Determinates of inner city butterfly and bee species richness

Kevin C. Matteson · Gail A. Langellotto

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Abstract Although urbanization is increasing worldwide, relatively few studies have investigated patterns of urban biodiversity outside of city parks and reserves, in urban neighborhoods where people live and work. We evaluated models including local and landscape factors that might influence the bee and butterfly richness of community gardens located within densely populated neighborhoods of the Bronx and East Harlem in New York City (>10,000 people/km²). The gardens were surrounded by buildings and limited amounts of green space (3,600–17,400 building units and 10–32% green space within a 500 m radius). Contrary to our initial prediction that landscape green space might be especially influential in this heavily urbanized setting, the most highly supported models for both bee and butterfly richness (based on Akaike Information Criterion) included just the local, within-garden variables of garden floral area and sunlight availability. There was marginal support for models of bee richness including the number of building units surrounding gardens within a 500 m radius (which exhibited a negative association with bee richness). In addition, perhaps because bees are central place foragers that may nest within or near gardens, supported models of bee species richness also included total garden area, canopy cover, and the presence of wild/unmanaged area in the garden. Generally, our findings indicate that sunlight and floral abundance are the major factors limiting local pollinator diversity in this setting. This suggests that rooftop and other “open” urban habitats might be managed to increase local pollinator diversity, even if seemingly “isolated” within heavily developed neighborhoods.

Keywords Community gardens · New York City · Green space · Urban biodiversity · Modeling species richness
Introduction

Urban areas are expanding worldwide (United Nations Population Division 2006), resulting in habitat alterations that are spatially vast (McKinney 2002). The changes associated with urbanization often alter the composition and distribution of a variety of species (McKinney 2008), including insects (McIntyre 2000). Specifically, native invertebrate diversity generally decreases with increasing urbanization, especially in central urban core areas (McKinney 2008). In addition, prevalence of generalist and/or exotic species tends to increase in heavily developed neighborhoods, which may alter the composition of functionally important groups, such as pollinating insects (Kitahara and Fujii 1994; Clark et al. 2007; Di Mauro et al. 2007; Matteson et al. 2008). Despite these difficulties, small urban habitats in densely populated neighborhoods have the potential to facilitate reconnection of humans and nature (Miller 2005), and provide ecosystem services to the urban populace (Bolund and Hunhammar 1999). Indeed, urban habitats located in the most densely populated neighborhoods may provide the greatest benefit to the most people but, ironically, are the most likely to suffer from biotic homogenization and reduced native species richness (McKinney 2006). Despite the potential value of small urban habitats (Bolund and Hunhammar 1999), relatively few studies have investigated the factors governing biodiversity in heavily developed and densely populated neighborhoods.

Human management of urban land may exert a profound influence on biodiversity of pollinating insects, both by determining the diversity of vegetation types and the abundance of floral resources. In addition, local species richness may be influenced by landscape-level variables external to urban habitats, such as the proportion or diversity of habitat in the surrounding landscape (Levins 1969; McIntyre and Hostetler 2001; Clark et al. 2007; Di Mauro et al. 2007; Öckinger et al. 2009). Urban vegetation is distributed among fragmented and heavily managed gardens, parks, greenways, and other green spaces, all of which may facilitate movement of insects and other wildlife, and increase local species richness. Conversely, densely populated neighborhoods often contain tall buildings that may limit insect richness by obstructing dispersal and/or reducing sunlight (Ruszczyk and De Araujo 1992).

New York City is the largest urban area in the United States, with 8.2 million people living in five boroughs (United States Census Bureau 2006). Within the city, there are over 700 community gardens (Council on the Environment of New York City 2006), most of which are open to the public and rely on the direct involvement of local citizens as garden managers. Many community gardens are located in high density and low income neighborhoods (Matteson 2007), providing an opportunity to evaluate biodiversity in an inner city environment. Specifically in the neighborhoods of this study (located in the central Bronx, and East Harlem in Manhattan), more than 70% of the land area is composed of buildings, sidewalks, roads, and other paved areas, and human density is 19,000 people/km² and 15,000 people/km², respectively (New Yorkers For Parks 2005). For this study, we investigated the relative influence of local and landscape variables on bee and butterfly species richness in community gardens located in these neighborhoods. We hypothesized that, in addition to local variables such as floral area, landscape variables (i.e., green space and building density) would strongly influence local species richness.

In addition to contributing to a theoretical understanding of factors influencing insect biodiversity at the far end of the spectrum of urbanization, this study has several practical implications. First, a variety of crops grown in community gardens and other urban green spaces are known to be dependent on (e.g. cucumbers, squashes) or to benefit from (e.g., tomatoes, raspberries) insect pollination (Matteson and Langellotto 2009). Therefore, a
diversity of pollinating insects may increase crop yield and contribute to urban agriculture, providing food security for residents of inner city neighborhoods. For example, the floral area of urban gardens was positively associated with deposition of conspecific pollen in cucumbers (Werrell et al. 2009), presumably because increased floral cover attracted a greater number of pollinators into urban gardens. Second, gardens include a variety of flowering plants (e.g. zinnias, marigolds, sunflowers, mint, basil, and more), some of which may be utilized by insects for pollen, nectar or both. These floral resources may be especially important for bees and butterflies moving through, or living within, developed urban landscapes where there are few alternate food resources. Finally, as mentioned above, the presence of “charismatic microfauna” (Connor et al. 2003), such as bees and butterflies, provides humans with opportunities for interaction with nature, potentially preventing what has been termed the “extinction of ecological experience” (Pyle 1978; Miller 2005) within urban neighborhoods.

Study location and methods

Community garden study sites

The 18 community gardens of this study were located in the central Bronx and East Harlem (Fig. 1) and varied in size (see Results), management and vegetative characteristics. Some gardens were managed primarily for the production of vegetables; others were used for non-

![Map of the locations (yellow circles) of 18 urban community gardens in East Harlem and the Bronx, New York City (note different spatial scales). The map was created using data provided by the USDA Forest Service, Northeastern Research Station.](image)
gardening recreation. Common crop plants grown in these community garden study sites include hot and sweet peppers, mints and other herbs, kale and collards, tomatoes and tomatillos, strawberries, and cucumbers (for the full list, see Matteson and Langellotto 2009). Specific locations (street addresses and GPS coordinates) of these gardens are presented in Matteson et al. (2008).

Local (within garden) variables

For each garden, we measured the following variables: total garden area, vegetable area, floral area, wild/unmanaged area, tree canopy cover, shrub cover, lawn area, area of paths/structures, garden age, and sunlight availability. We calculated garden age as the number of years from the creation of the garden (when soil and plants were added to a formerly abandoned lot) to 2003 (the first year of the study). Sunlight availability of individual gardens was measured using a photometer (Gempler’s 4 in 1 Weather Meter, Gempler’s, Madison, WI, USA). To account for daily and seasonal changes in sunlight availability, we used the average of 30 sunlight measurements taken at different times, days, and in different locations within each garden. Because bee and butterfly species vary in host plant and/or nesting substrate requirements, species richness might increase with the total diversity of land uses within the garden. Therefore, we used the Shannon Index to compute two measures of heterogeneity of land use within gardens. One heterogeneity measure (gardenH) included the proportion of seven within-garden land uses (vegetable area, floral area, wild/unmanaged area, tree canopy cover, shrub cover, lawn area, and area of paths/structures). A second measure (vegH) omitted the contribution of paths/structures and mowed lawn area to garden heterogeneity, which we believed would provide few resources for most pollinators.

Landscape variables

We assessed the proportion of green space surrounding individual gardens using a high spatial resolution map of New York City provided by David Nowak of the USDA Forest Service, Northeastern Research Station. The map included four land cover types (tree/shrub, grass/herbaceous, impervious surface, and water) with a pixel-size of 0.81 m², which were generated using a hybrid classification method (Myeong et al. 2001) applied to high resolution aerial photography taken in 2001 and 2002. In addition to large parks, this map included relatively small, vegetated areas of the city (i.e., street trees, vacant lots, community gardens, vegetated margins along roads, grass, and trees on the grounds of housing projects, etc.), which may contribute to butterfly and bee species richness.

We quantified the number of “building units” surrounding gardens, using a 2001 municipal data set provided by CENYC (Council on the Environment of New York City 2006). Building units include all residential, commercial, and office spaces in a building, providing a proxy for the vertical and horizontal spatial extent of buildings in the vicinity of the gardens. Tall buildings in the vicinity of the gardens often contained over 100 total building units, including commercial units on the ground level and multiple residential units above.

We used the buffer and clip toolbox functions in ArcDesktop 9.3 to quantify the number of building units and green space (combined tree/shrub and grass/herbaceous cover) surrounding each garden in circles with a 200 m radius and a 500 m radius. These distances were chosen because they generally correspond to the average daily movements of many butterflies (Scott 1975) and typical foraging distances of many bees (Gathmann and Tscharntke 2002; Greenleaf et al. 2007). Although some bees (e.g. bumble bees, Xylocopa...
virginica, Apis mellifera) (Greenleaf et al. 2007) and some butterflies (Öckinger et al. 2009) are capable of larger movements, we did not analyze green space surrounding gardens at larger spatial scales because we wanted to minimize spatial overlap among gardens. In addition, we were reluctant to conduct additional analyses that would increase the likelihood of erroneous results (i.e. Type I error), especially considering that measures of landscape connectivity are often highly correlated (Winfree et al. 2005).

Butterfly and bee sampling

We observed butterflies in community gardens weekly from July through September in 2003 and June through September in 2004 and 2005. Bees were sampled using hand-netting (once per month in May, June, July and August 2005 and 2006) and 355-ml yellow bowl traps (bowls: 15 cm in diameter, 4.5 cm in depth, Solo Cup Company, Urbana, IL; detergent: ~4 ml per bowl, Colgate-Palmolive Company, New York, NY) which were set twice a month from July to September in 2003 and May to September in 2004 and 2005. To reduce disturbances within these heavily trafficked gardens, bowls were clearly labeled as research (on the underside and lip of the bowl), suspended from trees, shrubs, or along fences (~1.5–2.5 m), and left out for just 24–48 h (same duration for all gardens) during calm weather conditions. Although this bowl trapping methodology differs from the standardized bee inventory plot (http://online.sfsu.edu/~beeplot/) (we were not aware of this protocol at the onset of the project), the cumulative number of traps collected in this study (>8,000) and the use of species richness estimators (see below) should ensure that the richness of bees and butterflies was sufficiently described. On each sampling date, for every 600 m² of garden area, we collected one yellow bowl trap, conducted 5 min of butterfly counts, and/or spent 10 min hand-netting bees. We scaled sampling effort to garden area in this manner because our goal was to assess the relative influence of local and landscape variables on species richness per garden (rather than per standardized unit area). All gardens were sampled the same number of days per year.

Butterflies were identified using Brock and Kaufman (2003). Following Clark et al. (2007) we classified butterflies by feeding specificity (broad vs. restricted), broods per year, and number of larval host plants used (>10 or ≤10 species). The number of broods per year, specific to the New York City area, was determined using Cech (1991). Habitat specificity and number of larval plants followed Clark et al. (2007) or, for butterflies unique to this study, Cech and Tudor (2005). The ecological characteristics of bees found in the gardens of this study are discussed separately in Matteson et al. (2008). Many bees in the genus Lasioglossum were identified by Sam Droege of the U.S. Geological Survey. All other bees were identified by John S. Ascher of the American Museum of Natural History, where a synoptic collection is maintained.

To determine total species richness (pooled across all sampling dates) in each of the 18 sampled gardens, we used EstimateS software (Version 8.0, R. K. Colwell, http://purl.oclc.org/estimates) to calculate Chao1 (for bees) and Chao2 (for butterflies) nonparametric species richness estimators. Chao2 is based on the incidence of species across sampling dates and thus can be used when a single, standardized sampling method is utilized, as was the case for our sampling of butterflies (bowl sampling was not used for butterflies). Because our sampling protocol for bees involved two separate sampling protocols (both bowl trapping and hand-netting), we used the Chao1 estimator, which is based on abundance of species relative to each other, rather than incidence of species across sampling dates. Both Chao estimators have been demonstrated to be more precise and less biased than using observed species richness or several other estimators (Brose and Martinez 2004; Walther and Moore 2005).
Assessing predictors of insect species richness in urban gardens

We measured 12 local, within-garden variables and four surrounding landscape variables for each garden of this study (Table 1). We constructed models including variables that we considered, a priori, to be most likely to have an impact on insect diversity in this setting. These included total garden area, floral area, and sunlight, all of which are widely known to influence bee and butterfly richness. Canopy cover and wild/unmanaged area per garden also were included because we believed that these variables might correlate with nest site availability (for bees) or host plants (for butterflies). The two measures of vegetation heterogeneity (gardenH and vegH) and garden age were evaluated because we believed diversity of plant types or colonization time might influence insect diversity. Finally, we included each of the four landscape variables (green space and number of building units at the 200 m and 500 m radius scales), resulting in a total of 12 candidate variables that we believed might influence bee or butterfly species richness.

To avoid data mining among so many independent variables and stepwise procedures that are prone to erroneous conclusions (Burnham and Anderson 1998), we utilized an objective approach to model selection. First, we used hierarchical partitioning (rand.hp function in the hier.part package of R statistical program 2.8.1) to identify the independent (uncorrelated) explanatory strength of each individual variable (for details regarding this procedure, see Mac Nally 2002). We then ranked all variables according to their Z-scores (Fig. 2), which indicate the independent influence of each variable on butterfly or bee species richness. Next, we constructed a global model that included all 12 independent variables for both butterfly and bee species richness. Each global model explained the majority of variation in species richness (unadjusted $R^2$ of 0.814 for butterfly richness and 0.900 for bee richness). Therefore, we proceeded to construct models of increasing simplicity (i.e., 11 variables, 10

Table 1 Values of local (within-garden) and surrounding landscape variables of the urban community gardens ($n=18$) of this study. Please refer to the methods for descriptions of all garden variables. Note that “green space” in this study largely consisted of parks, gardens and other human-managed sites, rather than forests or other forms of “less disturbed” habitat

| Variable                      | Units     | Mean ± SD     | Range          |
|-------------------------------|-----------|---------------|----------------|
| **Local variables**           |           |               |                |
| Total garden area             | Meters$^2$| 909±541       | 224–2,188      |
| Vegetable area                | %         | 23±17         | 0–71           |
| Floral area                   | %         | 12±9          | 1–40           |
| Wild/unmanaged area           | %         | 11±21         | 0–90           |
| Tree canopy cover             | %         | 27±15         | 1–71           |
| Shrub cover                   | %         | 5±4           | 0–13           |
| Lawn area                     | %         | 19±24         | 0–69           |
| Area of paths/structures      | %         | 35±26         | 1–80           |
| Garden age                    | Years     | 13±9          | 0–33           |
| Sunlight                      | Mean lux  | 61,000±20,000 | 21,000–81,000  |
| GardenH                       | Shannon Index | 1.38±0.25   | 0.77–1.72      |
| VegetationH                   | Shannon Index | 1.19±0.26   | 0.66–1.58      |
| **Landscape variables**       |           |               |                |
| Green space (200 m)           | Proportion| 0.15±0.05     | 0.08–0.23      |
| Green space (500 m)           | Proportion| 0.21±0.05     | 0.10–0.32      |
| Building units (200 m)        | Number    | 201±113       | 74–473         |
| Building units (500 m)        | Number    | 10,100±4,200  | 3,600–17,400   |
variables, etc.) by sequentially dropping independent variables with the lowest Z-score (least independent influence on the dependent variable) as identified by hierarchical partitioning. We used least squares regressions (Systat2004) to generate each of the 12 models. Akaike’s Information Criterion with a correction for small sample size ($AIC_c$) (Hurvich and Tsai 1989) was used to identify the most informative and parsimonious models (Burnham and Anderson 1998) for the two diversity measures. To enable comparison among models, evidence ratios were calculated as the ratio of two competing models’ Akaike weights (Burnham and Anderson1998). We considered an evidence ratio greater than 2.7 (which is equivalent to $\Delta AIC_c >2$) to indicate a better model (Burnham and Anderson1998).

To increase normality, the following variables were log10-transformed: total garden area, the number of building units at the 200 m and 500 m scale, Chao1 bee species richness, and Chao2 butterfly richness. Also to improve normality, vegetable area and tree canopy area per garden were square-root transformed, sunlight availability was cubed and green space proportions were arcsine square-root transformed (Sokal and Rohlf1995). After transformation, all above independent and dependent variables did not significantly deviate from a normal distribution (Shapiro-Wilk’s test, $p>0.05$ for all variables). The amount of wild/unmanaged area was not normally distributed because nearly half of the gardens had no wild/unmanaged space in the garden. Therefore, we characterized gardens with a dummy variable based on whether or not they contained any wild/unmanaged areas. The Durbin-Watson test indicated that all models met assumptions of linearity and homoscedasticity.

**Results**

Garden characteristics and insect composition

The community gardens of this study were created fairly recently (mean of 13 years old) and were relatively small (mean area of 909 m²; range of 224–2,188 m²) (Table 1). At the 500 m scale, just 21% of the landscape surrounding gardens consisted of vegetation (range of 10–
32%), whereas there was a mean of over 10,000 building units (range of 3,600–17,400) (Table 1). Within gardens, the largest use of land was paths/structures (mean = 35% of garden area), followed by vegetable beds (23%), lawn (19%), and floral area (12%). Gardens were largely open, with tree canopy covering just over a quarter of garden area (27%), on average.

Over 3 years, we documented 3745 individual butterflies, representing 24 species/taxa, in these gardens (Table 2). The introduced cabbage white butterfly (Pieris rapae) was found in all gardens and was by far the most common butterfly. Other common butterflies included spring azures, Celastrina ladon spp. and red admirals, Vanessa atalanta. Most of the 24 butterfly species/taxa observed were seasonal, habitat, or host plant generalists (Table 2). Specifically, just three species had single broods per year and only five species had restricted habitat requirements. Although nine butterfly species utilized 10 or fewer larval host plants, many of the required host plants used by these species are common in urban areas (e.g. nettles, elms, violets, crabgrass). Observed butterfly species richness ranged from 3 to 16 species/taxa per garden (mean ± SD=9±4). The Chao2 estimate of butterfly species richness ranged from 7 to 41 (mean ± SD=16±8) species per garden.

A total of 1,145 collected bees were identified to 54 species (including Apis mellifera). We more fully discuss the composition and ecological characteristics of these bee species in Matteson et al. (2008). Briefly however, the most abundant species were two exotic yellow-faced bees (Colletidae), Hylaeus leptocephalus and H. hyalinatus, and the native bumble bee Bombus impatiens. Native, small-bodied bees in the genus Lasioglossum were also prevalent. Observed bee species richness ranged from 7 to 29 species per garden (mean ± SD=18±7) whereas the Chao1 estimate of bee species richness ranged from 7 to 46 species per garden (mean ± SD=25±11).

Models of insect richness

For butterfly richness, the most informative model included just floral area and sunlight (ΔAICc=0.0; Wf=59%, adj. $R^2=0.601$; Table 3). There was also some support for a simple univariate model including only garden floral area (ΔAICc=1.39; Wf=30%, adj. $R^2=0.513$). The next most supported model included floral area, sunlight, and garden age (which exhibited a negative relationship with butterfly richness), but this model garnered just 10% of the support (Table 3). All other multivariate models of butterfly species richness were not well-supported (Akaike weights less than 5%).

For bee species richness, there was nearly equivalent support for two models. One model included total garden area, floral area, and sunlight (ΔAICc=0.0; Wf=35%, adj. $R^2=0.678$; Table 4). The other model included just total garden area and floral area (ΔAICc=0.18; Wf=32%, adj. $R^2=0.622$; Table 4). Together, these two models contained 67% of the support and each had more than two times the support of any subsequent model. The number of building units surrounding gardens at the 500 m scale was present in two models (one with 12% of the support, and one with 9% of the support) whereas garden canopy cover was included in a model that garnered 10% of the support.

Discussion

Influence of local variables on insect richness

As expected, floral area per garden had the greatest influence on both butterfly and bee species richness. Strong relationships between local plant/floral resources and butterfly and
| Common name                          | Family      | N     | Broods/year | Habitat specificity | Number (types) of larval host plant |
|-------------------------------------|-------------|-------|-------------|---------------------|-------------------------------------|
| Cabbage white (Pieris rapae)        | Pieridae    | 3,306 | >3          | Broad               | >10 (various crucifers)             |
| Spring azure (Celastrina ladon spp.)| Lycaenidae  | 83    | 1           | Broad               | >10 (ex. New Jersey tea, Spiraea)   |
| Red admiral (Vanessa atalanta)      | Nymphalidae | 70    | >3          | Broad               | ≤10 (nettles)                       |
| Common sootywing (Pholisora catullus)| Hesperiidae | 62    | 3           | Broad               | >10 (ex. pigweeds, lamb’s quarters) |
| Painted lady (Vanessa cardui)       | Nymphalidae | 46    | 2–3         | Broad               | >10 (ex. thistles, mallows)         |
| Question mark (Polygonia interrogationis) | Nymphalidae | 42    | 2           | Restricted          | ≤10 (nettles, elms, hackberries)    |
| Monarch (Danaus plexippus)          | Nymphalidae | 30    | <3          | Broad               | ≤10 (common milkweed, dogbane)      |
| Sulphur (Colias spp.)               | Pieridae    | 20    | >3          | Broad               | >10 (ex. legumes, esp. clovers)     |
| Eastern tiger swallowtail (Papilio glaucus) | Papilionidae | 16    | 2           | Restricted          | >10 (ex. cottonwood, ash, tulip)    |
| Eastern tailed blue (Everes comyntas) | Lycaenidae  | 16    | >3          | Broad               | >10 (ex. legumes, clovers)          |
| Black swallowtail (Papilio polyxenes) | Lycaenidae  | 12    | 2           | Broad               | >10 (ex. Queen Anne’s lace, fennel) |
| American lady (Vanessa virginiensis) | Nymphalidae | 9     | 2–3         | Broad               | >10 (ex. pearly everlasting, cudweeds) |
| Silver-spotted skipper (Epargyreus clarus) | Suididae    | 9     | 2–3         | Broad               | >10 (ex. wisteria, locusts)         |
| Spicebush swallowtail (Papilio troilus) | Papilionidae | 4     | 2–3         | Broad               | ≤10 (primarily spicebush)           |
| Red-banded hairstreak (Calycopis cecrops) | Lycaenidae  | 3     | 2           | Restricted          | Unknown (sumacs, oaks possibly others) |
| Gray hairstreak (Strymon melinus)    | Lycaenidae  | 3     | 3–4         | Broad               | >10 (ex. mallows, legumes)          |
| Unknown skipper (Hesperinae)         | Hesperiidae | 3     | –           | –                   | –                                   |
| American snout (Libytheana carineta) | Libytheidae | 2     | 2           | Restricted          | ≤10 (hackberries)                  |
| Sachem (Atalopedes campestris)       | Hesperiidae | 2     | 2           | Broad               | ≤10 (Bermuda grass, crabgrass, St. Augustine grass) |
| Great spangled fritillary (Speyeria cybele) | Nymphalidae | 1     | 1           | Broad               | ≤10 (violets)                      |
| Common buckeye (Junonia coenia)      | Nymphalidae | 1     | 2–3         | Broad               | >10 (ex. plantains, vervains)       |
| Pearl crescent (Phyciodes tharos)    | Hesperiidae | 1     | 3–4         | Broad               | ≤10 (asters)                        |
| Common/White Checkered Skipper (Pyrgus communis/albescens) | Hesperiidae | 1     | 3           | Broad               | >10 (ex. mallows, velvetleaf, hollyhock) |
| Hobomok skipper (Poanes hobomok)      | Hesperiidae | 1     | 1           | Restricted          | ≤10 (panic grass, blue grasses)     |

*Previously restricted; introduction of invasive host plants has expanded the variety habitat types where this species can be found (Cech and Tudor 2005)
Table 3 Models ranked by $\Delta AIC_c$ explaining butterfly species richness in urban community gardens of New York City. Reported statistics include $\Delta AIC_c$, $W_i$ (Akaike weight), AICc, RSS (residual sum of squares), $K$ (number of parameters including the constant and the estimate of residual variance) and Adj. $R^2$ (adjusted $R^2$). The sample size for all models was $n=18$. Regression equations with standardized coefficients for included variables are shown for models with >5% of the support based on Akaike weights [$W_i$] (shaded models). See Methods for descriptions of all model variables.

| Model Parameters | $\Delta AIC_c$ | $W_i$ | AICc | RSS | $K$ | Adj. $R^2$ |
|------------------|----------------|-------|------|-----|-----|-------------|
| Floral area + sunlight | log$_{10}(y) = 0.527 *$floral area + 0.387*sunlight$^3$ | 0.00 | 59% | -67.07 | 0.222 | 4 | 0.601 |
| Floral area | log$_{10}(y) = 0.736 *$floral area | 1.39 | 30% | -66.68 | 0.289 | 3 | 0.513 |
| Floral area + sunlight + garden age | log$_{10}(y) = 0.533 *$floral area + 0.356*sunlight$^3$ - 0.041*garden age | 3.55 | 10% | -64.52 | 0.217 | 5 | 0.499 |
| Floral area + sunlight + garden age + total garden area | 8.18 | 1% | -59.89 | 0.217 | 6 | 0.549 |
| Floral area + sunlight + garden area + total garden area + building units (500m) | 13.60 | <1% | -54.47 | 0.215 | 7 | 0.515 |
| Floral area + sunlight + garden area + total garden area + building units (500m) + presence of wild/unmanaged areas | 16.80 | <1% | -51.27 | 0.176 | 8 | 0.524 |
| All variables except garden canopy cover, gardenH, green space (500m), building units (200m) and green space (200m) | 25.30 | <1% | -42.77 | 0.176 | 9 | 0.524 |
| All variables except garden canopy cover, gardenH, green space (500m) and building units (200m) | 34.91 | <1% | -33.16 | 0.164 | 10 | 0.508 |
| All variables except garden canopy cover, gardenH, and green space (500m) | 47.59 | <1% | -20.48 | 0.147 | 11 | 0.502 |
| All variables except garden canopy cover and gardenH | 67.05 | <1% | -1.02 | 0.140 | 12 | 0.459 |
| All variables except garden canopy cover | 96.87 | <1% | 28.79 | 0.134 | 13 | 0.396 |
| Global (all 12 variables) | 145.45 | <1% | 77.38 | 0.117 | 14 | 0.367 |

Table 4 Models ranked by $\Delta AIC_c$ explaining bee species richness in urban community gardens of New York City. Reported statistics include $\Delta AIC_c$, $W_i$ (Akaike weight), AICc, RSS (residual sum of squares), $K$ (number of parameters including the constant and the estimate of residual variance) and Adj. $R^2$ (adjusted $R^2$). The sample size for all models was $n=18$. Regression equations with standardized coefficients for included variables are shown for models with >5% of the support based on Akaike weights [$W_i$] (shaded models). See Methods for descriptions of all model variables.

| Model Parameters | $\Delta AIC_c$ | $W_i$ | AICc | RSS | $K$ | Adj. $R^2$ |
|------------------|----------------|-------|------|-----|-----|-------------|
| Total garden area + floral area + sunlight | log$_{10}(y) = 0.00 + 0.437*$total garden area + 0.294*$floral area + 0.310*$sunlight$^3$ | 0.00 | 35% | -62.37 | 0.245 | 5 | 0.678 |
| Total garden area + floral area | log$_{10}(y) = 0.00 + 0.455*$total garden area + 0.449*$floral area | 0.18 | 32% | -62.18 | 0.307 | 4 | 0.622 |
| Total garden area + floral area + sunlight + building units (500m) + canopy cover | log$_{10}(y) = 0.093*$total garden area - 0.567*$floral area + 0.223*$sunlight$^3$ - 0.322*$log$_{10}$(building units (500m)) + 0.324*canopy cover$^2$ | 2.12 | 12% | -60.24 | 0.156 | 7 | 0.760 |
| Total garden area | log$_{10}(y) = 0.739*$total garden area | 2.41 | 11% | -59.95 | 0.419 | 3 | 0.517 |
| Total garden area + floral area + sunlight + building units (500m) | log$_{10}(y) = 0.338*$total garden area - 0.320*$floral area + 0.239*$sunlight$^3$ - 0.204*$log$_{10}$(building units (500m)) | 2.75 | 9% | -59.61 | 0.220 | 6 | 0.688 |
| Total garden area + floral area + sunlight + building units (500m) + canopy cover + wild/unmanaged area | 8.30 | 1% | -54.06 | 0.151 | 8 | 0.747 |
| All variables except building units (200m), green space (500m), VegH, GardenH and garden age | 16.80 | <1% | -45.57 | 0.151 | 9 | 0.722 |
| All variables except building units (200m), green space (500m), VegH, and GardenH | 26.63 | <1% | -35.73 | 0.142 | 10 | 0.710 |
| All variables except building units (200m) and green space (500m) | 58.27 | <1% | -4.09 | 0.118 | 12 | 0.689 |
| All variables except building units (200m), green space (500m), and VegH | 70.48 | <1% | 8.11 | 0.121 | 11 | 0.722 |
| All variables except building units (200m) | 87.85 | <1% | 25.48 | 0.111 | 13 | 0.658 |
| Global (all 12 variables) | 135.39 | <1% | 73.03 | 0.092 | 14 | 0.659 |
bee diversity have also been found for large parks of New York City (Giuliano et al. 2004), residential gardens of Berkeley (Frankie et al. 2005), grassland fragments in Boulder (Kearns and Oliveras 2009), and meadows habitats in the Greater Boston Area (Clark et al. 2007). We did not measure floral diversity, which has been shown to have a strong influence on pollinator diversity (Potts et al. 2004). All of the butterflies and most of the bees (89%) encountered in these gardens are generalist floral feeders as adults. Thus, it might be interesting to evaluate the influence of floral abundance relative to floral diversity in this or other urban settings characterized by generalist floral feeders.

The most highly supported models for both butterfly and bee species richness also included the availability of sunlight within gardens. In contrast to open meadow habitats or even single-family residential gardens, the gardens of this study were commonly obstructed by tall apartment buildings. Indeed, there were negative associations between sunlight availability and the number of building units surrounding gardens at both the 200 m ($r = -0.472$) and 500 m scales ($r = -0.458$). Shading due to buildings results in limited photosynthetic photon flux relative to shade from deciduous trees (Bell et al. 2000) and thus may more severely limit plant growth and/or opportunities for bees and butterflies to increase body temperature via passive basking in gardens. Although it is difficult to increase sunlight for ground-level urban gardens, there is increasing interest in using green roofs to augment habitat area for urban wildlife (Oberndorfer et al. 2007), including bees (Colla et al. 2009). Because sunlight is less limited on most rooftops, large, florally rich green roofs may provide important bee and butterfly habitat, especially in inner-city areas where ground-level habitats are often shaded by buildings.

While both bees and butterflies responded to sunlight and floral area, bees responded to several additional garden variables. Specifically, there was marginal support for a model of bee species richness including garden canopy cover, and bee species richness also responded positively to total garden area and the presence of wild/unmanaged areas in the garden (Fig. 2). The response of bees, but not butterflies, to these variables likely reflects general differences in the mobility and life history requirements of bees and butterflies. Specifically, many of the butterflies found in these gardens are highly mobile, and thus unlikely to remain in gardens for long periods. In contrast, bees forage for nectar and pollen within a fixed distance from a centrally located nest. Many of the bee species encountered in this study have limited foraging ranges (<500 m; body sizes of species listed in Matteson et al. 2008 applied to data provided in Greenleaf et al. 2007), suggesting that many of the bee species nest within or near gardens. Because most of the bee species encountered in our study gardens construct nests in hollow plant stems or in soil (Matteson et al. 2008), a larger total garden, or having more wild/unmanaged areas of the garden (where trampling/turning of soil and pruning of plant debris is limited) may better enable larva/pupa to develop (Buchmann and Nabhan 1997). In support of this idea, Smith et al. (2006) found bumble bee species richness to be negatively associated with intensity of garden management. None of the gardens of this study included bee or butterfly nesting blocks, which may be an additional method for increasing local species richness (Gaston et al. 2005).

Influence of landscape variables on insect richness

We initially hypothesized that surrounding green space would be especially influential in this setting, due to its limited availability (range of ~10–30% green space at the 500 m scale). Landscape green space, however, had little influence on bee or butterfly richness. This may have been because our measure of green space included closely mown lawns and
tree cover, which may be poor proxies for bee or butterfly larval, nesting or floral resources. Alternatively, it may be that landscape-sensitive species have already been locally extirpated from heavily developed neighborhoods of this study, as has been suggested for insect taxa in other cities (Kozlov 1996; Connor et al. 2003). For instance, despite sampling these gardens for 4 years, we encountered just five of the 16 bumble bee species known to inhabit southern New York state (John S. Ascher, personal communication), and similar reductions in bumble bees have been noted in San Francisco, USA (McFrederick and Lebuhn 2006) and Stockholm, Sweden (Ahrné et al. 2009). Overall, twice as many bee species were found in private gardens in less developed suburbs just 50 km north of our study sites (Fetridge et al. 2008). Similarly, we found just 24 of the 120 butterfly species reported to inhabit a 50-mile radius around New York City (Cech 1991) and most were seasonal or habitat generalists, or utilize host plants that are fairly common in the urban landscape (Table 2). In the Greater Boston Area, rare and specialist butterfly species exhibited the strongest decrease with increasing urbanization (Clark et al. 2007), suggesting that generalist species may be less responsive to limitations in landscape green space. Di Mauro et al. (2007), however, demonstrated that generalist butterflies also may decrease with urbanization (decreasing green space) in some garden types.

Collinge et al. (2003) and Kearns and Oliveras (2009) also failed to find a relationship between the proportion of the landscape that was developed and butterfly species richness or bee species richness. These authors, however, concluded that the degree of urbanization proximal to their study sites (grassland habitats located at the edge of urban development, outside of Boulder, Colorado) was insufficient to result in a measurable decrease in species richness. This suggests that traditional measures of landscape context (e.g. green space, semi-natural habitat) may have neutral effects both in highly developed areas (such as this study where there are few remaining landscape-sensitive species) as well as in relatively less developed areas at the outskirts of urban development (where few landscape-sensitive species may have been lost). Studies conducted across a wider range of urban development with local variables held close to constant (i.e., little variation in area or floral resources), may find a stronger impact of landscape green space (Winfree et al. 2007; Ahrné et al. 2009), which may exert a nonlinear influence on insect richness. For instance, Öckinger et al. 2009 found butterfly richness of urban parks to increase with green space in a system with 10–80% green space at the 1,000 m scale (in contrast this study, where there was a range of just ∼10–30% green space at the 500 m scale).

We found some support for the hypothesis that the number of building units surrounding gardens at the 500 m scale negatively influences bee species richness. The number of building units was not correlated with green space at the 500 m scale ($r=0.02$), indicating that there are mechanisms other than a reduction in green space involved in this relationship. The number of building units at the 500 m scale was, however, negatively correlated with garden sunlight availability ($r=-0.458$), suggesting that the effect may be due to shading of both the focal garden (as discussed above) and surrounding landscape. Another possibility is that floral resources are reduced, independent of our measure of “green space”, around large apartment buildings relative to single-family residences. In support of this idea, green space surrounding several large housing projects proximal to these gardens was largely composed of lawns interspersed with sycamore trees and Taxus bushes, and few floral resources (KCM, personal observation). This suggests that different types of urban housing development may have very different effects on bee or butterfly species richness. Specifically, bee and butterfly diversity might be optimized near low-density or single-family homes with large, florally rich gardens (Winfree et al. 2007; Fetridge et al. 2008; Frankie et al. 2009), but limited in high-density neighborhoods with few floral resources.
Conclusion

The results of this study indicate that sunlight and floral abundance are major factors limiting pollinator diversity in urban gardens located in heavily developed neighborhoods. The general patterns hold true in post hoc analyses of observed, rather than estimated, species richness and in analyses of specific-species groupings such as just native or "large-bodied" bees (KCM, unpublished data). This suggests that rooftop gardens and other "open" urban habitats might be managed to benefit local pollinator diversity by increasing floral area, even if seemingly "isolated" within heavily developed neighborhoods. Such actions may be difficult to implement, however, in heavily developed neighborhoods where sunlight is limited by the vertical extent of tall buildings. In addition, the maximum level of species richness may be restricted to the pool of species that linger within heavily developed neighborhoods. While we did not find the amount of green space in a 500 m radius to have a major influence in this setting, additional bee and butterfly species may only persist in neighborhoods with a green space threshold greater than the 10–30% range (at the 500 m radius scale) of this study. Therefore, larger scale efforts to implement floral plantings into areas of the urban landscape that receive full sun exposure (e.g. rooftop gardens, building terraces with southern exposure), in addition to increasing connectivity to species rich peri-urban areas (Snep et al. 2006), may be required to attract and benefit a greater diversity of bees and butterflies.

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