Functional diversity and habitat preferences of native grassland plants and ground-dwelling invertebrates in private gardens along an urbanization gradient

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Abstract
Urbanization is occurring around the globe, changing environmental conditions and influencing biodiversity and ecosystem functions. Urban domestic gardens represent a small-grained mosaic of diverse habitats for numerous species. The challenging conditions in urban gardens support species possessing certain traits, and exclude other species. Functional diversity is therefore often altered in urban gardens. By using a multi-taxon approach focused on native grassland plants and ground-dwelling invertebrates with overall low mobility (snails, slugs, spiders, millipedes, woodlice, ants, rove beetles), we examined the effects of urbanization (distance to city center, percentage of sealed area) and garden characteristics on functional dispersion, functional evenness, habitat preferences and body size. We conducted a field survey in 35 domestic gardens along a rural–urban gradient in Basel, Switzerland. The various groups showed different responses to urbanization. Functional dispersion of native grassland plants decreased with increasing distance to the city center, while functional dispersion of ants decreased with increasing percentage of sealed area. Functional evenness of ants increased with increasing distance to the city center and that of rove beetles decreased with increasing percentage of sealed area. Contrary to our expectation, in rove beetles, the proportion of generalists decreased with increasing percentage of sealed area. Functional evenness of ants increased with increasing distance to the city center and that of rove beetles decreased with increasing percentage of sealed area. Contrary to our expectation, in rove beetles, the proportion of generalists decreased with increasing percentage of sealed area. Functional evenness of ants increased with increasing percentage of sealed area in the surroundings, and the proportion of species preferring dry conditions increased with increasing distance to the city center. Body size of species increased with distance to city center for slugs, spiders, millipedes, ants, and rove beetles. Local garden characteristics had few effects on functional diversity and habitat preferences of the groups examined. Our study supports the importance of using multi-taxon approaches when examining effects of environmental change on biodiversity. Considering only a single group may result in misleading findings for overall biodiversity. The ground-dwelling invertebrates investigated may be affected in different ways from the more often-studied flying pollinators or birds.
1 | INTRODUCTION

Urbanization is currently one of the major drivers of global change (Güneralp et al., 2020) with multifarious consequences for biodiversity and ecosystems (McDonald et al., 2020). Among others, negative impacts derive from habitat loss and fragmentation due to the increase of impervious surfaces and barrier effects of the urban structure (Beninde et al., 2015; Fenoglio et al., 2020), an increase in temperature due to the urban heat island effect (Hamblin et al., 2018; Piano et al., 2017; Youngsteadt et al., 2017), and direct anthropogenic disturbance (Delgado de la flor et al., 2019). Despite these negative effects, cities may also offer unexploited opportunities for biodiversity (Samways et al., 2020; Soanes & Lentini, 2019). For example, urban habitats can serve as secondary habitats or refuges, respectively, for rare and endangered species, whose primordial habitats are degraded by the intensification of agriculture or even destroyed (Hall et al., 2016; Ives et al., 2016).

Urban gardens constitute an important part of urban green space and despite their small size, they cover large parts of urban area (Gaston et al., 2013). Home or community gardens are distributed throughout the city and provide valuable stepping-stones for numerous species and thus are an essential part of the urban green infrastructure (Cameron et al., 2012). At a local scale, gardens provide a heterogeneous small-grained mosaic of diverse habitats formed through different user management practices and individual owner preferences (Braschler et al., 2021; Lin & Egerer, 2020; Smith et al., 2005). Gardens may harbor a considerable variety of vascular plants, which in turn increases biodiversity of herbivores and decomposers (Adams et al., 2020). Therefore, invertebrate biodiversity of urban gardens could be remarkable as has been shown for cities in the United Kingdom (Smith, Chapman, & Eggleton, 2006; Smith, Gaston, et al., 2006), the United States (Egerer et al., 2017; Philpott et al., 2014), and Switzerland (Braschler et al., 2020).

While studies on biodiversity aspects in cities have increased for certain taxa (e.g., spiders, ground beetles or pollinators including wild bees; Martins et al., 2017; Piano, Bona, & Isiaa, 2020; Piano, Souffreau, et al., 2020; Threlfall et al., 2015), other, often less conspicuous taxa have rarely been the focus of urban biodiversity research (e.g., millipedes, rove beetles, for an exception see Smith, Chapman, & Eggleton, 2006; Smith, Gaston, et al., 2006). Multi-taxon studies considering urban gardens are even rarer (but see, e.g., Braschler et al., 2020; Egerer et al., 2017; Tresch, Tresch, Frey, Le Bayon, Mäder, et al., 2019; Tresch, Frey, Le Bayon, Zanetta, et al., 2019).

Furthermore, studies on the effects of the urban matrix on functional diversity and the distribution of biological traits across several taxa are scarce (but see Melliger et al., 2017). This may impede our understanding of biodiversity changes along urbanization gradients as functional approaches incorporating species traits provide a mechanistic understanding of how species communities are shaped by the environment (Scheiner et al., 2017). In this context, analyzing relationships between species’ biological traits and environmental factors can help to identify which species can thrive in certain urban habitats (Wong et al., 2019). For example, urban habitats can act as environmental filters for species because some biological traits may be beneficial for inhabiting cities, while others may lead to the segregation of species (Buchholz & Egerer, 2020).

Building on this concept, we aimed to investigate how functional diversity is changing along urbanization gradients and which combination of biological traits is favored in highly urbanized areas. We examined these questions using a multi-taxon approach focusing on native grassland plants and several groups of ground-dwelling invertebrates (snails, slugs, spiders, millipedes, woodlice, ants, and rove beetles). As measures of functional diversity we considered functional dispersion (FDIs) and functional evenness (FEve). FDIs is a measure of functional richness, which considers the species’ relative abundances by estimating their dispersion in a multi-dimensional trait space (Laliberté & Legendre, 2010). FEve describes the evenness of abundance distribution in a functional trait space.

In our study, we tested the following four hypotheses:

1. Due to increased homogenization, which is often assumed for urban environments (Knop, 2016), we expected that functional diversity expressed as functional dispersion (Laliberté & Legendre, 2010) will decrease with increasing urbanization. Similarly, we expected that functional evenness, which measures the regularity of the distribution of species in functional space (Mason et al., 2005), will also be reduced, as only a few species sharing similar traits can cope with urban conditions. However, these species may then become very abundant.

2. Based on the assumption that in cities habitat generalist species are favored (Concepción et al., 2015, 2016), we expected an increase of habitat generalists in urban habitats.

3. The urban heat island causes higher temperatures in cities when compared with rural surroundings. Many thermophilic species also show preference for dry conditions (xerothermic species) (Horváth et al., 2012; Seifert, 2007). Following Menke et al. (2011) and Piano et al. (2017), who reported an enhanced proportion of thermophilic ant and carabid species respectively in urban sites, and Horváth et al. (2012), who reported higher numbers of xerophilic spider species in urban forest sites than in suburban or rural forest sites, we hypothesized that the abundance of dry adapted species will increase with increasing urbanization and that this will be consistent across all examined groups. This might be the case in spite of frequent irrigation in most private gardens along the urbanization gradient. We expected more xerothermophilic species in gardens within highly urbanized surroundings,

KEYWORDS
body size, domestic gardens, dry-adapted species, functional dispersion, functional evenness, urban ecology
although irrigation may partly dampen the urban heat island effect in some microhabitats of the gardens.

4. Within a taxonomic group, increased temperature may lead to a decline in body size (Gardner et al., 2011). Several studies provided evidence that this can result in small-sized species being more numerous in urban assemblages, as shown in bumblebees (Eggenberger et al., 2019), wild bees (Hamblin et al., 2018), spiders, carabids and weevils (Merckx et al., 2018) and in habitat specialists among carabids (Magura et al., 2020). Hence, we expected that for some of the examined groups of ground-dwelling invertebrates the proportion of small-sized species would increase with degree of urbanization.

2 | MATERIAL AND METHODS

2.1 | Garden selection

The study was conducted in the city of Basel, its suburbs and nearby villages in North-western Switzerland (47°34’N, 7°36’E). Total annual precipitation averages 842 mm and annual mean temperature is 10.5°C in the city (records from 1981 to 2010, www.meteoswiss.admin.ch). We selected 35 gardens from a pool of 65 candidates offered in response to public calls. The chosen gardens reflect a rural–urban gradient, encompassing wide ranges of garden sizes and management types. Further criteria for the garden choice were acceptance of the intended sampling methods by the garden owners and guaranteed daytime access to the gardens. All gardens had a grassland area of at least 4 m², allowing us to set up traps and hay baits, but they differed in the composition of other habitat types (see below).

2.2 | Plant and ground-dwelling invertebrate surveys

We recorded all native plant species (including woody species) occurring in the grassland by slowly walking in a zigzag line over the grassland area of a garden (hereafter native plant species richness in grassland). Plants were recorded when touching this line. Thus, sampling effort was proportional to the size of the grassland area in a garden. For total native plant species richness used as explanatory variable (see below), we complemented native plant species richness in grassland by recording the native plant species in the other habitat types by slowly walking along transect lines. These lines ran along the long axis of garden features (e.g., flower beds, vegetable plots, and hedges). Furthermore, we considered plants at intervals of 2 m along the transect line to measure the height of the vegetation, which we used to calculate structural diversity of the vegetation as a covariate for models on invertebrate functional diversity. Sampling effort for native plant species richness was thus proportional to the area with vegetation. Plant diversity was assessed in all gardens between July 24 and August 20, 2018.

We surveyed seven groups of ground-dwelling invertebrates (for details of sampling see Braschler et al., 2020). The groups cover a wide range of feeding strategies and included phylogenetically distant taxa: two groups of Gastropoda (snails and slugs), Araneae (spiders), Diplopoda (millipedes), Isopoda (woodlice), Formicidae (ants), and Staphylinidae (rove beetles) excluding the subfamily Pselaphinae. Some years ago, the Pselaphinae was considered a distinct family (Newton & Thayer, 1995), therefore our expert did not determine this group. Members of this group have a very different morphology from other rove beetles. Furthermore, very little information on habitat preferences was available for this subfamily. We used pitfall traps (plastic beakers, 5.8 cm diameter, containing salt-water as a preservative, that were buried flush with the ground) and hay bait traps (moist hay in coarse plastic netting placed in direct contact with the ground; Tuf et al., 2015) to sample all groups. Hay bait traps complement pitfall traps by providing a spot of humid conditions and thus attracting less-mobile, frequently in the leaf-litter or upper soil layers living invertebrates, as well as acting as a food resource attracting many detritivores and their predators (Tuf et al., 2015). We placed five pitfall traps and five hay bait traps in the grassland of each garden. Pitfall traps and hay bait traps were operated three times for 7 days each from early to late summer 2018. We employed additional techniques for four groups (snails and slugs (active search and sieving of soil samples), millipedes and ants (active search); see Braschler et al. (2020) for details). Invertebrate surveys were conducted between May 31 and October 18, 2018.

In the grassland area of the gardens, we recorded a total of 157 native plant species (Braschler et al., 2020). For spiders and woodlice, we only considered adult individuals and for ants, only workers. In the remaining invertebrate groups, we included all individuals that could be determined to the species level (99.4% of all individuals sampled). References for identification keys used and nomenclature followed can be found in Braschler et al. (2020). In all, we considered 1744 snail individuals (34 species), 1671 slug individuals (6 species), 1079 adult spider individuals (52 species), 6864 millipede individuals (21 species), 2582 adult woodlice individuals (9 species), and 1265 individuals of rove beetles (85 species). Furthermore, we found 28 species of ants (abundance data for this group was not considered because of the aggregated nature of ant colonies).

2.3 | Traits and habitat preferences

As measures of functional diversity, we considered functional dispersion (FDIs) and functional evenness (FEve). FDIs is a measure of functional richness, which considers the species’ relative abundances by estimating their dispersion in a multi-dimensional trait space (Laliberté & Legendre, 2010). FDIs has no upper limit, and high values correspond to large numbers of functionally different species. FEve describes the evenness of abundance distribution in a functional trait space. High FEve-values indicate a balanced niche occupancy, which occurs in species communities with balanced trait frequencies (Mason et al., 2005). To calculate FDIs and FEve, we used morphological and life-history
traits and microhabitat preference of the species recorded in the various gardens. The set of traits varied among groups (Table 1).

Urbanization alters environmental conditions (Gilbert, 1989; Sukopp & Wittig, 1998) and increases frequency, magnitude and type of disturbance (Niemelä, 2011). This may be to the detriment of some habitat specialists and thus increase the proportion of generalist species. We therefore divided species in those that are habitat generalists and those that are special to a particular habitat type (Table 1). Similarly, we divided species into those showing a preference for dry conditions and those showing either a preference for moist or wet conditions or without a preference (Table 1). In the various invertebrate groups, body size is measured in a different way (Table 1).

### 2.4 Local garden characteristics and landscape variables

We assessed 11 local garden characteristics: total garden area, area with vegetation, grassland area, percentage of grassland in a garden, area of shrubs and trees, percentage shrub and tree cover in a garden, habitat richness, length of non-permeable garden border, percentage length of non-permeable garden border, index of permeable garden border. In addition, for analyses of invertebrate functional diversity, we used the two characteristics total native plant species richness and structural diversity of the vegetation as explanatory variables. However, due to collinearity, we omitted several of the above variables from the data analyses, retaining only total native plant species richness, habitat richness, structural diversity and index of permeable border (Table 2).

We could not measure management intensity in a precise way for the following reasons. Firstly, some garden owners have changed in the recent past. The history of their gardens’ management (e.g., previous herbicide and pesticide applications) was thus not known. Secondly, within gardens the management is spatially heterogeneous (e.g., intensively managed flowerbeds and lawn with nearby wild-growing hedges and high-turf grassland). Our sampling considered all these differently managed habitat types. Thirdly, we received insufficient information on management decisions from the garden owners. For example, most did only provide vague statements on mowing frequency. Similarly, garden owners had quite different views on the meaning of the term “pesticide.” Snail baits were frequently not considered as a poison. Fourthly, turf height, which is sometimes used as a measure related to management intensity, was measured, but was also unreliable as we discovered that several garden owners did mow their lawns in anticipation of our visits. We therefore decided to use the number of native plant species as a proxy for naturalness of the gardens in models analyzing FDis and FEve of invertebrates (see below). We assume that the naturalness of a garden is inversely related to the management intensity of the garden.

As landscape variables we determined for each garden the percentage of sealed area and percentage of green area in the surroundings (both within a radius of 200 m) and the distance to the city center (Table 2). In data analyses, we considered only percentage of sealed area and distance to the city center because percentage of green area was correlated with both.

### 2.5 Statistical analyses

Statistical analyses were performed in R (ver. 3.3.3 and ver. 3.6.1; R Core Team, 2015) and were carried out separately for the different taxonomic groups with the 35 gardens as replicates. We used various traits, depending on the plants or the invertebrate group examined, to calculate FDis and FEve (Table 1) using the FD package in R (Laliberté & Legendre, 2010). For native grassland plants and for ants, we used presence/absence data, and for the remaining groups we used abundance data to calculate FDis and FEve. We applied generalized linear models (GLM) with Gaussian error distribution to examine potential effects of landscape variables, garden size (vegetated garden area) and various local garden characteristics on species richness of different taxonomic groups. The three main variables (distance to city center, percentage sealed area and vegetated garden area) were retained in all models, while a step-wise procedure was followed with the other four garden characteristics (total native plant richness, structural diversity of the vegetation, habitat type richness and index of permeable border) to obtain the minimal adequate models (Crawley, 2007). Three variables (total native plant richness, structural diversity of the vegetation and habitat type richness) were correlated with total garden area. Therefore, we used residuals of the relationships between the variable and total garden area for the GLM models. Distance to city center and percentage of sealed area were slightly correlated \( r = -0.46, p = .006 \). However, we retained both in the model because they represent different aspects of urbanization. The same GLM model was used to analyze urbanization-related effects on body size for invertebrate groups.

To examine whether the proportion of generalist species was affected by urbanization, we used GLM models with the three variables distance to city center, percentage of sealed area within 200 m and vegetated garden area (all log-transformed). The same model was used to examine the effect of urbanization on the proportion of species showing a preference for dry conditions. We applied a binominal error distribution and logit link function. To detect overdispersion, we compared the residual deviance with the residual degrees of freedom, and if the former were much higher (ratio >> 1), then we used quasi-binominal error distribution instead (Crawley, 2007).

### 3 RESULTS

#### 3.1 Functional dispersion

In the 35 gardens investigated, we found a negative effect of distance to city center on FDis of native grassland plants (Figure 1;
### TABLE 1
Plant and invertebrate traits used in the analyses. Measures of body length were used both for calculating FDis and FEve and as a dependent variable in analyses of the effects of urbanization on body size. Not all traits or habitat preferences were available for all taxonomic groups.

| Trait | Type           | Specification                                                                 | References |
|-------|----------------|-------------------------------------------------------------------------------|------------|
|       | (a) Traits used to calculate FDis and FEve<sup>a</sup> |                                               |            |
| **Plants** |                      |                                               |            |
| Life form | Categorical     | Macrophanerophyte; nanophanerophyte; chamaephyte; hemicryptophyte; geophyte; therophyte | 1          |
| Reproduction type | Categorical | Sexual; mixed; self-fertilization | 1          |
| Ecological strategy | Categorical | C; CR; CS; CSR; S; SR (categories after Grime, 1979) | 1          |
| Pollination syndrome | Categorical | Insects; wind | 1          |
| Seed dispersal type | Categorical | Zoochor; anemochor; hemerochor; autochor; hydrochor | 2          |
| Seed mass | Continuous | Mean of seed mass (mg) | 1          |
| **Snails** |                      |                                               |            |
| Shell size | Continuous     | The longer of shell height or shell breadth (mm) | 3          |
| Age at sexual maturity | Ordinal | <1 year; 1 year; >1 year | 4–6         |
| Longevity | Ordinal         | <1 year; 1–2 years; >2 years | 5, 6        |
| Shell shape (snails) | Categorical | Depressed; globose/conical; oblong | 3          |
| **Spiders** |                      |                                               |            |
| Body size | Continuous | Mean male body length (mm) | 7          |
| Hunting mode | Categorical | Ambush; ground hunter; orb web; sensing web; sheet web; space web; specialist; other | 8          |
| Shading width | Continuous | Measure for the niche width | 9          |
| Humidity width | Continuous | Measure for the niche width | 9          |
| **Millipedes** |                      |                                               |            |
| Body length | Continuous | Mean of male and female body length of first adult stage (mm) | 10–15      |
| Breadth | Continuous | Maximum body breadth at mid length of individual (mm) | 10–15      |
| Eye morphology | Continuous | Number of ocelli | 10–13, 16  |
| Feeding guild | Categorical | Detritivore; facultative scavenger; algivore | 10–12, 17  |
| **Ants** |                      |                                               |            |
| Body size | Continuous | Maximum total length of workers, including major workers in species where these forage (mm) | 18–21      |
| Main food | Categorical | Carbohydrates; animal matter; carbohydrates & animal matter; grains | 18–20      |
| Main nest stratum | Categorical | Wood & litter; soil & crevices; both | 18–20      |
| Colony founding mode | Categorical | Independent; social parasite | 18–20      |
| Number of queens | Categorical | Monogynous; polygynous (species where only some nests have more than one queen are considered polygynous) | 18–20      |
| **Rove beetles** |                      |                                               |            |
| Body size | Continuous | Mean of mean body length of males and females (mm) | 10, 22–30  |
| Microhabitat preference | Categorical | No specific microhabitat preference; saprophilous; coprophilous; thermophilous but no further preference; hygrophilous but no further preference; phytodetricolous | 31          |
| **(b) Measures for habitat preferences** |                      |                                               |            |
| Native grassland plants |                      |                                               |            |
| Habitat preference | Binomial | Habitat specialist (grassland); habitat generalist (occurring in two or more habitat types, e.g., open land, forest or agricultural land) | 32          |
| Preference for dry conditions | Binomial | Preference for wet or moist habitats (indicator value above 3); preference for dry habitats (indicator value up to 3) | 33          |

(Continues)
Percentage of sealed area negatively influenced FDis of ants (Figure 1; Table 3; Figure S1). No significant effects of distance to city center or percentage of sealed area on FDis were found for the other invertebrate groups. Habitat type richness was the only local garden characteristics to affect FDis, which was increasing in snails with increasing habitat type richness (Figure 1; Table 3; Figure S1).

### 3.2 Functional evenness

Distance to city center positively affected FEve of ants (Figure 2; Table 3; Figure S2). In rove beetles, percentage of sealed area within 200 m negatively affected FEve (Figure 2; Table 3; Figure S2). Vegetated garden area positively influenced the FEve of ants (Figure 2; Table 3; Figure S2). Other local garden characteristics did not significantly affect FEve in any group (Figure 2; Table 3; Figure S2).

### 3.3 Habitat preference

The proportion of generalists among rove beetles declined with increasing percentage of sealed area in the surroundings of the gardens in contrast to our hypothesis (Table S1). In the other groups, the proportion of generalists was not affected by either distance to the city center or percentage of sealed area (Table S1).

Also contrary to our expectation, the proportion of rove beetle species preferring dry conditions increased with increasing distance to the city center (Table S1). For the other groups examined neither distance to the city center nor percentage of sealed area influenced the proportion of species preferring dry conditions (Table S1).

### 3.4 Body size

Distance to city center affected body size in all invertebrate groups except snails and woodlice (Figure 3; Table S2; Figure S3). Body size increased with distance to city center for slugs, spiders, millipedes, ants, and rove beetles. In millipedes, body size also increased with index of permeable garden border (Figure 3; Table S2; Figure S3).

### 4 DISCUSSION

Urbanization can affect functional diversity of organisms and the distribution of biological traits (Williams et al., 2009). We used a
A multi-taxon approach, including plants and seven groups of ground-dwelling invertebrates, which revealed that the observed effects of urbanization—either at landscape or at habitat level—can be multifarious and are often ambivalent. In some groups, effects of urbanization on certain aspects of functional diversity may even be positive, while other groups do not show a response to the increasing degree of urbanization at all.

We found no evidence for an overall homogenization in terms of a reduced FDis across the investigated taxa, except for native grassland plants and ants. The responses of FEve were also sporadic and only significant for ants and rove beetles. The first hypothesis is therefore not supported by our findings. The second hypothesis assuming that the proportion of generalists is higher in urban than in rural gardens, could not be confirmed either as the proportion of generalists did not change over the urbanization gradient in all groups except in rove beetles. The latter even showed the opposite trend with a decreased proportion of generalists in urban gardens.

Our third hypothesis assumed that the proportion of xerophilic species is increased in urban gardens due to the urban heat island effect. Our results could, however, not confirm this hypothesis. Contrary to our expectations, only xerophilic rove beetle species responded, but in the opposite direction from our expectation making up a lower proportion of the assemblage in urban gardens. Finally, we found strong evidence confirming our fourth hypothesis that body size decreases with increasing urbanization. In most invertebrate groups examined, the average size of the species decreased with decreasing distance to the city center.

### 4.1 Functional dispersion

The environmental conditions in urban areas can act as filters, which have the potential to reduce plant and invertebrate species richness, change species composition and thus functional diversity (Buchholz et al., 2020; Williams et al., 2009). This does not necessarily lead to a decrease in biodiversity. Urban areas can also harbor high species richness, sometimes higher than in the rural surroundings (Kühn et al., 2004; Wania et al., 2006).

In plants, species composition has been reported to be more homogeneous in urban habitats than rural ones (Zeeman et al., 2017), which may result in a decreased functional diversity. We therefore hypothesized that FDis decreases along the rural–urban gradient. However, we recorded higher FDis in plants in highly urbanized areas than in rural ones. This contrasts the findings from other habitat types. Knapp et al. (2012) and Melliger et al. (2018) demonstrated that FDis in plants decreased with increasing degree of urbanization in grassland and forest. Our unexpected result could be explained by species composition not being affected by distance to the city center. However, the higher FDis of native grassland plant species at short distance to the city center may be due to the observed
higher similarity among plant assemblages of the grasslands in the city center than in the rural surroundings (H.-P. Rusterholz, unpublished data). This was mainly caused by gardens in the city center having slightly higher proportions of annual plants in their grasslands than gardens in the rural surroundings (H.-P. Rusterholz, unpublished data). Furthermore, grasslands in gardens are in most cases not naturally grown plant communities (Gilbert, 1989). Seed mixtures and management actions influence plant species composition of grasslands in gardens and thus FDis (Knapp et al., 2012).

Ants were the only group that responded as expected; their FDis decreased with increasing urbanization, expressed as percentage of sealed area in the surroundings. We do not have any explanation for this result, and did not find any literature on ant FDis changes along urbanization gradients. However, species composition of ants is generally considered as a good indicator of altered environmental factors and of land-use change (Andersen, 2000; Kaspari & Majer, 2000). Therefore, functional groups of ants are frequently considered in assessments of land-use change (Andersen, 2000). Thus, one may expect that responses to land-use change accompanying urbanization may be captured by ant FDis, as shown in our study.

For the other invertebrate groups examined we found no effect of urbanization on functional dispersion. Tóth and Hornung (2020) reported decreased FDis with increasing urbanization for millipedes in forests and other types of woodland (e.g., parks) in Budapest, Hungary. The decrease in this other habitat type was mainly due to a reduced number of forest specialist species in highly urbanized areas. This is consistent with the finding of Bogyó et al. (2015) for millipedes in urban forests. Similarly, Nagy et al. (2018) reported that functional diversity (measured as functional richness) in woodlice decreased along the urbanization gradient in forests. However, gardens

| Studied group  | Distance to city center | Percentage of sealed area | Vegetated garden area | Native plant richness | Habitat richness | Structural diversity | Index of permeable garden border |
|---------------|------------------------|---------------------------|-----------------------|----------------------|------------------|---------------------|--------------------------------|
| Native plants | ns                     | ns                        | ns                    | –                    | –                | –                   | –                              |
| Snails        | ns                     | ns                        | ns                    | –                    | ns               | –                   | ns                             |
| Spiders       | ns                     | ns                        | ns                    | –                    | ns               | –                   | –                              |
| Millipedes    | ns                     | ns                        | ns                    | –                    | –                | –                   | –                              |
| Ants          | ns                     | ns                        | ns                    | –                    | –                | –                   | ns                             |
| Rove beetles  | ns                     | ns                        | ns                    | –                    | –                | –                   | –                              |
### TABLE 3  Summary of GLM analyses examining the effects of two measures of urbanization (distance to city center and percentage of sealed area in the surroundings) and garden size (vegetated garden area), total native plant species richness, habitat type richness, structural diversity of the vegetation, index of permeable border on functional dispersion (FDis) and functional evenness (FEve) of various organism groups

| Organism group | Variable                        | FDis df | F   | p    | FEve df | F   | p    |
|----------------|---------------------------------|---------|-----|------|--------|-----|------|
| Native grassland plants | Distance to city centera | 1.33 | 7.60 | .010 | 1.33 | 2.26 | .14 |
|                     | Percentage sealed areaa | 1.32 | .05  | .82  | 1.32 | .16  | .69 |
|                     | Vegetated garden areaa | 1.31 | .11  | .74  | 1.31 | 3.64 | .066 |
|                     | Habitat type richnessb | -      | -    | -    | -      | -    | -    |
|                     | Index of permeable border | -      | -    | -    | -      | -    | -    |
| Snails | Distance to city centera | 1.33 | 2.94 | .097 | 1.32 | .01  | .94 |
|                     | Percentage sealed areaa | 1.32 | .01  | .91  | 1.31 | .39  | .54 |
|                     | Vegetated garden areaa | 1.31 | .26  | .61  | 1.30 | .49  | .49 |
|                     | Total native plant species richnessb | - | -  | -    | - | -    | -    |
|                     | Habitat type richnessb | 1.30 | 5.68 | .024 | 1.29 | 1.05 | .31 |
|                     | Structural diversity of the vegetationb | - | -  | -    | - | -    | -    |
|                     | Index of permeable border | 1.29 | 1.90 | .18  | - | -    | -    |
| Spiders | Distance to city centera | 1.33 | .04  | .85  | 1.33 | .43  | .52 |
|                     | Percentage sealed areaa | 1.32 | .84  | .37  | 1.32 | 1.39 | .25 |
|                     | Vegetated garden areaa | 1.31 | 2.57 | .12  | 1.31 | <.01 | .99 |
|                     | Total native plant species richnessb | - | -  | -    | - | -    | -    |
|                     | Habitat type richnessb | 1.30 | 3.84 | .059 | 1.29 | 3.73 | .063 |
|                     | Structural diversity of the vegetationb | - | -  | -    | - | -    | -    |
|                     | Index of permeable border | - | - | -    | - | -    | -    |
| Millipedes | Distance to city centera | 1.33 | .14  | .71  | 1.30 | .89  | .35 |
|                     | Percentage sealed areaa | 1.32 | .01  | .92  | 1.29 | 4.07 | .053 |
|                     | Vegetated garden areaa | 1.31 | <.01 | .98  | 1.28 | .96  | .33 |
|                     | Total native plant species richnessb | - | -  | -    | - | -    | -    |
|                     | Habitat type richnessb | - | -  | -    | - | -    | -    |
|                     | Structural diversity of the vegetationb | - | -  | -    | - | -    | -    |
|                     | Index of permeable border | - | - | -    | - | -    | -    |
| Ants | Distance to city centera | 1.33 | 1.16 | .29  | 1.33 | 16.67 | <.001 |
|                     | Percentage sealed areaa | 1.32 | 4.41 | .044 | 1.32 | 1.58 | .22 |
|                     | Vegetated garden areaa | 1.31 | .39  | .54  | 1.31 | 22.66 | <.001 |
|                     | Total native plant species richnessb | - | -  | -    | - | -    | -    |
|                     | Habitat type richnessb | - | - | -    | - | -    | -    |
|                     | Structural diversity of the vegetationb | - | -  | -    | - | -    | -    |
|                     | Index of permeable border | 1.30 | 2.75 | .11  | - | -    | -    |
| Rove beetles | Distance to city centera | 1.33 | .29  | .59  | 1.33 | .08  | .78 |
|                     | Percentage sealed areaa | 1.32 | .09  | .76  | 1.32 | 6.75 | .014 |
|                     | Vegetated garden areaa | 1.31 | .32  | .58  | 1.31 | .12  | .73 |
|                     | Total native plant species richnessb | - | -  | -    | - | -    | -    |
|                     | Habitat type richnessb | - | -  | -    | - | -    | -    |
|                     | Structural diversity of the vegetationb | - | -  | -    | - | -    | -    |
|                     | Index of permeable border | - | - | -    | - | -    | -    |

Note: Significant p-values (<.05) are in bold. FDis and FEve are based on abundance data except for native grassland plants and ants, for which presence/absence data was used. "-" variable was excluded from the model by step-wise reduction.

aLog-transformed.

bDue to correlation with total garden size, residuals of the regression of the variable on total garden size were used for analyses.
provide other environmental conditions than urban forests. In contrast to forests, gardens may, independent of their location along the rural–urban gradient, provide relatively similar conditions for ground-dwelling invertebrates due to similar management (see below).

### 4.2 Functional evenness

Decreasing FEve with increasing urbanization as found for rove beetles may indicate temporally variable habitat conditions due to disturbance or environmental stressors, which cause an unbalanced niche occupancy (Schleuter et al., 2010). This can be a result of the dominance of few species that may have adapted to urban environments, while specialized species may disappear as found for rove beetles by Magura et al. (2013).

Similar to the findings on FDis, FEve of ants changed with urbanization in our study, being lower close to the city center. Additionally, FEve of ants increased with increasing vegetated garden area. Increasing the size of a garden may thus counteract some of the negative effects on FEve, as ant colonies with their durable nests are potentially more severely affected by local garden characteristics than more mobile species. Indeed, ant species richness increased in gardens with larger vegetated area (Braschler et al., 2020).

For the other ground-dwelling invertebrate groups examined we recorded no effects of urbanization on FEve. This matches the few other studies, which for the most part found no changes in invertebrate FEve along urbanization gradients (e.g., Banaszak-Cibicka and Dylewski (2021) for bees and Correa et al. (2021) for dung beetles).

| Studied group          | Distance to city center | Percentage of sealed area | Vegetated garden area | Native plant richness | Habitat richness | Structural diversity | Index of permeable garden border |
|------------------------|-------------------------|---------------------------|-----------------------|-----------------------|------------------|----------------------|---------------------------------|
| Native plants          | ns                      | ns                        | ns                    | ns                    | –                | –                    | –                               |
| Snails                 | ns                      | ns                        | ns                    | –                     | ns               | –                    | –                               |
| Spiders                | ns                      | ns                        | ns                    | ns                    | ns               | –                    | –                               |
| Millipedes             | ns                      | ns                        | ns                    | –                     | –                | –                    | –                               |
| Ants                   | –                       | –                         | –                     | –                     | –                | –                    | –                               |
| Rove beetles           | ns                      | –                         | –                     | ns                    | –                | –                    | ns                               |

**FIGURE 2**  Effects of urbanization, garden size and local garden characteristics, habitat type richness, structural diversity of the vegetation, and index of permeable border, on FEve of native grassland plants and five groups of invertebrates. For detailed explanations, see caption to Figure 1.
4.3 | Habitat preference

Contrary to our expectations, the proportion of habitat generalists increased in rove beetles in rural gardens, while no change in the proportion of generalists was found in the other groups examined. However, Melliger et al. (2018) reported an increase in the share of habitat generalists with increasing percentage of sealed area in ants and spiders in urban forests in Basel. Similarly, Magura et al. (2013) found that the proportion of forest-associated rove beetles was significantly lower in urban forest, when compared with suburban and rural forest. Forests, which are less intensively managed than private gardens, may show steeper rural–urban gradients for abiotic environmental factors and/or levels of disturbance. Unlike the more frequently studied forests or grasslands, urban gardens may harbor less species with strong habitat bindings. Urban gardens consist of a mosaic of patches of different habitat types. These patches may be too small to harbor certain habitat specialist species. Furthermore, as a result of regular management activity, variation in some environmental conditions may be dampened in gardens independent of the location of the garden. While we could not directly study the effects of management intensity on functional diversity in the examined gardens, because of challenges including heterogeneous management within gardens and imprecise information provided by owners, our observations gave us no reason to expect garden management to change along the rural–urban gradient. The similar environmental conditions in gardens in both rural and urban settings might be more powerful in filtering species composition than other factors such as the composition of the surrounding matrix, which change along the rural–urban gradient (Braschler et al., 2020). For example, repeated watering excludes pronounced periods of drought. This may also explain why—contrary to our expectations—we did not find an increased proportion of xerophilous species in gardens in highly urbanized areas, despite the existing urban heat island effect. Menke et al. (2011) suggested that urban areas may serve as habitat and corridors for dry-adapted and heat-tolerant species, and provided some evidence for this in ants along a rural–urban gradient including different habitat types in Raleigh, North Carolina, USA, which is similar in population to the greater Basel area.

In plants, factors other than management including watering may lead to homogenization of species assemblages in gardens along the rural–urban gradient. Importantly, grasslands in gardens typically originate from commercial seed mixtures (Gilbert, 1989). Grassland plant species assemblages in gardens are also strongly affected by mowing regimes and trampling (Bertoncini et al., 2012). Furthermore, grassland plant communities are commonly characterized by a low proportion of generalist species (Ellenberg, 1986). A combination of these factors may explain why we did not find a change in the proportion of generalist plant species in the grasslands of the gardens. Similarly, the percentage of generalist species in the other ground-dwelling invertebrate groups did not change along the rural–urban gradient.

4.4 | Body size

We found that urban gardens compared with rural ones harbored more small-sized species in five out of the seven groups of invertebrates examined. One explanation for this pattern is the well-known heat island effect, which is also well documented in the city of Basel (Wicki et al., 2018). Increased temperatures may influence the body size distributions both within species and at the species assemblage level with larger species becoming rarer at higher temperatures (Gardner et al., 2011; Verberk et al., 2021). In general, this could be related to the temperature–size rule (Atkinson, 1994), for which an ecological explanation could be a larger requirement for resources (food and oxygen) under warmer conditions preventing animals from growing larger (Verberk et al., 2021). An alternative explanation is that animals consist of smaller cells in warm environments (Verberk et al., 2021). The trend towards smaller body size will continue due to global warming and may lead to significant changes in the diversity and species composition of animals in cities. This in turn will affect body size-dependent ecosystem services in cities, especially as smaller species may not be as effective predators, decomposers, seed dispersers or pollinators as larger species are.

The decrease in millipede body size with decreasing distance to the city center mirrors the findings from other studies (Bogyó et al., 2015; Tóth & Hornung, 2020). Bogyó et al. (2015) hypothesized that the decreased size in millipedes in urban areas is a result of lower food quality, while Tóth and Hornung (2020) suggested that it may be related to a combination of low food quality, deteriorated soils, reduced soil moisture, and increased soil contamination. For example, reduced soil moisture is likely to affect smaller species more because of a disadvantageous surface area to body volume ratio and lower capacity of storing water. Indeed, smaller millipede species usually inhabit colder and less arid places (Enghoff, 1992). In our study, most of the small millipede species recorded live buried in the soil. In this way, they are less influenced by above-ground management activities, but may be strongly negatively affected by soil degradation and soil contaminants.

Body size is frequently related to the species’ dispersal mode and their ability to disperse (Biedermann, 2002; Jenkins et al., 2007; Kuussaari et al., 2014). In some groups of ground-dwelling invertebrates, smaller species are mainly passively dispersed among habitats including gardens (e.g., slugs or millipedes transported in soil, attached to garden plants or green waste; Dörge, 1999; Stoev et al., 2010). In contrast, larger species of the same groups have to disperse mainly actively through the urban matrix. Thus, larger species may be less likely to reach isolated gardens in the city center. Open gardens with permeable borders enhance colonization by actively dispersing larger species, as shown by the correlation between millipede body size and index of permeable garden border in our study (Figure 3).

Body size did not decrease with increasing urbanization in only two groups: snails and woodlice. In contrast to our observation, Ooms et al. (2020) reported that woodlice body size increased with urbanization in Amsterdam and explained their findings by the
advantages of larger body size to reduce water loss under dry and warm conditions (cf. Csonka et al., 2018; Merckx et al., 2018).

5 | CONCLUSIONS

Our study showed different responses to urbanization by the various groups examined. This confirms that environmental change affects taxonomic groups differently and highlights the need for using multi-taxon approaches to avoid basing conservation decisions and land-use management on the particular response by just a single group. We studied small ground-dwelling, less conspicuous species, which, however, are of key importance for certain ecosystem functions. Generally, ground-dwelling invertebrates are little studied and thus not much is known about their population trends and vulnerabilities to environmental change. This precludes management tailored to the needs of such taxonomic groups.

Independent of their location along their rural-urban gradient, domestic gardens are more intensively managed than semi-natural areas (meadows, forest, and hedges). Therefore, species inhabiting gardens may have traits suitable for the special environmental conditions in this habitat, which in turn may not vary considerably along the rural-urban gradient. The variation in environmental conditions in gardens along the urbanization gradient may thus be lower than those in frequently studied habitats such as urban forests or grasslands. This may also explain why we found few effects of local

| Studied group | Distance to city center | Percentage of sealed area | Vegetated garden area | Native plant richness | Habitat richness | Structural diversity | Index of permeable garden border |
|---------------|-------------------------|---------------------------|----------------------|----------------------|------------------|---------------------|---------------------------------|
| Snails        | ns                      | ns                        | ns                   | –                    | –                | ns                  | ns                              |
| Slugs         |                         |                           |                      |                      |                  |                     |                                 |
| Spiders       |                         |                           |                      |                      |                  |                     |                                 |
| Millipedes    |                         |                           |                      |                      |                  |                     |                                 |
| Woodlice      | ns                      | ns                        | ns                   | ns                   | –                | ns                  | ns                              |
| Ants          |                         |                           |                      |                      |                  |                     |                                 |
| Rove beetles  |                         |                           |                      |                      |                  |                     |                                 |

**FIGURE 3** Effects of urbanization, garden size and local garden characteristics, habitat type richness, structural diversity of the vegetation, and index of permeable border, on body size of seven groups of invertebrates. For detailed explanations, see caption to Figure 1.
garden characteristics on the functional diversity of the groups examined. Typical traits of species inhabiting gardens may also be generally helpful in coping with the intense anthropogenic use of green space characteristic of urban areas.

Our findings demonstrate that gardens in highly urbanized areas have similar functional diversity for several organism groups as gardens in rural surroundings. This should motivate urban garden owners to promote native biodiversity in the future.

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CONFLICT OF INTEREST
The authors declare no conflict of interest.

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DATA AVAILABILITY STATEMENT
The abundance data per species and garden (presence/absence for plants and ants), the trait values used for each species and the sources used, as well as the landscape variables and garden characteristics for each garden are available from the Dryad Digital Repository (https://doi:10.5061/dryad.fqz612j).

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REFERENCES
Adam, L. (2010). Remarks on some European Aleocharinae, with description of a new Rhopalates species from Croatia (Coleoptera: Staphylinidae). Travaux Du Museum National D'Histoire Naturelle "Grigore Antipa", 53, 191–215. https://doi.org/10.2478/v10199-010-0015-6
Adams, B. J., Li, E., Bahli, C. A., Meineke, E. K., McGlynn, T. P., & Brown, B. V. (2020). Local- and landscape-scale variables shape insect diversity in an urban biodiversity hot spot. Ecological Applications, 30, e02089. https://doi.org/10.1002/eap.2089
Andersen, A. N. (2000). A global ecology of rainforest ants: Functional groups in relation to environmental stress and disturbance. In D. Agosti, J. D. Majer, L. E. Alonso, & T. R. Schultz (Eds.), Ants: Standard methods for measuring and monitoring biodiversity (pp. 25–34). Smithsonian Institution Press.
Anderson, R. (1996). Annamastigona pulchellum Silvestri (Chordeumatida: Anthroleucosomatidae) as an addition to the fauna of northern Europe, recorded from Ireland. Bulletin of the British Myriapod Group, 12, 7–11.
Atkinson, D. (1994). Temperature and organism size — A biological law for ectotherms? Advances in Ecological Research, 25, 1–58.
Banaszak-Cibicka, W., & Dylewski, Ł. (2021). Species and functional diversity — A better understanding of the impact of urbanization on bee communities. Science of the Total Environment, 774, 145729. https://doi.org/10.1016/j.scitotenv.2021.145729
Baur, B. (1994). Parental care in terrestrial gastropods. Experientia, 50, 5–14. https://doi.org/10.1007/BF01992042
Bengtsson, J., & Baur, B. (1993). Do pioneers have r-selected traits? Life-history patterns among colonizing terrestrial gastropods. Oecologia, 94, 17–22. https://doi.org/10.1007/BF00317295
Beninde, J., Veith, M., & Hochkirch, A. (2015). Biodiversity in cities needs space: a meta-analysis of factors determining intra-urban biodiversity variation. Ecology Letters, 18, 581–592. https://doi.org/10.1111/ele.12427
Bertoncini, A. P., Machon, N., Pavoine, S., & Muratet, A. (2012). Local garden practices shape urban lawn floristic communities. Landscape and Urban Planning, 105, 53–61. https://doi.org/10.1016/j.landurbplan.2011.11.017
Biedermann, R. (2002). Mating success in the spittlebug Cercopis sanguinolenta (Scopoli, 1763) (Homoptera, Cercopidae): The role of body size and mobility. Journal of Ethology, 20, 13–18. https://doi.org/10.1007/s10164-002-0048-9
Blower, J. G. (1995). Millipedes: keys and notes for the identification of the species. In D. M. Kermack & R. S. K. Barnes (Eds.), Synopses of the British Fauna, New Series, Vol. 35 (pp. 242). Linnean Society.
Bogó, D., Magura, T., Simon, E., & Tóthmérész, B. (2015). Millipede (Diplopoda) assemblages alter drastically by urbanisation. Landscape and Urban Planning, 133, 118–126. https://doi.org/10.1016/j.landurbplan.2014.09.014
Braschler, B., Gilgado, J. D., Zwahlen, V., Rusterholz, H.-P., Buchholz, S., & Baur, B. (2020). Ground-dwelling invertebrate diversity in domestic
forest in Hungary. *Ecological Research*, 27, 1069–1077. https://doi.org/10.1007/s11284-012-0988-7

Ives, C. D., Lentini, P. E., Threlfall, C. G., Ikin, K., Shanahan, D. F., Garrard, G. E., Bekessy, S. A., Fuller, R. A., Mumaw, L., Rayner, L., Rowe, R., Valentine, L. E., & Kendall, D. (2016). Cities are hotspots for threatened species. *Global Ecology and Biogeography*, 25, 117–126. https://doi.org/10.1111/geb.12404

Jenkins, D. G., Bresciani, C. R., Duxbury, C. V., Elliott, J. A., Evans, J. A., Grablow, K. R., Hillegass, M., Lyon, B. N., Metzger, G. A., Olandese, M. L., Pepe, D., Silvers, G. A., Suresch, H. N., Thompson, T. N., Trexler, C. M., Williams, G. E., Williams, N. C., & Williams, S. E. (2007). Does size matter for dispersal distance? *Global Ecology and Biogeography*, 16, 415–425. https://doi.org/10.1111/j.1466-8238.2007.00312.x

Kaspari, M., & Mayer, J. D. (2000). Using ants to monitor environmental change. In D. Agosti, J. D. Mayer, L. E. Alonso, & T. R. Schultz (Eds.), *Ants: Standard methods for measuring and monitoring biodiversity* (pp. 25–34). Smithsonian Institution Press.

Kerney, M. P., Cameron, R. A. D., & Jungbluth, J. H. (1983). Die Landschnecken Nord- und Mitteleuropas (pp. 384). Paul Parey.

Kime, R. D., & Enghoff, H. (2011). Atlas of European Millipedes (Class Diplopoda), vol. 1: *Orders Polyxenida, Glomerida, Platyselida, Siphonocryptidae, Polysoniida, Callipodida, Polydesmidia* (pp. 282). Pensoft Publishers.

Kime, R. D., & Enghoff, H. (2017). Atlas of European millipedes, vol. 2: order Julida (class Diplopoda). *European Journal of Taxonomy*, 346, 1–299. https://doi.org/10.5852/ejt.2017.346

Klotz, S., Künn, I., & Durka, W. (2002). BIOFLOR – Eine Datenbank mit biologisch-ökologischen Merkmalen zur Flora von Deutschland. Landwirtschaftsverlag. https://www2.ufz.de/biolflor

Knapp, S., Dinsmore, L., Fissore, C., Hobbie, S. E., Jakobsdottir, I., Kattge, J., King, J. Y., Klotz, S., McFadden, J. P., & Cavender-Bares, J. (2012). Biotic homogenization of three insect groups due to gaps in knowledge of the impact of urban growth on biodiversity. *Nature Sustainability*, 3, 16–24. https://doi.org/10.1038/s41389-017-0043-6

Knell, R. L., Braschler, B., Rusterholz, H.-P., & Baur, B. (2018). Diverse effects of urbanisation and forest size on species richness and functional diversity of plants, and ground surface-active ants and spiders. *PLoS One*, 13, e0199245. https://doi.org/10.1371/journal.pone.0199245

Kühn, I., Brandl, R., & Klotz, S. (2004). The flora of German cities is naturally species rich. *Evolutionary Ecology Research*, 6, 749–764.

Kutter, H. (1977). Hymenoptera, Formicidae. *Insecta Helvetica: Fauna*, 6, 1–298.

Kuussaari, M., Saarinen, M., Korpela, E.-L., Pöyry, J., & Hyvönen, T. (2014). Higher mobility of butterflies than mites connected to habitat suitability and body size in a release experiment. *Ecology and Evolution*, 4, 3800–3811. https://doi.org/10.1002/ece31187

Laiberté, É., & Legendre, P. (2010). A distance-based framework for measuring functional diversity from multiple traits. *Ecology*, 91, 299–305. https://doi.org/10.1890/08-2244.1

Lee, S.-G., & Ahn, K.-J. (2017). A taxonomic review of the Korean species of the subgenus *Datamica* Mulsant & Rey of the genus *Atheta* Thomson, with description of a new species (*Coleoptera, Staphylinidae, Aleocharinae*). *Zootaxa*, 4268, 508–522. https://doi.org/10.11646/zootaxa.4268.4.3

Legrand, J. J. (1948). Les isopodes terrestres des environs de Paris. II. Liste des espèces observées. *Remarques écologiques. Bulletin Du Muséum National D’histoire Naturelle (Paris)*, 20, 154–158.

Lin, B. B., & Egerer, M. H. (2020). Global social and environmental change drives the management and delivery of ecosystem services from urban gardens: A case study from Central Coast. *California. Global Environmental Change*, 60, 102006. https://doi.org/10.1016/j.gloenvcha.2019.102006

Luka, H. (2004). Ökologische Bewertung von Landschaftselementen mit Arthropoden. *Opuscula Biogeographica Basileaenia*, 4, 1–253.
Wicki, A., Parlow, E., & Feigenwinter, C. (2018). Evaluation and modeling of urban heat island intensity in Basel. Switzerland. Climate, 6, 55. https://doi.org/10.3390/cli6030055

Williams, N. S. G., Schwartz, M. W., Vesk, P. A., McCarthy, M. A., Hahs, A. K., Clemants, S. E., Corlett, R. T., Duncan, R. P., Norton, B. A., Thompson, K., & McDonnell, M. J. (2009). A conceptual framework for predicting the effects of urban environments on floras. Journal of Ecology, 97, 4–9. https://doi.org/10.1111/j.1365-2745.2008.01460.x

Wong, M. K. L., Guénard, B., & Lewis, O. T. (2019). Trait-based ecology of terrestrial arthropods. Biological Reviews, 94, 999–1022.

Youngsteadt, E., Ernst, A. F., Dunn, R. R., & Frank, S. D. (2017). Responses of arthropod populations to warming depend on latitude: evidence from urban heat islands. Global Change Biology, 23, 1436–1447. https://doi.org/10.1111/gcb.13550

Zeeman, B. J., McDonnell, M. J., Kendal, D., & Morgan, J. W. (2017). Biotic homogenization in an increasingly urbanized temperate grassland ecosystem. Journal of Vegetation Science, 28, 550–561. https://doi.org/10.1111/jvs.12507

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