Local and landscape features constrain the trait and taxonomic diversity of urban bees

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Abstract
Context There is concern that urbanization threatens bees, a diverse group of economic importance. The impact of urbanization on bees is likely mediated by their phenotypic traits.

Objectives We examine how urban cover and resource availability at local and landscape scales influences both species taxonomic and functional diversity in bees.

Methods We used a combination of aerial netting and pan traps across six sampling periods to collect wild bees in 18 urban gardens spanning more than 125 km of the California central coast. We identified 3537 specimens to genus and, when possible, to species to obtain species richness and abundance at each site. For each species we measured a suite of bee traits, including body size, sociality, nesting location, nesting behavior, pollen-carrying structure, parasitism, and lecyn.

Results We found that increased garden size was positively associated with bee species richness and abundance. Somewhat counterintuitively, we found that urban cover surrounding gardens (2 km) was positively associated with bee species richness. Urban cover was also associated with the prevalence of certain bee traits, such as bees that excavate nests over those who rent, and bees with non-corbiculate structures. We suggest that urban habitats such as gardens can host a high number of bee species, but urbanization selects for species with specific traits.

Conclusions These findings illustrate that local and landscape features both influence bee abundance, species richness, and the frequency of specific traits. We highlight the importance of trait-based approaches for assessing biodiversity in urban landscapes, and suggest conceptualizing urbanization as a process of habitat change rather than habitat loss.

Keywords Urban agriculture · Biodiversity · Functional traits · Landscape ecology · Bees
Introduction

Functional diversity can be used to infer how environmental stressors influence complex ecosystem processes (Laakso and Setälä 1999; Norberg et al. 2001; Cadotte et al. 2011). Functional diversity refers to traits, the components of an organism’s phenotype that are important for an ecosystem-level process (Tilman 2001; Petchey and Gaston 2006). Trait-based metrics of community functional structure include, for example, feeding types, habitat preferences, body size, and dispersal ability. Trait-based metrics may be more important than species taxonomic identity for predicting ecological processes (Petchey and Gaston 2006). For example, in insect pollinator guilds, traits related to sociality and nesting behaviors may be more critical than species identity for predicting responses to environmental disturbance (Winfree et al. 2009; Williams et al. 2010).

The composition of functional traits in a community may mediate the impact of habitat loss on species persistence (Bommarco et al. 2010; Maire et al. 2012; Tscharntke et al. 2012). Local features, such as resource availability, and landscape features, such as natural habitat availability or regional climate, may filter which species are present because they possess traits suitable for a given habitat (Keddy 1992; Diaz et al. 1998) or because species exhibit variation in life history strategies and tolerance to stress (e.g., Waser et al. 1996; Hulot et al. 2000). This is seen across multiple taxa, including bacteria (e.g., Langenheder and Prosser 2008), algae (e.g., Fowler-Walker et al. 2006), plants (e.g., Lavorel et al. 2007), and insects (e.g. Philpott et al. 2019) and birds (e.g., Tscharntke et al. 2008; Clough et al. 2009). For example, larger-bodied species may be more likely to decline than small-bodied species (Bartomeus et al. 2013) or more tolerant to the loss of resources than small-bodied species due to dispersal ability (Bommarco et al. 2010; Öckinger et al. 2010). Feeding generalists may be more resilient to fluctuations in resource availability than specialists with narrow diets (Root 1973; Kassen 2002; Rand et al. 2006), and social species may be more adaptive to environmental stress than solitary species due to risk spreading among group members (Gadagkar 1990; Hoiss et al. 2012).

Bees (Superfamily: Apoidea) are a species-rich group of mobile insects threatened by habitat loss (Potts et al. 2010; Goulson et al. 2015). While bee taxonomic (Klein et al. 2003; Gomez et al. 2007) and functional (Hoehn et al. 2008; Blüthgen and Klein 2011) diversity are both tied to the delivery of pollination, trait and taxonomic diversity may respond differently to land-use (Murray et al. 2009; Roulston and Goodell 2011; Hoiss et al. 2012). For example, bees who forage further may be able to persist in fragmented landscapes (Gathmann and Tscharntke 2002; Bommarco et al. 2010; Härterl and Steffan-Dewenter 2014; Redhead et al. 2016), but they may also be exposed to agrochemicals and pollutants (Hladik et al. 2016; Long and Krupke 2016). Higher reproductive capacity may buffer bee losses against disturbances (De Palma et al. 2015; Persson et al. 2015), or result in greater resource requirements (Schmid-Hempel and Durrer 1991; Woodard and Jha 2017). To date, investigations into how environmental landscapes structure bee functional diversity have focused on bee communities in natural or agricultural systems (e.g., Hoiss et al. 2012; Kennedy et al. 2013). We know less about bee functional diversity in urban systems (but see Martins et al. 2017; Normandin et al. 2017; Bulchholz et al. 2020).

Urbanization—the conversion of natural habitat into impervious land cover—often drives biotic homogenization, wherein a few species with competitive traits become dominant while many native species are lost (McKinney 2006). Urbanization impacts bee species richness and abundance, often due to losses of floral or nesting resources at local and landscape scales (e.g., Fischer et al. 2016; Ballare et al. 2019; Guenat et al. 2019). These local and landscape features may filter which bee traits are dominant within the community (Hung et al. 2019; Büchholz et al. 2020). For example, habitat and feeding specialists across multiple insect taxa often respond negatively to urbanization (Threlfall et al. 2015; Knop 2016), while generalist responses are highly variable (Burbkan and Gardiner 2014). It is less clear if these generalizations apply to urban bees. Urbanization may also affect the relative representation of bee species characterized by their nesting behavior; Cane et al. (2006) found that ground-nesting specialists were underrepresented and less abundant than cavity-nesting species in smaller urban habitat fragments, and Quistberg et al. (2016) found that mulch cover in urban gardens is associated with fewer ground-nesting bees. Landscape-level features around urban sites are also important if remnant and restored habitat provide...
resources for species with particular traits (Buchholz et al. 2020).

Within cities, gardens are increasingly planted to support pollinators. Urban gardens may harbor high levels of bee trait diversity in comparison to other urban habitat types such as parks and cemeteries (Normandin et al. 2017). In this study, we evaluate and compare how local and landscape features influence bee taxonomic diversity and functional traits in community gardens spanning more than 125 km of central California, with the goal of understanding how gardens can be managed to support bees. We ask if bee richness, bee abundance, and functional traits of bees vary with local garden management and landscape features.

Methods

Study system

The study took place in 18 urban community gardens in the California central coast in Monterey, Santa Clara, and Santa Cruz counties (Fig. 1). The gardens are surrounded by natural, agricultural, and urban land cover, reflecting a gradient in landscape diversity and intensity. The gardens are > 2 km apart in proximity from one another, have been in cultivation for 5 to 47 years, and are 444 to 15,525 m² in size.

Bee surveys

We collected bees in each garden during summer 2015 using aerial netting and pan trapping methods. Sampling occurred over 5 periods between mid-June and mid-September 2015 (approximately every 3 weeks). We aerial netted within a 20 × 20 m plot placed at the center of each garden for 30 min. We also pan trapped using elevated color pan traps (yellow, white, and blue). Traps were mounted atop 1.2 m tall PVC pipes and the pan bowls were filled with a water and 1.5% soap solution. On sampling days, we positioned traps (one yellow, one white, one blue) in a triangular formation, 5 m apart within the 20 × 20 m survey plot. Traps were left for a total of 9–11 h, set out at 8–9 AM and collected the same day between 5 and 7 PM. We identified bees using online resources, image databases, books, and dichotomous keys (see Online Appendix A for full list and citations). We identified bees to species (or morphotaxon for certain genera such as Lasioglossum (Dialictus) where species identification is difficult). Identifications were verified by researchers trained in bee identification according to Michener et al. (1994) at the 2014 Bee Course (American Museum of Natural History).

Trait data

We selected six bee traits recognized for their functional role in bee assemblages (Violle et al. 2007; Williams et al. 2010): body size, sociality, feeding behavior, nesting location, nesting behavior, and pollen-carrying structure (Tables 1, 2). We refer to these as ‘functional traits.’ We searched the literature (see Online Appendix A, Table S1) to determine which attributes each species exhibits in terms of sociality (solitary, social), feeding behavior (polyleptic, oligolectic, variable), nesting location (above ground, below ground, both), nesting behavior (excavate, rent), and pollen-carrying structure (scopa on leg, scopa on abdomen, corbicula, crop). For social behavior, we grouped solitary, sub-social, and communal species as “solitary,” and advanced eusocial and primitively eusocial species as “social.” For parasitic species, we categorized their sociality as “parasitic,” pollen-carrying structure as “no structure,” their nesting location as the location of their host species, nesting behavior as “NA.” We used bee identification guides and peer-reviewed literature for references (see Online Appendix A, Table S1).

Local and landscape features

We surveyed garden habitat characteristics at each period within a 20 × 20 m plot at the center of each garden. We counted and identified all trees and shrubs, and noted whether they were in flower. Then, in four, randomly placed, 1 × 1 m quadrants within the plot, we measured the height of the tallest herbaceous
vegetation, identified all herbaceous plants and classified them as crops, ornamental plants or weeds, counted all flowers and the number of species in flower, and estimated the percent ground cover from
mulch, bare soil, herbaceous plants, and leaf litter. We also measured the size of each garden. We averaged values collected over the sample periods for each habitat measurement for each garden. For landscape features, we used the 2016 National Land Cover Database (NLCD, Homer et al. 2020) in ArcGIS to calculate the proportion of urban, agricultural, and natural land use cover within (2 km) buffers surrounding each garden—a scale at which bees respond to landscape scale features in our system (Egerer et al. 2017) and in others (Kremen et al. 2004).

Analysis

We tested the response of bee abundance and richness to local and landscape features. We then evaluated the role of these features on functional diversity. Because many local and landscape features were correlated, we prioritized including those previously found to be important in describing pollinator diversity in these gardens (Quistberg et al. 2016; Plascencia and Philpott 2017). We identified collinearity of features in our models by calculating the variance inflation factor (VIF) for each variable using the car package (Fox et al. 2018) in R (R Core Team 2020). We used a cutoff score of 3 and removed the predictor features with the highest value in a stepwise approach until all features received a score below the cutoff. Our models included the response variables: garden size (acres), the number of flower individuals, the number of tree and shrub individuals, proportion bare soil, the number of herbaceous plant species (richness of crops, weeds, and ornamentals), and urban cover within 2 km. We additionally calculated Pearson’s correlations for

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### Table 1: Description of traits described

| Trait                        | Value range       | Description                                                                 | Example species          |
|------------------------------|-------------------|----------------------------------------------------------------------------|--------------------------|
| Intertegular span (ITD)      | 1 to 7.64 mm      | Only data for females were used in final analyses                          | NA                       |
| Sociality                    | Social            | Eusocial bees or primitively eusocial bees                                | Apis mellifera           |
| Solitary                     | Bees that forage, nest, and reproduce individually for the majority of their lifespan | Megachile apicalis      |
| Parasitic                    | Bees that invade nests of other species | Nomada edwardsii             |
| Nesting location Above ground | All nest types, from cavity nests to tunnels burrowed into wood | Ceratina acantha         |
| Nesting location Below ground | Tunnels, chambers, cavities in substrate | Eucera actuosa             |
| Nesting behavior Excavate    | Bees that dig into a substrate | Agapostemon texanus           |
| Nesting behavior Rent        | Bees that utilize pre-existing cavities, stems, and other nesting materials | Hylaeus mesillae         |
| Pollen-carrying structure   | Corbicula         | Structural “pollen basket” on hind tibia                                 | Bombus caliginosus       |
| Crop                         | Bees that transport pollen in their digestive tract and regurgitate at the nest | Hylaeus punctatus        |
| Nesting behavior Abdomen     | Dense mass of branched hairs on abdomen | Bombus vosmenenskii         |
| Nesting behavior Scopa       | Dense mass of branched hairs on hind leg                                 | Melissodes robustor      |
| Feeding                      | Polylectic        | Specialists = oligolectic                                                | Xylocopa tabiniformis    |
| No structure                 | Parasitic bees lack pollen-carrying structures                          | Coelioxys rufitarsis     |
| Feeding                      | Oligolectic       | Generalists = polylectic                                                 | Dufourea spp.            |
| Variable                     | When trait varies within taxon                                          | Lasioglossum spp.        |
Table 2  Bees trait values for each species collected

| Species                          | ITD (mm) | Sociality | Nest location | Nesting behavior | Pollen-carrying structure | Feeding behavior   | Abundance Included in trait analyses |
|----------------------------------|----------|-----------|---------------|------------------|---------------------------|--------------------|--------------------------------------|
| Agapostemon texanus              | 1.98     | Solitary  | Belowground   | Excavate scopa.leg | Polylectic                | Unknown            | 76 Yes                               |
| Andrena sp.                      | 1.77     | Solitary  | Belowground   | Excavate scopa.leg | Polylectic                | Unknown            | 1 No                                 |
| Anthidium manicatum             | 4.89     | Solitary  | Aboveground   | Rent scopa.abdomen | Polylectic                | 2 Yes              |
| Anthophora urbana               | 2.98     | Solitary  | Belowground   | Excavate scopa.leg | Polylectic                | 7 Yes              |
| Apis mellifera                  | 3.14     | Social    | Aboveground   | Rent Corbicula     | Polylectic                | 1778 Yes           |
| Ashmeadiella buconis            | 1.52     | Solitary  | Aboveground   | Rent scopa.abdomen | Oligolectic               | 6 Yes              |
| Bombus caliginosus              | 3.58     | Social    | Both          | Rent Corbicula     | Polylectic                | 146 Yes            |
| Bombus vosnesenskii            | 5.17     | Social    | Both          | Rent Corbicula     | Polylectic                | 144 Yes            |
| Ceratina acantha                | 1.15     | Solitary  | Aboveground   | Excavate scopa.leg | Polylectic                | 70 Yes             |
| Ceratina manicata               | 1.02     | Solitary  | Aboveground   | Excavate scopa.leg | Polylectic                | 27 Yes             |
| Coelioxys rufitarsis            | 3.316    | Parasitic | Aboveground   | NA no.structure    | Polylectic                | 2 Yes              |
| Colletes sp.                    | 2.49     | Solitary  | Belowground   | Excavate scopa.leg | Polylectic                | Unknown            | 7 No                                 |
| Eucera sp.                      | 3.31     | Solitary  | Belowground   | Excavate scopa.leg | Polylectic                | Unknown            | 1 No                                 |
| Eucera actuosa                  | 3.9      | Solitary  | Belowground   | Excavate scopa.leg | Polylectic                | 4 Yes              |
| Halictus confusus               | 1.341    | Social    | Belowground   | Excavate scopa.leg | Polylectic                | 3 Yes              |
| Halictus ligatus                | 1.53     | Social    | Belowground   | Excavate scopa.leg | Polylectic                | 49 Yes             |
| Halictus sp.                    | 1.41     | Social    | Belowground   | Excavate scopa.leg | Polylectic                | 4 No                |
| Halictus rubicundus             | 2        | Social    | Belowground   | Excavate scopa.leg | Polylectic                | 1 No                |
| Halictus tripartitus            | 1.2      | Social    | Belowground   | Excavate scopa.leg | Polylectic                | 454 Yes            |
| Holcopasites sp.                | NA       | Parasitic | Belowground   | NA no.structure    | Polylectic                | 1 No                |
| Hylaeus leptocephala            | 1.13     | Solitary  | Aboveground   | Rent Crop          | Polylectic                | 21 Yes             |
| Hylaeus mesillae                | 1.17     | Solitary  | Aboveground   | Rent Crop          | Polylectic                | 99 Yes             |
| Hylaeus panamensis              | 1        | Solitary  | Aboveground   | Rent crop          | Polylectic                | 13 Yes             |
| Hylaeus polifolli               | 1.43     | Solitary  | Aboveground   | Rent Crop          | Polylectic                | 7 Yes              |
| Hylaeus punctatus               | 1.46     | Solitary  | Aboveground   | Rent Crop          | Polylectic                | 42 Yes             |
| Hylaeus rudbeckiae              | 1.14     | Solitary  | Aboveground   | Rent Crop          | Polylectic                | 76 Yes             |
| Hylaeus sp.                     | 1.15     | Solitary  | Aboveground   | Rent CROP          | Polylectic                | 19 No              |
| Lasioglossum Dialictus sp.      | 1.14     | Social    | Belowground   | Excavate scopa.leg | Polylectic                | 104 Yes            |
| Lasioglossum Spechodoagastria   | 1.3      | Social    | Belowground   | Excavate scopa.leg | Polylectic                | 9 Yes              |
| Lasioglossum Lasioglossum sp.   | 1.22     | Solitary  | Belowground   | Excavate scopa.leg | Polylectic                | Variable 33 Yes    |
| Lasioglossum sp.                | 1.26     | Unknown   | Belowground   | Excavate scopa.leg | Polylectic                | Variable 101 No    |
| Megachile apicalis              | 2.31     | Solitary  | Both          | Rent scopa.abdomen | Oligolectic               | 1 No                |
| Megachile brevis                | 2.77     | Solitary  | Both          | Rent scopa.abdomen | Polylectic                | 4 Yes              |
| Megachile perihirta             | 4.13     | Solitary  | Belowground   | Excavate scopa.leg | Polylectic                | 30 Yes             |
these variables (Online Appendix A, Table S2) and tested for collinearity between bee abundance and bee richness using the Hmisc package (Harrell and Dupont 2019). We included site as a random effect. To model total bee abundance, we fit a negative binomial error model to account for overdispersion. To model richness, we fit a Gaussian error model.

We fit linear and generalized linear mixed models (LMM and GLMM, respectively) to determine which local and landscape features influence bee richness and abundance. Our response variables were total bee abundance at each site at each sampling period and total bee richness at each site at each sampling period, calculated by merging pan trap data and aerial net observations. We included all specimens identified to species and morphotype. We used a model selection framework using the MuMln package (Barton 2018) and averaged the top models within two AICc points of the top model. Our approach averages a set of coefficients that are conditional on a specific set of covariates (Cade 2015; Banner and Higgs 2017). While we ensured little correlation between variables, it is controversial if averaging approaches should be used in cases where there is any multicollinearity between variables as this can lead to spurious results (Gunst 1983; Walker 2017). We used standard model assessment techniques to determine whether the final model met all the assumptions of a GLMM/LMM. To compute the unconditional variance, we used the method proposed by Burnham and Anderson (2004), as implemented in MuMIn. To determine the goodness-of-fit of the best models, we calculated a conditional pseudo-$R^2$ value using the r.squareGLMM function.

We used the combined fourth corner and RLQ modelling approach in the ade4 package (Dray and
Dufour 2007) in R to determine which local and landscape features of urban gardens filter bee traits. The RLQ method summarizes the joint relationships between local and landscape features (R matrix), bee species distribution among gardens (L matrix), and bee traits (Q matrix). The fourth corner approach tests for direct correlations between local and landscape features and bee traits (Dray and Legendre 2008; Dray et al. 2014). For the analysis, we removed singleton species and morphotypes, as well as morphotypes that were missing trait data (those not identified to species) from the species abundance matrix (L) (19 species removed from trait analyses, 142 individuals). We included the same local and landscape features as in the analyses above. Following Dray et al. (2014), we first performed the RLQ method that consisted of a correspondence analysis on the species (L) matrix and principal component analysis on the R and Q matrices and used permutation models to evaluate (a) if garden features influence the distribution of bee traits (“model 2”), and (b) if traits influence the composition of species assemblages in relation to the local and landscape features (“model 4”). We visualized the RLQ results using a biplot to assess the global relationships between bee traits and local and landscape features. Second, we used the fourth corner analysis to determine the significance of each bee trait-factor relationship. For the analyses, we used Monte-Carlo permutations (n = 999) to test for correlations between quantitative variables and used the ‘D2’ correlation coefficient at a 0.95 level of significance to test for associations between quantitative variables and each categorical variable separately (Dray et al. 2014).

Results

Local and landscape features and bee richness and abundance

We documented 3537 individuals from 59 species and morphotypes of bees representing various traits (Table 1). The five most common bees recorded (74.2% all individuals) included honey bees (Apis mellifera), sweat bees (Halictus tripartitus and Lasiosglossum (Dialictus) sp.), and bumble bees (Bombus vosnesenskii and Bombus caliginosus). 28.8% of species or morphotypes were rare and were recorded only once across sites. These include species with abdominal scopae (e.g., five species in Megachile) but also cleptoparasitic bees (Holcopasites sp.), mining bees (Andrena sp.), and others.

We found that both local and landscape features predicted bee species abundance, richness and traits in gardens. Bee abundance was significantly associated with garden size (Fig. 2) and marginally associated with urban cover (2 km), where larger gardens had more bee individuals (p < 0.05) and gardens surrounded by more urban cover had fewer individuals (p = 0.056). Bee richness was significantly associated with the number of flowers, garden size, and urban cover (2 km). Larger gardens (p < 0.01) and gardens surrounded by more urban cover (p < 0.01) had more bee species (p < 0.01), and gardens with more bare soil (p < 0.05) and more flowers (p < 0.001) had fewer species (Table 3 for test statistics for each model) (Fig. 3).

Local and landscape features and bee traits

The gardens supported bees with a diverse range of body sizes, social behaviors, nesting locations, pollen-carrying structures, lecty, and parasitic behaviors (Tables 1, 2). The fourth corner test for bivariate relationships between traits and local and landscape factors found that below ground nesting, excavating, and legs with scopae were more positively associated with sites surrounded by more urban cover within
2 km, while above-ground nesting, renting, and corbicula were negatively associated with urban cover (all relationships \( p < 0.05 \), Fig. 4a).

The combined RLQ and fourth-corner analysis revealed that, overall, local and landscape features did not influence the distribution of bee species (Model 2, \( P_2 = 0.141 \)) and bee traits did not influence the composition of species found in sites with certain local and landscape features (Model 4, \( P_4 = 0.319 \)). However, individual features were still significantly related to changes in certain bee traits. The first trait axis was driven by differences in the number of flowers (\( p < 0.05 \)), where gardens with fewer flowers had fewer species exhibiting above-ground nesting, renting, and corbicula, and more species exhibiting below-ground nesting, excavating, and scopa on legs. The second trait axis was driven by urban cover (\( p < 0.01 \)), where gardens surrounded by more urban cover were negatively associated with social bees and positively associated with solitary behavior and pollen transport in the crop (Fig. 4b, c).

### Table 3

The estimates, adjusted standard errors, test statistics, and \( p \) values for the top model for wild bee abundance and top model for bee richness

| Model         | Predictor                  | Estimate | Std. error | Z value | \( p \) value | DF | Pseudo R-sq |
|---------------|----------------------------|----------|------------|---------|--------------|----|-------------|
| Bee abundance | Garden size                | 0.193    | 0.083      | 2.303   | 0.021*       | 17 | 0.355       |
|               | Urban cover (2 km)         | -0.061   | 0.032      | 1.905   | 0.057        |    |             |
|               | No. flowers                | -0.178   | 0.099      | 1.789   | 0.0736       |    |             |
|               | Bare (1 m)                 | -0.124   | 0.077      | 1.606   | 0.108        |    |             |
|               | Herbaceous plant spp.      | 0.114    | 0.073      | 1.553   | 0.121        |    |             |
|               | Trees and shrubs           | 0.097    | 0.083      | 1.168   | 0.243        |    |             |
| Bee richness  | Garden size                | 1.267    | 0.433      | 2.923   | 0.003**      | 17 | 0.479       |
|               | Urban cover (2 km)         | 0.514    | 0.1866     | 2.757   | 0.006**      |    |             |
|               | No. flowers                | -1.537   | 0.5422     | 2.835   | 0.005**      |    |             |
|               | Bare (1 m)                 | -0.790   | 0.387      | 2.039   | 0.061        |    |             |
|               | Herbaceous plant spp.      | -0.587   | 0.3887     | 1.511   | 0.131        |    |             |
|               | Trees and shrubs           | 0.682    | 0.446      | 1.529   | 0.126        |    |             |

Estimates and errors are conditional model averages calculated from those models within delta 2 AICc of the top model. Intercepts are scaled values

Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

Fig. 3 The variables significantly associated with wild bee richness. Garden size (acres) and urban cover (2 km) were positively related to richness. The number of flowers was negatively related to richness. The black line indicates the slope estimate and the gray is the 95% confidence interval around the estimate. Points represent wild bee abundance at a site at each survey period.
Discussion

Local and landscape features and bee richness and abundance

Our study adds to the growing evidence that urban environments can support bee species (Matteson et al. 2008; Verboven et al. 2012), and that bee richness, abundance, and traits are driven by local and landscape features of the urban environment. In this system, percent urbanization ranged from 7.85 to 93.15% and we found that gardens surrounded by more urban land cover supported a higher richness of bee species. This is possibly because the urban landscape in this region is a mosaic of suburban and non-industrial areas such as gardens, parks, and semi-natural habitat that can serve as refugia and resources for a diverse assemblage of species within the urban matrix (Hall et al. 2017). Alternatively, gardens may act as resource-rich islands within an inhospitable habitat, attracting species who have nowhere else to go. Interestingly, we found that bee abundance declined with more urban cover, echoing the findings of similar studies (Fortel et al. 2014). In other words, while the urban environment supported a high diversity of bees, their abundance was low. If the urban matrix indeed includes diverse resources for pollinators, one speculative explanation is that gardens may not attract large numbers of bees from nearby home gardens and parks, which have been found to host an abundance of bees (Lowenstein et al. 2014; Hülsmann et al. 2015). These findings suggest that urbanization should be conceptualized as a process of habitat change, rather than habitat loss, with implications for the distribution of biodiversity.

In this system, urban cover was negatively co-linear with tree and shrub abundance, and therefore the positive impact of urban cover on bee richness may
indicate that tree and shrub abundance had a negative impact on bee richness. This was unexpected, as was the finding that the number of flowers was negatively associated with bee richness. We suggest that increases to floral abundance and tree and shrub abundance in the gardens are associated with a dilution effect. Dilution has previously been observed for pollinators visiting sites with increase floral resources, especially in sites with high landscape diversity (Sha and Vandermeer 2009; Riedinger et al. 2014; Wenneniger et al. 2016). This finding further builds the case against the assumption that all urban landscapes are bereft of resources for pollinators, and we propose that future studies in this region employ detailed, fine-scale assessments of urban landscape-level features. An alternative explanation is that an increased abundance of flowers may have not attracted bee species if they were the “wrong” type. It is well established that pollinator preference plays a role in community assembly. Because bees have preferences for certain plant groups (e.g., Asteraceae, Lamiaceae), flower identity and composition are important for shaping bee communities in urban gardens (Frankie et al. 2005), even more so than floral abundance (Lowenstein et al. 2019). Furthermore, although it is often assumed that native plants promote bee species in gardens, this has not been found previously in this system (Egerer et al. 2020), where the average native plant richness across gardens is 24.4% of all plants cultivated and ranges from 0 to 100%. In other systems, native plants may (Pardee and Philpott 2014) or may not (Matteson and Langelloto 2011) influence bee community composition. We therefore suggest that more research on plant preference is critical for promoting urban pollinators.

The only local factor important for both bee richness and abundance was garden size. Larger gardens supported both greater bee richness and greater abundance. Our observation supports previous research on the importance of habitat patch size for promoting bee diversity in agricultural and natural systems (Steffan-Dewenter 2003; Krauss et al. 2009; Hopfenmüller et al. 2020). Moreover, our findings are similar to other urban bee studies showing a positive effect of garden size on bees (Frankie et al. 2005; Cane et al. 2006; Matteson and Langellotto 2011; Pardee and Philpott 2014; Quistberg et al. 2016). However, some studies have found no effect of garden size on bees (Potter and LeBuhn 2015; Lanner et al. 2019). Garden size is likely associated with a greater number and diversity of resources for bees that could impact richness and abundance, though this may be region-dependent.

Local and landscape features and bee traits

While habitat change is often associated with colonization by generalist species across multiple systems (Rand and Tscharntke 2007; Rocha and Fellowes 2020), we did not find an association between lecty and urbanization or between lecty and any other factor we measured. Bee diet breadth can explain sensitivity to land use changes in agricultural (Steffan-Dewenter et al. 2006; Moretti et al. 2009), natural systems (Hopfenmüller et al. 2014), and urban systems (Fetridge et al. 2008; Banaszak-Cibicka and Żmihorski 2012), yet our results do not support any generalizations between diet and urbanization. We did find a link between urbanization and sociality—urban land cover supported more solitary bees than social bees. It has been observed that bees exhibiting social traits may be more impacted by disturbances (Winfree et al. 2009) and isolation from natural habitat (Williams et al. 2010). For example, Wilson and Jamieson (2019) found that social Bombus and Lasioglossum were negatively affected by urbanization, though other work has found either positive or no association between urbanization and social behaviors (Carper et al. 2014; Guenat et al. 2019). One possible explanation is that species that exhibit solitary behaviors may be able to utilize urban environments in ways that social bees cannot and in ways not captured by our trait measurements.

Bees with scopa were positively associated with urban land cover. This finding is possibly driven by bees in the genus Halictus, which are a highly abundant group in our garden sites that feature scopa and are known to tolerate and utilize a wide range of floral resources (Cane 2015). Finally, like Buchholz et al. (2020), we found that urbanization was associated with nesting behavior and location, with gardens surrounded by more urban cover associated with below ground nesting species and excavators. This could be because these gardens provide the necessary soil substrate that is otherwise limited in highly urbanized areas with greater impervious land cover. Nesting location has previously been associated with responses to isolation from natural habitat, where
species that nest above ground were most sensitive to habitat loss than below ground species (Williams et al. 2010). Clearly, more work is needed to make strong generalizations about the impact of urbanization, but this study suggests that urbanization is a strong determinant of trait composition in urban gardens.

We observed that local floral resources were negatively associated with solitary bee traits and with bees that carry pollen internally in their crop (a specialized behavior specific to bees such as those in the genus *Hylaeus*). This is surprising, because it is often assumed by conservation initiatives that flowers promote a variety of wild bees with different traits. But this study shows that not all bees benefit universally from increased floral abundance in gardens. However, our finding may be biased because we did not observe many social bee species, and the most abundant social species in our system was the honey bee (*Apis mellifera*), a generalist known to utilize a very wide range of plant species and that has been hypothesized to outcompete native species (Steffan-Dewenter and Tscharntke 2000). Indeed, the abundance of honey bees has previously been negatively correlated with wild bee species richness in our study sites (Plascencia and Philpott 2017). In urban gardens, Threlfall et al. (2015) found that the presence of exotic, non-native plants had a positive impact on honey bees, but not wild bees, suggesting that there is an interaction between floral identity and the prevalence of social insects in the garden that our study could not pinpoint. More research is needed on which floral types support a diverse assemblage of pollinators (see Rollings and Goulson 2019).

We expected to find a relationship between body size and local or landscape features because body size is directly related to foraging range (Greenleaf et al. 2007). Other urban bee studies have shown that urbanization filters for smaller bees (Hamblin et al. 2017), potentially due to heat island effects (Eggenberger et al. 2019), where larger bodied bees are more vulnerable to overheating than smaller species. We suggest that body size does not consistently respond to urban landscape-scale fragmentation, as some studies have found that small-bodied bees are more sensitive (e.g., Cane et al. 2006) while other studies have concluded that small-bodied bees respond positively to urbanization (e.g., Wray et al. 2014). We expect that this variability reflects the fact that cities vary widely across regions. There may also be a trade-off between the ability to access widely distributed resources and the ability to thrive on locally scarce resources (Harrison and Winfree 2015).

Our study results may also reflect some limitations of our methodologies. In our analyses, we pooled species with the same life-history traits for the fourth corner analysis. One limitation of this approach is that it may mask general responses (Williams et al. 2010). For example, if most large bees increased in response to urbanization but one abundant, small bee decreased, decline of the abundant species could obscure increases in the number of the others. We also acknowledge the difficulty of categorizing species by their functional traits. We justified our choices based on previous analyses. For example, following Forrest et al. (2015), we described species as either solitary or social, simplifying categories together. This means that socially polymorphic species such as *Halictus rubicundus*, which can have both eusocial and non-eusocial relatives within populations, was classified as social.

**Conclusion: ecological applications**

We confirm that bee abundance and richness respond to urbanization (Guenat et al. 2019; Ballare et al. 2019). Urbanization was associated with increases to bee richness and decreases to bee abundance, suggesting that while multiple species can utilize urban systems, their populations are low in abundance. While landscape-scale urbanization is not easily managed by local community gardeners, examining trait diversity reveals that local-scale management (e.g., creating nesting habitat) could affect the assemblage of the bee community. While we contribute to a growing number of studies that suggest traits can be used to understand how bees respond to habitat management, we also highlight that the trait-based literature has found mixed impacts of urbanization on bee traits (Tonietto et al. 2011; Normandin et al. 2017; Fitch et al. 2019; Wilson and Jamieson 2019). These discrepancies are possibly due to how urbanization and traits are measured. Approaches that utilize distance-based frameworks to measure trait diversity and conceptualize species traits relative to a community average (Laliberté and Legendre 2010; Coux et al. 2016) may be more predictive of how bee communities respond to change landscapes and offer a
promising future methodological direction. Understanding the impact of anthropogenic change on bee communities is important because changes in the composition of species and traits likely influences the functioning of urban ecosystems (Faeth et al. 2001). This is especially relevant in urban agricultural systems like gardens, where bees support pollination services of urban crops (Potter and LeBuhn 2015; Lowenstein et al. 2015). For urban pollinators, we need a greater understanding of which bee traits influence species vulnerability to disturbance and which influence their ability to provision ecosystem services.

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Author contributions HC led study design, participated in fieldwork, coordinated lab work and insect identification, and coordinated manuscript writing. ME analyzed the data and contributed to the manuscript. S–ST collected samples and identified them. SMP led site selection, field research design, fieldwork logistics and contributed to manuscript writing.

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Data availability Data is available on a public repository at github.com/hamutahlc/beetraits.

Code availability Code is available on a public repository at github.com/hamutahlc/beetraits.

Declarations

Conflict of interest We have no competing interests.

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