hectare field in the presence of a ladybird predator population on days 0, 30, 60, and 90 of the season.
Figure 14: Heatmaps of the aphid population density over a 50 hectare field in the presence of a carabid predator population on days 0, 30, 60, and 90 of the season.
Figure 15: Heatmaps of the aphid population density over a 50 hectare field in the presence of a ladybird predator population on days 0, 30, 60, and 90 of the season.

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