Urbanization alters the abundance and composition of predator communities and leads to aphid outbreaks on urban trees

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Abstract
Urbanization can affect arthropod abundance in different ways. While species with narrow habitat range and low dispersal ability often respond negatively to urban environments, many habitat generalist species with good dispersal ability reach high densities in city centers. This filtering effect of urban habitats can strongly influence predator-prey-mutualist interactions and may therefore affect the abundance of predatory and phytophagous species both directly and indirectly. Here, we assessed the effect of urbanization on aphids, predatory arthropods, and ants on field maple (Acer campestre) trees in and around the city of Budapest, Hungary. We used the percentage of impervious surfaces within a 500 m radius of each site as an index of the degree of urbanization. We found that the abundance of aphids increased with increasing level of urbanization. However, abundance of predatory arthropods and occurrence of poorly dispersing species within the predator community were negatively related to urbanization, and we identified these two independent factors as significant predictors of aphid abundances. The abundance of ants decreased with urbanization, and contrary to our expectations, did not affect the abundance pattern of aphids. Our results suggest that urbanization, by altering the abundance and composition of predator communities, can disrupt biological control of aphid populations, and thus may contribute to the aphid outbreaks on urban trees.

Keywords Urban landscape · Predator-prey interactions · Pest control services · Community composition · Dispersal ability

Introduction
The number of people living in cities continues to grow, and the expansion of urban land cover is predicted to increase by 1.2 million km² by 2030 (Seto et al. 2012). As a result, a landscape with heterogeneous elements, from semi-natural remnant forests and public green areas to concrete surfaces and artificial structures is created, which has a wide range of functional characteristics and environmental conditions. This landscape provides an opportunity for studying complex ecological systems operating under strong human influence (Alberti 1999; McDonnell and Hahs 2008; McDonnell and Pickett 1990).

Although the number of studies examining the effects of urban landscapes on ecological patterns and processes has increased considerably in the past years, fewer studies have focused on arthropods compared to other taxa (Muderere et al. 2018). Arthropods are an ideal choice for studying the effects of urbanization as they (1) show quick responses to changed conditions, (2) have great biological diversity, (3) have relatively short generation time, and (4) represent a broad spectrum of trophic levels (McIntyre 2000). Arthropods show various responses to increasing levels of urbanization. While species that are less mobile or habitat specialist decrease in abundance or disappear altogether with increased urbanization, species that are highly mobile, or habitat generalists can reach high densities (Comont et al. 2014; McIntyre 2000; Niemelä and Kotze 2009). This process can lead to biotic
homogenization (McKinney 2006). Urbanization can also influence the abundance (Corcos et al. 2019; Denys and Schmidt 1998; Rocha and Fellowes 2020) and composition (Argáñaraz et al. 2018; Burkman and Gardiner 2014; Lüvei et al. 2019) of species at higher trophic levels, which might affect the top-down control of some herbivorous species, such as sap-sucking insects, and result in outbreaks of their populations (Meineke et al. 2017; Raupp et al. 2010; Shrewsbury and Raupp 2006).

Within the sap sucking insects, aphids (Hemiptera, Aphididae) are important worldwide pests of more than 400 plant species, including agricultural crops and ornamental plants (Blackman and Eastop 2000). They are also appropriate model organisms for studying how urbanization might affect predator-prey-mutualist systems because they (1) can reach high densities in urban areas (Denys and Schmidt 1998; Honenk et al. 2018; Parsons and Frank 2019), (2) have a wide range of natural enemies (Nagy et al. 2015), and (3) often have a mutualistic relationship with ants (Dixon 1998).

Aphidophagous arthropods have a strong tendency to aggregate on aphid colonies, and they can play a key role in setting the density of aphid populations (Dixon 1998; Markó et al. 2013; Piñol et al. 2009b; Roy et al. 2016). However, the level of predation pressure on aphids depends on several factors including the spatial and temporal distribution of aphids, their natural enemies, and mutualistic ants. Predatory arthropods can be more detrimental to aphids at the beginning of aphid population development when colonies are small, and the per capita predation risk is high (Boreau de Roincé et al. 2013; Gómez-Marco et al. 2016; Nagy et al. 2015). Highly mobile predators usually arrive in greater numbers when aphid colonies are already large, and are therefore often tracking rather than driving their abundance, while more sedentary predators that have a lower dispersal ability and higher degree of habitat fidelity might stay at the sites even when aphid density is low, and can thus suppress the aphid population increase more effectively (Markó et al. 2013; Piñol et al. 2009b; Welch and Harwood 2014).

Aphid abundance can also be affected by the presence of ants, which can act as mutualists by protecting aphid species that provide sugar-rich honeydew for them (Dixon 1998). Skinner and Whittaker (1981), for example, showed that the abundance of ant-tended species, such as Periphyllus testudinaceus (Fermie), increased in the presence of the ant Formica rufa L. on sycamore trees (Acer pseudoplatanus L.). Predators and ants might respond differently to urbanization, which could lead to different outcomes for aphid species. However, despite the importance of natural enemies in shaping herbivorous insect communities, only a few studies have investigated the effects of urbanization on predator-prey-mutualist systems (Eötvös et al. 2018; Rocha and Fellowes 2018, 2020).

Turrini et al. (2016) studied how the abundance of a non-ant-tended (non-myrmecophilous) aphid species, the vetch aphid (Megoura viciae Buckton), changed on potted broad bean (Vicia faba L.) plants in agricultural versus urban ecosystems. They applied treatments with and without predator exclusion and found that aphid abundance was determined by predation, which was less intense in city centers than in agricultural fields. As a consequence, aphids showed a pattern opposite to predators, being more abundant in the city centers. In a similar study Rocha and Fellowes (2018) examined an ant-tended (myrmecophilous) aphid species, the black bean aphid (Aphis fabae Scopoli), on the same host plant and found that the presence of mutualist ant species (Myrmica rubra L. and Lasius niger L.) overrode the effect of predators and, therefore, aphid abundance showed no difference between highly-urbanized and semi-natural areas.

In this study, we examined the abundance pattern and interactions of aphids, predatory arthropods, and ants in the canopy of field maple (Acer campestre L.) trees along an urbanization gradient. We chose A. campestre because it is highly tolerant of urban conditions (Roloff et al. 2009) and common both in rural and urban habitats with a rather uniform distribution in Hungary (Bartha et al. 2018). Moreover, A. campestre is fed on by several aphid species, which are preyed on by a large number of predatory arthropods. More specifically, we tested the following hypotheses: along the urbanization gradient (1) the abundance of aphids, predators, and ants would increase, decrease and not change, respectively; (2) considering the whole growing season, the predatory arthropods would affect negatively while ants, through their negative effect on predators, would affect positively the abundance of aphids; (3) during the peaks in aphid abundance, the populations of predatory arthropods and ants would track the abundance of aphids; (4) the composition of the predator community would change along the urban gradient. Finally, we also hypothesized (5) that a shift in predator community composition towards species with high dispersal ability would lead to higher aphid abundance.

Methods

Study area and arthropod collection

We selected 22 sites for this study in or near the city of Budapest, Hungary (Fig. S1), representing a range of semi-natural rural, suburban, and urban areas. We chose three field maple (A. campestre) trees at each site for arthropod collection by having similar trunk and canopy diameters and the shortest distance between them. The distance between the trunks of the sampled trees was 2–15 m. There was no significant effect of urbanization (defined as the proportion of impervious surfaces) on the trunk diameters (diameter at breast height)
selected trees (Pearson $r = 0.24, p = 0.28$) and on the distance between the selected trees (Pearson $r = 0.14, p = 0.55$).

A beating funnel 70 cm in diameter, 60 cm in depth, together with a 120 cm long beating stick, was used to collect arthropods from the canopy of the trees (Basset et al. 1997). In order to standardize sampling effort, ten branches were sampled per tree. Branches were sampled monthly on seven dates in 2016 (April 26, May 25, June 22, July 18, August 15, September 13, and October 14). In 2017, we also sampled on three additional dates (May 5, July 16, and September 27). On all sample dates, we identified and counted all collected aphids (Aphididae) to species. Predatory arthropods (Hemiptera, Coccinellidae, Dermaptera, Neuroptera, and Araneae) were identified to species or, if not possible, to genus or family, using characters of the exoskeleton or genitalia. The collected arthropods were deposited at the Department of Entomology of Szent István University. Because of the low aphid parasitism rates observed, parasitoids were not included in this study.

Dispersal ability of predators

Aphidophagous predator species were classified into five dispersal groups following the methodology described by Gossner et al. (2015) using data of different literature sources (Bell et al. 2005; Blandenier 2009; Duffey 1956; Gossner et al. 2015; Wachmann et al. 2004–2012). We considered wing morphology for true bugs, ladybirds, and lacewings (always brachypterous, predominantly brachypterous, equally brachypterous and macropterous, predominantly macropterous, and always macropterous; coded 0.00, 0.25, 0.50, 0.75 and 1.00 respectively), and ballooning ability for spiders (non-ballooning species [code: 0.00], species with weak ballooning propensity [code: 0.25], ballooning species observed outside the main habitat less frequently [code: 0.50] and frequently [0.75], and species with the highest ballooning propensity and dispersal distances [code: 1.00]). For juvenile spider individuals which were identified to genus or species group (e.g. Philodromus aureolus and rufus group) level, we used the dispersal trait values of adult individuals representing the same genus or species group weighted by their relative abundances. The collected dermapteran species were considered as species with the lowest dispersal category (code: 0.00) as Apterygida media (Hagenbach) and Chelicurella acanthopygia (Géné) are apterous species, and although Forficula auricularia L. has wings, it hardly ever uses them (Crumb et al. 1941) and the mobility of this species is very low (Moerkens et al. 2010). For the assigned dispersal trait values, see Table S1.

Landscape structure

We calculated landscape composition around each sampling site within a 500 m radius buffer with the OpenStreetMap landcover map, using OpenLayers plugin of Quantum GIS 2.16 software (Quantum GIS Development Team 2018). This radius has been found to be appropriate spatial scale for analyzing the relationship between landscape pattern and aphid-ant-predator relationships (Schüepp et al. 2014; Stutz and Entling 2011). The geographic center (centroid) of a triangle formed by the sampled trees was used as the center of each study site. To characterize the landscape composition of the study sites, we considered six land cover types (buildings, roads, forests, gardens, meadows, and parks) and calculated their percentage cover in a buffer around each site (Table S2). We excluded water bodies from our analyses due to their low abundance within the sampled landscapes.

Analyses

First, to explore the correlations among the six landscape variables, we calculated the Kendall rank correlation coefficients for all pairs of the six land cover variables. We found significant correlations between most landscape variables (Table S3); therefore, we used only the proportion of impervious surfaces (pooled proportion of buildings and roads) as an explanatory variable in further analyses. Finally, a gradient was created from this variable among the sites that ranged between 2 and 95% surface with impervious cover (Table S2) indicating the degree of urbanization. For all analyses, we pooled the number of individuals collected (from three trees) at each site. For analysis of annual abundance, we pooled data across sampling dates in both years and analyzed them separately. The spatial autocorrelation in the residuals of our models was tested by Moran’s I tests with inverse distance weighting using ‘spdep’ package version 1.1–3 (Bivand and Wong 2018). There was no evidence of residual spatial autocorrelation in any model ($p > 0.21$ in all cases), and therefore we did not add an auto-covariate to our models. We used R 3.4.4 statistical environment (R Core Team 2018) for all analyses.

Hypothesis 1: Effects of urbanization on the abundance patterns of aphids, predators and ants

To test how annual and monthly abundance of aphids, predators, and ants (response variables) depended on the proportion of impervious surfaces (explanatory variable), we used Poisson-distributed generalized linear models (GLMs). The models included the percentage of impervious surfaces as a fixed factor. Because we detected overdispersion in the abundance data, we corrected the standard errors using generalized linear models with quasi-Poisson distribution (quasi-GLMs) (Zuur et al. 2009). In case of monthly analyses, $p$ values were adjusted using the method of Benjamini and Hochberg (1995).
Hypotheses 2 and 3: Aphid-predator-ant interactions

Since the effects of predators and mutualistic ants on aphid population development might be accumulated and delayed, we tested how annual aphid abundance (response variable) depended on annual and monthly abundance of predators and ants (explanatory variables) using quasi-GLMs. We fitted the models separately for each sample year and month. The full models included predator and ant abundance as fixed factors, and model selection retained only the models with significant variables ($p < 0.05$). In the analyses, where monthly abundances of predators were considered as explanatory variables, $p$ values were adjusted with the Benjamini-Hochberg method.

To determine whether there is any relationship between predator and ant abundances in each year, we built quasi-GLMs including the abundance of predators as response variable and the abundance of ants as explanatory variable (fixed factor).

We also tested how abundance of predators and ants was influenced by the abundance of aphids (tracking) in different years and seasons using generalized linear mixed effects models with quasi-Poisson errors (penalized quasi-likelihood GLMM, GLMMpQL) in the package ‘MASS’ version 7.3–51.4 (Venables and Ripley 2002). In order to obtain a meaningful quantitative response, only the months of peak aphid abundance were considered (April, May and October in 2016; May and September in 2017). In the model, the response variable was the abundance of predators or ants, and abundance of aphids, season, and year were entered as fixed factors. To control for the possible effect of sampling location on the response variables, we included sampling site as a random intercept factor.

Hypothesis 1 and 4: Species of aphids and predators and taxonomic composition of predator community

To analyze how the observed number of aphid or predatory species, the number of species groups, and genera (response variables) responded to the proportion of impervious surfaces (explanatory variable), we used quasi-GLMs. Only those arthropod groups (species, species groups, or genera) represented by at least 50 individuals in each year’s samples were included in the analyses (for species groups and genera, see Table S4). In the case of Harmonia axyridis Pallas, juvenile and adult stages were analyzed separately. The full model included percentage of impervious surfaces as a fixed factor. The data from the two years (2016 and 2017) were analyzed separately. In these analyses, $p$ values were adjusted using the Benjamini-Hochberg method.

We performed non-metric multidimensional scaling (NMDS) to examine the differences in composition for the predator community and within the three largest predatory assemblages: true bugs (Heteroptera), aphidophagous ladybirds (Coccinellidae), and spiders (Araneae), each separately. We used the Bray-Curtis distance measure to compute the resemblance matrix among sites. The resulting matrix was the basis for creating a NMDS ordination, in which sites were ranked based on their similarity and then plotted in 2-dimensional ordination space. The analyses were based on species-level abundance data. Because we collected samples in both years only in May, July, and September, the abundance data of these three months were pooled across years for the NMDS analyses of true bugs, aphidophagous ladybirds, and spiders. We generated smooth surfaces along impervious surface gradient with generalized additive models (GAMs). This function automatically selects the degree of smoothing by generalized cross-validation and interpolates the fitted values on the NMDS plot. NMDS was performed using ‘metaMDS’ function, while GAMs were run using ‘odisurf’ function of the package ‘vegan’ version 2.5–3 (Oksanen et al. 2018).

Hypothesis 5: Effect of dispersal ability of predators on aphids

We calculated community weighted means (CWM) for dispersal trait of predators using the averages of trait values weighted by the relative abundances of each predator species at each study site ($n = 22$) (Ricotta and Moretti 2011). We used the ‘FD’ package version 1.0–12 to calculate CWM indices (Laliberte and Legendre 2010). In case of some spider individuals, dispersal trait classification was not possible, due to the family level identification ($n = 23$) or the lack of adult representatives of the genus ($n = 24$) (Table S1), thus these individuals were excluded from the analyses. We used general linear models to test whether percentage of impervious surfaces (explanatory variable, fixed factor) had a significant effect on CWM of dispersal trait (response variable) in 2016 and 2017 separately.

Quasi-GLMs were used to assess whether predators with lower dispersal ability would have a more important role in aphid suppression than those with higher dispersal ability. The models included annual abundance of aphids as a response variable and CWM of dispersal ability of predator community as an explanatory variable (fixed factor) for each year.

Finally, predators were classified into eleven groups based on their taxonomic status and abundance: (1) Dermaptera, (2) Deraeocoris lutescens, (3) other heteropterans (all Heteroptera except D. lutescens), (4) H. axyridis larvae, (5) H. axyridis adults, (6) other coccinellids, (7) Neuroptera, (8) web building spiders, (9) the Philodromus aureolus group, (10) the Philodromus rufus group, and (11) other hunting spiders (for detailed data see Table S1). We used NMDS to analyze how these species and groups were associated to aphid abundance using ‘metaMDS’ function. For this analysis, we summed data for the two sample years for the abundance of aphids and for each predator species or group. We
used Euclidean distance measure for comparing similarities and generated smooth surfaces along aphid infestation with GAMs using ‘odisurf’ function.

**Results**

**Taxa recorded**

In total, we collected 10,197 individual aphids (Sternorrhyncha, Aphididae), 8955 predators, and 3555 ants (Hymenoptera, Formicidae) from the canopy of A. campestre trees. The collected aphids belonged to four genera and nine species. Among these, two ant-tended species, *Periphyllus testudinaceus* and *Periphyllus obscurus* Mamontova, were collected in the largest numbers (59.5% and 30.9% of total aphid abundance, respectively) followed by two non-ant-tended species, *Drepanosiphum aceris* Koch and *Drepanosiphum platanoidis* (Schrank), which occurred only in small numbers (2.1% and 1.8% of the total aphid abundance, respectively). The most abundant ant species were *Prenolepis nitens* (Mayr), *L. niger*, and *Lasius emarginatus* (Olivier) (32.7%, 21.6%, and 10.4% of the ants collected, respectively). These ant species are known for honeydew utilization and having mutualistic relationships with aphids (Miñarro et al. 2010; Purkart et al. 2019), thus we assumed that they have mutualistic interactions with the myrmecophilous aphid species (90.5% of aphids collected).

The predators we collected comprised 24 families, 103 genera, and 145 species. They consisted mainly of spiders (Araneae; 32.4%), coccinellids (28.2%), mirid bugs (25.9%), forficulid earwigs (9.3%), and, to a lesser degree, lacewings (Neuroptera, Chrysopidae, and Hemerobiidae; 1.6%), anthocorid flower bugs (1.6%), and other species (1.3%). The most abundant predator species were *Deraeocoris lutescens* (Schilling) (Miriidae, n = 1450), *Harmania axtys* (Coccinellidae, n = 1439), *Forficula auricularia* (Forficulidae, n = 675), and two species groups of spiders—*Philodromus aureolus* group (n = 752) and *Ph. rufus* group (n = 692) (Philodromidae) (Tables S1 and S4).

**Effects of urbanization on the abundance of aphids, predators and ants**

Aphids were abundant in spring and autumn, while in summer they were almost absent. Predator abundance was lowest in spring and increased throughout the growing season. The monthly abundances of ants showed a less clear pattern. They were abundant in April and June in 2016 and in May and September in 2017 (Fig. 1). For the monthly abundances of each aphid species and predatory groups, see Fig. S2.

The annual abundance of aphids increased significantly with increase in the percentage of impervious surfaces in both years (2016: $p = 0.008$; 2017: $p < 0.001$). In contrast, the abundance of predators and ants was negatively affected by the percentage of impervious surfaces, and these relationships were also significant for both groups ($p = 0.039$ and 0.013, respectively) in 2016 and nearly significant for ants ($p = 0.073$) in 2017 (Table 1, Fig. 2).

The pattern of insects along the urbanization gradient also varied over time. Aphid abundance was positively affected by the percentage of impervious surfaces in spring and autumn, but not in May 2016, when this relationship was negative but not significant ($p = 0.054$) (Figs. 3 and 4, Table S5). We found only a weak correlation between aphid abundances in autumn 2016 and spring 2017 (Spearman $r = 0.39$, $p = 0.07$). For predators, abundance increased slightly with increasing levels of urbanization in spring, but this pattern changed from the beginning of summer, and the abundance was higher in seminatural habitats for the rest of the season. This temporal pattern was more pronounced in 2016 than in 2017. The abundance of ants slightly decreased with the increasing percentage of impervious surfaces on all sampling occasions. Surprisingly, predators and ants did not follow the spatial abundance pattern of aphids in spring and even less in autumn (Figs. 3 and 4, Table S5).

**Aphid-predator-ant interactions**

According to the best models, the presence of predators negatively affected the total abundance of aphids in 2016 ($p = 0.013$, Table S6). We found significant negative relationship between monthly predator and total aphid abundance in September and October in 2016. However, there was no association between the yearly or monthly abundances of ants and the total aphid density, either in 2016 or in 2017 (Tables 2 and S7). We did not find any significant relationship between yearly abundances of predators and ants in either year (Quasi-GLMs: 2016: $t = 0.593$, $p = 0.560$; 2017: $t = -1.176$, $p = 0.253$).

During the months of peak aphid abundance, number of aphids had no significant effect on the abundance of predators ($p = 0.483$) or ants ($p = 0.408$), i.e. predators and ants did not track aphid abundance (Table S8).

**Species of aphids and predators and taxonomic composition of predator community**

The main aphid species showed a positive response to percentage of impervious surfaces. This relationship was significant in the cases of *P. testudinaceus* in 2016 and *P. obscurus* in 2017 (Table 3, Fig. S3).

The earwig *Aptyrygida media* was strongly associated with sites with the lowest percentage of impervious surfaces (where there was a high percentage of forests). In contrast, *F. auricularia* were most abundant at sites with high
percentages of gardens or urban parks and showed no relationship to the degree of urbanization (Table 3, Fig. S4, for the percentage of each landscape element see Table S2).

Among the true bugs, the abundance of *Deraeocoris flavilinea* (A. Costa) increased with the increasing level of impervious surfaces. Urbanization had no significant effect on the abundance of the rest of the true bug species (Table 3, Fig. S4).

Among aphidophagous coccinellids, the density of *H. axyridis* larvae increased significantly with urbanization, while that of *H. axyridis* adults decreased slightly, in 2016. Urbanization had no effect on the abundance of the rest of the coccinellid species (Table 3, Fig. S5).

Most of the spider species responded negatively to the increase of impervious surfaces. This was particularly true for *Anyphaena accentuata* (Walckenaer), *Araniella* spp. [mainly *A. cucurbitina* (Clerck), and *A. opisthographa* (Kulczyński)], and for the groups ‘other Theridiidae’ (mostly juveniles), ‘other Thomisidae’, but to a lesser extent for *Ph. aureolus* group. Only *Xysticus* spp. (mostly juveniles) had significantly higher abundance in built-up areas. *Anelosimus viitatus* (CL Koch), *Cheiracanthium* spp. (mainly *C. mildei* L. Koch) and the *Ph. rufus* group showed no response to urbanization (Table 3, Fig. S6, for further information on the species groups and genera, see Table S4).

Lacewing abundance (mostly of *Chrysoperla carnea* [Stephens]) decreased slightly with increasing levels of urbanization in 2016 (Table 3, Fig. S5).

NMDS ordination revealed that the community composition of predators changed along the urbanization gradient in
both 2016 and 2017 (NMDS, Stress = 0.186, $R^2 = 0.674$; Stress 0.203, $R^2 = 0.715$, respectively) (Fig. 5a–b). This pattern was most typical for the spider assemblage (NMDS, Stress = 0.173, $R^2 = 0.539$) and to a lesser extent for true bugs and coccinellids (NMDS, Stress = 0.203, $R^2 = 0.600$; Stress = 0.198, $R^2 = 0.547$, respectively) (Fig. 5c–e). The composition of coccinellids did not separate clearly along the first axis, although it showed differences between the most natural (impervious surface <20%) and highly urbanized sites (impervious surface >60%) (Fig. 5d).

**Effect of dispersal ability of predators on aphids**

We found a significant shift in the CWM trait values for predator dispersal ability along the urbanization gradient in both years, where the proportion of predatory species with higher dispersal ability increased with the increasing level of impervious surfaces (GLMs: 2016: $t = 3.156, p = 0.005$; 2017: $t = 2.448, p = 0.024$) (Fig. 6), and the annual abundance of aphids decreased significantly when the predator community contained more low-dispersing predators (Quasi-GLMs: 2016: $t = 3.05, p = 0.006$; 2017: $t = 2.385, p = 0.027$).

According to the NMDS ordination based on the association between the abundance of aphids and predatory groups earwigs (mean dispersal value of the group: 0.0, $n = 831$) and web building spiders (0.61, 649) were associated with the sites with the lowest, while *H. axyridis* larvae (1.0, 255) and other coccinellids (1.0, 1082) were associated with the sites with the highest aphid abundances (NMDS, Stress = 0.113, $R^2 = 0.227$; Stress = 0.155, $R^2 = 0.359$, respectively) (Fig. 7).

**Discussion**

The aphid-predator-ant system we studied changed markedly both in space, along the urbanization gradient, and in time, across the season, in both study years. Aphids and predatory arthropods showed species-specific abundance patterns. There was a negative relationship between predator and aphid abundances in one of the two study years and the abundance of aphids decreased significantly with increasing number of low-dispersing individuals in the predator community in both years.

**Effects of urbanization on the abundance of aphids, predators and ants**

Aphids exhibited the most consistent pattern in this study. In line with our first hypothesis, we found high aphid
abundances in highly urbanized areas, in contrast to semi-natural landscapes where their abundance remained low (Table 1, Fig. 2). Aphids benefit from urbanization disturbance due to the environmental conditions and characteristics of their host plants in cities (Braun and Flückiger 1985; Butler and Trumble 2008; Hall and Ehler 1980) coupled with their enormous reproductive potential and multiple generations per year, which allows them to rapidly take advantage of favorable conditions (Durak et al. 2016; Harrington et al. 2007; Parsons and Frank 2019).

In contrast to aphids, the overall number of predators decreased towards the city center (Table 1, Fig. 2), which can be explained by several factors. Highly urbanized areas are generally characterized by fragmented habitats with reduced plant density, diversity, and structural complexity (Raupp et al. 2010). This may create an unfavorable environment for predators due to reduced numbers of prey (and alternative resources), refuges, and suboptimal climatic conditions (Langellotto and Denno 2004). Moreover, the increasing proportion of human-made structures might act as a dispersal
barrier for predatory arthropods (Denys and Schmidt 1998; Mader et al. 1990; Turrini et al. 2016).

Lessard and Buddle (2005) found the highest ant abundance in residential backyards compared to forests while Rocha and Fellowes (2018) showed that ants were present in high numbers even at the most urbanized sites. Contrary to their findings, we observed that ants had the highest abundance in semi-natural areas and the lowest in the city center (Table 1, Fig. 2). Although we did not determine the species composition of ants along the gradient, we assume that highly urbanized areas are less suitable habitats for most of the ant species. The explanation for this could be colonization limitation, as individual ants may have more limited opportunities to spread to more isolated areas from source populations (Pacheco and Vasconcelos 2007), but also may be due to a lack of necessary resources for nesting sites (Vepsäläinen et al. 2008) and a decline in soil moisture (Clarke et al. 2008).

Generally, the activity of tree-dwelling aphids is characterized by two seasonal peaks (Dixon 1977; Piñol et al. 2009a, 2009b), and we found the same pattern here (Fig. 1). Periphyllus species aestivate during summer months as first instar nymphs, while Drepanosiphum species aestivate as adults (Dixon 1977; Dixon 1998; Furuta 1985; Junkiert et al. 2011). Although the total abundance of aphids increased towards the city center in both years, their monthly abundance showed a less consistent pattern. The abundance of aphids greatly increased in April, but slightly decreased in May towards the city center in 2016 (Fig. 3, Table S5). Highly urbanized areas generally have higher temperatures than the surrounding semi-natural habitats due to the urban heat island effect caused by the high proportion of impervious surfaces (Long et al. 2019; Rizwan et al. 2008). Since temperature is considered one of the most important factors in aphid development (Durak et al. 2016; Harrington et al. 2007), it is possible that warmer temperatures resulted in faster nymphal development (phenological shift) at these sites compared to semi-natural habitats (Fig. 3, Table S5). Higher temperature can also lead to delayed appearance of sexual generations and reproduction of aphids at the end of summer (Durak et al. 2016). Based on these factors, we expected an earlier peak in aphid abundance in semi-natural habitats compared to highly urbanized ones in this period, but such a pattern was not observed (Fig. 3). The lack of a phenological shift in the later part of the season may be explained by greater predator

Table 2 Results of the best quasi-GLMs (after model selection) for the annual abundance of aphids (response variable) depending on the monthly abundance of predators and ants (explanatory variables)

| Explanatory variable | Month | Estimate | SE  | Exp. Estimate | Adj. p | Unadj. p | Pseudo-$R^2$ |
|----------------------|-------|----------|-----|---------------|--------|----------|--------------|
| Predators            | June  | −0.013   | 0.007| 0.987         | 0.148  | 0.089    | 14.9         |
| Predators            | September | −0.024 | 0.008| 0.976         | 0.030  | 0.006    | 34.4         |
| Predators            | October | −0.010  | 0.004| 0.990         | 0.035  | 0.014    | 27.5         |

Only significant ($p<0.05$, numbers in bold) and marginally significant ($p\geq 0.05$ and $<0.1$, number in italics) results are shown for the best models (in 2016). The main outcomes of the best and full models are summarized in Table S7.
Table 3  Relationships between impervious surface percentage and the abundance of aphid and predator taxa

| Taxon                         | 2016 Estimate | SE  | Exp. Estimate | Adj. p | Pseudo-R²  | 2017 Estimate | SE  | Exp. Estimate | Adj. p | Pseudo-R²  |
|-------------------------------|--------------|-----|---------------|--------|------------|--------------|-----|---------------|--------|------------|
| Sternorrhyncha                |              |     |               |        |            |              |     |               |        |            |
| Drosophila aceris             | 0.008        | 0.012| 1.008         | 0.514 (0.514) | 3.1         |              |     |               |        |            |
| Drosophila platanioides       | 0.018        | 0.008| 1.018         | 0.099 (0.048) | 23.9        |              |     |               |        |            |
| Derpyllus obscurus            | 0.008        | 0.005| 1.008         | 0.192 (0.144) | 11.0        | 0.032        | 0.005| 1.033         | 0.000 (0.000) | 74.5 |
| Derpyllus testudinaceus       | 0.011        | 0.004| 1.011         | 0.049 (0.012) | 28.0        | 0.011        | 0.006| 1.011         | 0.111 (0.111) | 13.2 |
| Dermaptera                    |              |     |               |        |            |              |     |               |        |            |
| Apterygida media              | −0.112       | 0.049| 0.894         | 0.068 (0.034) | 53.3        | −0.133       | 0.046| 0.876         | 0.019 (0.010) | 55.6 |
| Forficula auricularia         | −0.008       | 0.007| 0.992         | 0.270 (0.270) | 6.3         | −0.002       | 0.009| 0.998         | 0.820 (0.820) | 0.3  |
| Heteroptera                   |              |     |               |        |            |              |     |               |        |            |
| Deraeocoris flavilinea        | 0.021        | 0.007| 1.021         | 0.035 (0.005) | 34.8        |              |     |               |        |            |
| Deraeocoris lutescens         | −0.005       | 0.005| 0.995         | 0.490 (0.350) | 4.9         | −0.003       | 0.009| 0.997         | 0.761 (0.761) | 0.6  |
| Orius spp.                    | 0.000        | 0.007| 1.000         | 0.992 (0.992) | 0.0         | −0.003       | 0.007| 0.997         | 0.761 (0.733) | 0.5  |
| Phytoecoris tiliae            | 0.018        | 0.008| 1.018         | 0.103 (0.044) | 18.7        |              |     |               |        |            |
| Pilophorus perplexus          | −0.036       | 0.016| 0.964         | 0.103 (0.034) | 33.7        |              |     |               |        |            |
| Psyllus assimilis             | 0.006        | 0.005| 1.006         | 0.424 (0.242) | 6.3         | 0.029        | 0.015| 1.030         | 0.199 (0.066) | 27.5 |
| Reuteria marqueti             | −0.007       | 0.014| 0.993         | 0.708 (0.606) | 1.8         |              |     |               |        |            |
| Coleoptera                    |              |     |               |        |            |              |     |               |        |            |
| Coccinella septempunctata     | −0.005       | 0.014| 0.995         | 0.755 (0.717) | 1.2         |              |     |               |        |            |
| Ecochosoma quadripustulatus   | −0.002       | 0.007| 0.998         | 0.755 (0.755) | 0.5         | 0.004        | 0.006| 1.004         | 0.490 (0.490) | 2.2  |
| Harmonia axyridis larvae      | 0.015        | 0.004| 1.015         | 0.019 (0.003) | 35.1        |              |     |               |        |            |
| Harmonia axyridis adults      | −0.011       | 0.005| 0.998         | 0.121 (0.047) | 21.5        | −0.004       | 0.004| 0.996         | 0.490 (0.380) | 4.2  |
| Oenopia congobalata           | 0.014        | 0.007| 1.014         | 0.121 (0.072) | 15.7        | 0.018        | 0.012| 1.018         | 0.466 (0.155) | 12.2 |
| Scymnus spp.                  | 0.013        | 0.007| 1.013         | 0.121 (0.080) | 13.0        |              |     |               |        |            |
| Neuroptera                    |              |     |               |        |            |              |     |               |        |            |
| Chrysoperla spp.              | −0.016       | 0.008| 0.984         | 0.078 (0.078) | 15.3        |              |     |               |        |            |
| Araneae                       |              |     |               |        |            |              |     |               |        |            |
| Anelosimus vitatus            | −0.017       | 0.005| 0.983         | 0.012 (0.001) | 41.7        | −0.020       | 0.009| 0.981         | 0.072 (0.040) | 26.8 |
| ‘other Theridiidae’            | −0.087       | 0.039| 0.917         | 0.091 (0.040) | 70.2        | −0.091       | 0.030| 0.913         | 0.023 (0.008) | 62.3 |
| Anyphaena accentuata          | −0.022       | 0.010| 0.978         | 0.091 (0.038) | 26.2        | −0.032       | 0.010| 0.968         | 0.018 (0.004) | 48.0 |
| Araniella spp.                | 0.005        | 0.006| 1.005         | 0.470 (0.418) | 3.1         | −0.003       | 0.005| 0.997         | 0.624 (0.485) | 1.9  |
| Chaetocnemis spp.             |              |     |               |        |            |              |     |               |        |            |
| Philodromus spp. aureolus gr. | −0.007       | 0.003| 0.993         | 0.091 (0.035) | 20.2        | −0.009       | 0.004| 0.991         | 0.078 (0.052) | 18.1 |
| Philodromus spp. rufus gr.     | 0.004        | 0.008| 1.004         | 0.585 (0.585) | 1.9         | 0.003        | 0.008| 1.003         | 0.820 (0.729) | 0.7  |
| Xysticus spp.                 |              |     |               |        |            |              |     |               |        |            |
| ‘other Thomisidae’             | −0.088       | 0.058| 0.916         | 0.213 (0.142) | 56.0        | −0.052       | 0.016| 0.949         | 0.018 (0.004) | 56.4 |

Summary of the quasi-GLM results by species and years. Numbers with bold indicate significant (< 0.05) and numbers with italic indicate marginally significant (≥ 0.05 and < 0.1) p values. Numbers in parentheses are the unadjusted p values.

Abundances and therefore higher predation rates in semi-natural areas that might obscure the impact of climate on aphids.

The overall number of predators was relatively low and only increased slightly towards the city center in spring (mostly due to H. axyridis adults), and this pattern was most pronounced in autumn (mostly due to spiders and coccinellids), including H. axyridis adults, Predatory insects with high dispersal ability and ants exhibit positive density-dependent numerical responses to aphid density (Brown 2004; Leigh and van Emden 2017; Markó et al. 2013; Stutz and Entling 2011). However, in our study, predatory arthropods and ants showed a reverse pattern...
compared to aphids, i.e. the abundance of predators and ants did not track the abundance of aphids along the urbanization gradient even at the aphid peaks in spring and autumn (Figs. 3 and 4, Tables S5 and S8). This finding indicates that urbanization may lead to a spatial mismatch between aphids, their predators, and mutualistic ants.

**Aphid-predator-ant interactions**

In line with our second hypothesis, high predator numbers were accompanied by low aphid abundance in 2016, but contrary to our expectations (and hypothesis 3), there was no relationship between ant and aphid abundance (Tables 2 and S7), and predatory arthropods and ants varied independently of each other. One explanation for the observed higher aphid densities in more urbanized areas may be that urbanization disrupted predators, resulting in weakened top-down control on aphids (Turrini et al. 2016). Predators reached their highest abundance in autumn (Fig. 1; Table S8), and predator abundances were the best predictors of the total aphid abundance in this period in 2016 (Tables 2 and S7). This suggests that number of aphids is primarily regulated by predators before and during their second activity peak in semi-natural areas. Furuta (1985) pointed out that aestivating populations of *Periphyllus californiensis* (Shinji) on maple trees seem to face high predation in the second half of the growing season, and this may have a stronger effect on their survival than other factors such as unfavorable climatic conditions.

The highest aphid densities were coincided with the lowest predator densities at the beginning of the growing season. This
implies that aphid species with early egg hatch (especially *P. testudinaceus*) are exposed to lower predation in spring (Dixon 1998), and therefore the peak in aphid density early in the season might be determined more by aphid abundance in the previous autumn than by predation in early spring. Rocha and Fellowes (2018) reported that instead of predation, the abundance of mutualistic ants drove the abundance of myrmecophilous aphids along the urbanization gradient. Contrary to their results and to those of Skinner and Whittaker (1981), we found that the supposed positive effect of ants on aphids did not override the predation pressure on aphids (almost exclusively ant-tended *Periphyllus* spp.) in semi-natural areas.

**Species of aphids and predators and taxonomic composition of predator community**

The ant-tended aphid species (*P. obscurus*, *P. testudinaceus*) responded positively, while the non-ant-tended species (*D. aceris*, *D. platanoidis*) showed no or weak responses to increasing level of urbanization. In contrast, and in line with hypothesis 4, predatory species showed definitive species-specific responses (Table 3). Seven predatory groups responded negatively, whereas three groups responded positively to urbanization in at least one of the studied years. Fourteen groups showed no response to urbanization and were occasionally abundant in moderately built-up sites but less often in highly built-up areas.

One earwig species and most of the spider groups were negatively affected by urbanization. In contrast, most true bugs and coccinellids showed no response or responded positively to increasing levels of urbanization (Table 3, Figs. S4 and S5).

In accordance with species-specific responses, the composition of spiders changed more along the urbanization gradient than that of true bugs or coccinellids (Fig. 5c-e), and based on this, spiders contributed most significantly to the changing composition pattern of the total predator community in both years (Fig. 5a-b). These results suggest that although some spider groups have a wide habitat range and good dispersal ability (Alaruikka et al. 2002; Blandenier 2009; Duffey 1956), they are less capable of target-oriented movement than the winged species with relatively high mobility and short residency time such as coccinellids and true bugs (Egerer et al. 2016; Piñol et al. 2009b). Therefore, spiders can be highly sensitive to habitat isolation and its consequences caused by urbanization (Argañaraz et al. 2018; Langellotto and Denno 2004; Meineke et al. 2017).

**Effect of dispersal ability of predators on aphids**

We found that the dispersal capacity of predator community significantly increased with increasing level of
urbanization, suggesting that urbanization filters predatory species based on their dispersal ability (Merckx and Van Dyck 2019; Piano et al. 2017). Furthermore, our observations are consistent with hypothesis 5 and suggest that generalist predators with low dispersal ability such as earwigs and some spiders may play a major role in the biological control of aphids. Some of these predators can be present constantly on the trees, even at early aphid population growth and low aphid density (Piñol et al. 2009b), which can be an important prerequisite for successful aphid suppression (Bureau de Roincé et al. 2013; Gómez-Marco et al. 2016; Nagy et al. 2015).

Earwigs are voracious predators of aphids and play a key role in the top-down control of these pests (Carroll and Hoyt 1984; Mueller et al. 1988; Piñol et al. 2009a). They are characterized by low dispersal ability and a preference for habitats with higher humidity (Kirstová et al. 2019). Therefore, increasing isolation of trees from forests is associated with lower earwig numbers and, as a consequence, greater density of aphids (Stutz and Entling 2011). There are a few examples where hunting spiders (e.g. Anyphaena, Philodromus spp.) contributed to the early biological control of aphids (Bureau de Roincé et al. 2013; Lefebvre et al. 2017). Moreover, studies showed that web-building spiders can substantially reduce the number of aphids, especially in autumn when density of their webs is high and winged aphids are present. This can lead to fewer aphid fundatrices the following spring (Cahenzli et al. 2017; Wyss et al. 1995).

Our results are in accordance with the finding that predators with high dispersal capacity such as lady beetles and true bugs can be generally less effective at keeping aphid populations at low levels, as they mostly track aphid density rather than control it (Piñol et al. 2009a, 2009b).

Conclusions

The most important finding of our study is that an increasing level of urbanization resulted in increasing aphid infestations on urban maple trees, while predators showed opposite abundance pattern. Overall dispersal ability of predator community was positively related to urbanization, i.e. predatory species with low dispersal but high aphid control ability (such as earwigs and some spiders) decreased in abundance toward city centers. In contrast, species with higher dispersal potential but a lower ability to prevent aphid outbreaks such as coccinellids and true bugs did not aggregate in sufficient number on the abundant aphid colonies to compensate for the lack of early aphid control in the city centers by other predators. We concluded that aphid populations in semi-natural habitats are primarily regulated at low aphid densities by predators with low dispersal ability. In this study, ants were less abundant in urban than in the semi-natural habitats and did not drive aphid density.

Our results suggest that pest control services provided by predatory arthropods decrease markedly towards city centers and are associated with aphid outbreaks on urban trees. To confirm this, further field studies combined with experimental approaches are needed.

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