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Conservation biology

Does urbanization favour exotic bee species? Implications for the conservation of native bees in cities

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A growing body of research indicates that cities can support diverse bee communities. However, urbanization may disproportionately benefit exotic bees, potentially to the detriment of native species. We examined the influence of urbanization on exotic and native bees using two datasets from Michigan, USA. We found that urbanization positively influenced exotic but not native—bee abundance and richness, and that this association could not be explained by proximity to international ports of entry, prevalence of exotic flora or urban warming. We found a negative relationship between native and exotic bee abundance at sites with high total bee abundance, suggesting that exotic bees may negatively affect native bee populations. These effects were not driven by the numerically dominant exotic honeybee, but rather by other exotic bees. Our findings complicate the emerging paradigm of cities as key sites for pollinator conservation.

1. Introduction

Exotic species introductions have accelerated in recent decades alongside increases in international commerce [1–3]. Exotic species can cause biodiversity loss [4,5], biotic homogenization [6] and changes to ecosystem function [4]. Among the most widely established exotic species is the European honeybee— *Apis mellifera* Linn.—which has been introduced worldwide for pollination and honey production [7,8]. Other bee species have also been introduced beyond their native ranges, both purposefully and accidentally [9,10]. While exotic bees can provide important pollination services, they may also compete with native species for resources or spread pathogens to native species [6,10–13]. This is concerning given that native bee populations are declining in some areas [14–16]. To date, studies of the effects of exotic bees on native bees have focused primarily on eusocial exotics, and less so on solitary exotics (but see [10]).

Little is known about the ecological determinants of exotic bee colonization and spread. However, there are several ways urbanization could facilitate exotic bee establishment. First, exotic bee introductions often occur accidentally via international commerce [10]. As trading nodes with international ports of entry (IPOE), cities may be a frequent introduction point for exotic bees, thus increasing establishment via high propagule pressure [17]. Second, most exotic bees are cavity-nesting [9], presumably because this nesting strategy facilitates longdistance transport and introduction via the movement of occupied nest substrate [10]. Multiple studies have found increased prevalence of cavity-nesting bees with urbanization [18–20], likely because the built environment provides suitable nest

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sites for these species [18]. Thus, exotic bees may thrive in urban areas due to their nesting behaviour. Third, urban warming [21] may favour a suite of bee species different from that present in less urbanized areas [22]. This could allow for increased dominance of exotic species whose thermal tolerance better matches warmer urban conditions (e.g. [23]). Finally, exotic bees often show foraging preference for exotic flowering plants [11], and cities can be richer in exotic plants than their surroundings [24–26]. However, the abundance of managed species in cities, such as honeybees, is likely more related to management intensity, rather than preferred resource availability. Nonetheless, cities might support non-honeybee exotics via increased abundance of preferred floral resources.

Despite evidence that cities support exotic bees [27,28], limited research has shown that urbanization positively influences exotic bee populations (but see [29]), or explored the underlying drivers of this association. Recently, several authors have suggested that cities may be important sites for bee conservation [30–32]. Given this emerging paradigm, and the fact that cities are predicted to continue expanding worldwide [33], we urgently need to understand how urbanization influences the establishment and spread of exotic bees, and associated impacts on native bees.

In this study, we address the following questions:

- (1) Are exotic bee abundance and richness positively associated with increased urbanization?
- (2) Is the relationship between exotic bees and urbanization mediated by (i) proximity to registered IPOE, (ii) prevalence of exotic plants, (iii) urban warming and/or (iv) nesting strategy?
- (3) Do different exotic bee species demonstrate similar responses to urbanization?
- (4) Is there evidence for negative effects of exotic bees on native bees?

2. Material and methods

Our dataset was compiled from two field studies conducted within southeast Michigan, USA [34,35]. Bees were sampled at 41 farms and community gardens, with 26 sites sampled in 2014 and 15 in 2017. Monthly from June to August, we sampled bees using bowl traps and netting and conducted floral surveys. Data loggers at each site collected hourly temperature data. For details of survey methods, see electronic supplementary material.

We assessed urbanization at each site as the proportion of developed land within each of four concentric buffers, using the National Land Cover 2011 database [36] following the approach described in [37] (see electronic supplementary material, table S1). Among the four radii, 500 m was the most predictive for exotic bee abundance and richness and was used in all subsequent analyses. We also measured the distance from each site to the nearest registered IPOE (electronic supplementary material).

Statistical analyses were conducted in R v. 3.5.1 [38]. Because honeybees occur mainly in managed colonies in the study region, we considered them separately from other exotic bees in all analyses. Since honeybees were treated differently in netting protocols between studies (electronic supplementary material), we used only trap data in analyses that included honeybees.

We tested the effect of urbanization on per-site exotic and native bee richness and abundance using generalized linear models (GLMs) fit with a Poisson distribution (negative binomial if data were overdispersed) and log-link function. Study year and proximity to IPOE were included as predictors in these models. We evaluated the relationship between urbanization and the abundance of four widespread non-honeybee exotics (species found at \geq 10 sites) using the same framework. We were interested in whether urbanization disproportionately favoured exotics, so we evaluated proportional abundance and richness of exotic bees and exotic flowering plants. To do so, we included total abundance or richness as an offset in the model. We evaluated the effects of additional putative drivers using likelihood ratio tests and AICc.

To assess the effect of exotic floral resources on the bee fauna, we considered each monthly observation separately and used generalized linear mixed models (GLMMs) with site as a random effect and proportional richness or cover of exotic plants as a fixed effect. Total floral richness, urbanization and year were also included as predictors in these models (electronic supplementary material).

To test whether nesting strategy could account for the correlation between exotic bees and urbanization, we assessed the responses of native cavity-nesting and non-cavity-nesting bee abundance and richness to urbanization using GLMs.

We evaluated the effect of exotic bee abundance on native bee abundance using GLMMs with native bee abundance as the response variable; exotic bee abundance, urbanization and floral richness as fixed effects; and site as a random effect. Effects of exotic bees on natives are likely to be density-dependent [39], with stronger negative effects when total population density is higher [40]. Thus, we assessed the relationship between native and exotic bee abundance separately at sites with high versus low bee abundance (electronic supplementary material). To test the robustness of our findings, we considered a range of cutoffs for separating high-abundance from low-abundance sites [25-50 bees/sampling period for comparing non-honeybee exotics and natives; 10-40 bees/sampling period for comparing honeybees and natives (smaller numbers because only trap data are included; electronic supplementary material)], and, for each cut-off, fit models for both high- and low-abundance sites.

3. Results

We found 14 exotic bee species, comprising 20% of all bee species collected [41]. The percentage of exotic bees collected per site ranged from 0 to 40% (mean \pm s.d. = 16 \pm 14%; figure 1), while exotic bee species richness ranged from 0 to 9 (mean \pm s.d. = 3 \pm 2) per site (electronic supplementary material, table S2). The European honeybee—the only actively managed bee species in the study area—was the most abundant exotic species, comprising 58% of all exotics, and occurring at 18 of 41 sites (30 of 41 when including netting data). Non-honeybee exotics belonged to four families: Colletidae (two species, 50% of individuals), Megachilidae (eight species, 34%), Andrenidae (one species, 11%) and Halictidae (two species, 5%) (electronic supplementary material, figures S1 and S2). Proportional abundance of exotic bees was higher in 2017 (29%) than 2014 (14%).

The absolute and proportional abundance and richness of exotic bees significantly increased with urbanization, while native and total bee abundance and richness were unaffected (figure 2a and table 1). Abundance of each widespread wild exotic showed a qualitatively similar response to urbanization, indicating that the overall relationship between urbanization and exotic bee abundance was not driven by a single species (table 1; electronic supplementary material, figure S3). However, honeybees showed no response to urbanization (figure 2a).

Cavity-nesting bees comprised 6% of native bees and 57% of non-honeybee exotics. More urbanized sites supported



Figure 1. Map of study sites with proportional abundance of native and exotic bees. Greyscale represents degree of urbanization. Geographical location of nearby sites is offset to prevent overlap of pie charts. (Online version in colour.)

more exotic and native cavity-nesting bees, although this relationship was much stronger for exotics. While exotic cavity-nester richness was positively related to urbanization, native cavity-nester richness was not (table 1).

Urbanization was positively related to proximity to the nearest IPOE (LM: $R^2 = 0.53$, $F_{1,39} = 45.8$, p < 0.001). When both proximity to IPOE and urbanization were included as model predictors, only urbanization had a significant effect on exotic bee richness and abundance and including both measures increased AICc. However, proportional non-honeybee exotic abundance was negatively related to proximity to IPOE (table 1). Similarly, urbanization and minimum temperature were positively related (LM: $R^2 = 0.32$, $F_{1,39} = 17.9$, p < 0.001), but minimum temperature had no independent effect on exotic bee richness or abundance (electronic supplementary material, table S3).

The positive relationships between urbanization and exotic bee richness and abundance were not mediated by exotic plants. Neither raw nor proportional richness or cover of exotic plants were correlated with urbanization (electronic supplementary material, table S5). Moreover, while exotic bee abundance and richness were positively related to total floral resource availability, they were not influenced by proportional richness or cover of exotic plants (table 1).

Exotic bee abundance had no effect on native bee abundance $(\beta = 0.01 \pm 0.02, p = 0.39)$. However, at high-abundance sites, there was a significant negative relationship between non-honeybee exotic and native bee abundance not seen at low-abundance sites (figure 2*b*; electronic supplementary material, table S5). The relationship between honeybee abundance and native bee abundance was qualitatively similar; however, the negative relationship between honeybees and

native bees was generated by a single observation (figure 2*c*; electronic supplementary material, table S6) and should be interpreted with caution. The negative relationship between native and wild exotic bee abundance at high-abundance sites was not due to correlation between exotic abundance and any other measured driver of native bee abundance (electronic supplementary material, table S6).

4. Discussion

Urbanization alters the composition of biotic communities by creating a matrix of habitats distinct from natural ecosystems [42,43]. In this study, urbanization correlated with increased prevalence of exotic bees, via increases in exotic bee abundance and richness rather than declines in native bees. The association between urbanization and exotic bees was not mediated by exotic floral resource availability, proximity to IPOE or urban warming.

The lack of relationship between exotic bees and exotic plant prevalence contradicts other studies suggesting that exotic bees preferentially visit exotic plants [44–46]. However, 96% of exotic bees we collected were from generalist species; their success in their introduced range may derive from the ability to feed on a wide range of plants [47]. Because we did not assess bee diet, our findings do not demonstrate that exotic bees do not prefer exotic plants. They do, however, indicate that the success of exotic bees in cities is not due to increased abundance of exotic floral resources.

Most of the exotic bees we collected nest in cavities; the additional nesting substrate provided by urbanization may facilitate these species [18]. Indeed, the abundance of native cavity-nesting bees also increased with urbanization, though



Figure 2. (*a*) Relationship between urbanization and exotic bee proportional abundance. (*b*,*c*) Relationship between native and exotic bee abundance for (*b*) non-honeybee exotics and (*c*) honeybees. Grey points represent low-abundance sites; black points represent high-abundance sites (in (*b*), high abundance: \geq 40 bees collected; in (*c*) \geq 20 bees). Outlined point in (*c*) indicates outlier. Trendlines derived from GLMs. Significance codes: ^{n.s.}*p* > 0.05; **p* < 0.05; **p* < 0.01. (Online version in colour.)

this response was weaker than that of exotic bees, suggesting that nesting preferences alone cannot account for exotic bee success in cities. In sum, the increased prevalence of exotic bees in cities is largely attributable to trait-matching between exotics and urban environments (e.g. cavity-nesting habit). We found no evidence that proximity to IPOE increases exotic bee abundance or richness, but the limited scope of our study does not allow us to definitively evaluate the role of propagule pressure in exotic bee success in cities.

The observed negative correlation between native and exotic bees at sites supporting high total bee abundance but not at low-abundance sites—suggests density-dependent effects of exotics on natives. Negative effects of exotic bees on natives may be due to competition for food or nest sites [9,48,49], or apparent competition mediated by shared pathogens [9,50]. Intriguingly, we found that wild exotic bee abundance accounted for more deviance in native bee abundance than did honeybee abundance. The strength of the relationship between wild exotic and native bee abundance is surprising, given that (1) studies measuring the effect of exotic honeybees on native bees rarely demonstrate population-level consequences [12] and (2) effects of non-eusocial exotic bees on native bees are understudied. Alternatively, environmental filtering, operating differently on native versus exotic bees, may be responsible for the observed relationship. Conclusively determining whether the observed relationship indicates that wild exotic bees exert more influence than honeybees on natives, or results from collinearity with some unassessed driver of bee abundance requires further study.

Recent findings that cities can maintain diverse bee communities [31,32,37,51] has increased interest in cities as targets for bee conservation [30–32]. While promoting beefriendly management of urban land is vital to protecting pollinators, this study highlights the need to think critically Table 1. Effects of potential drivers of native and exotic bee abundance and richness. Dash indicates that the predictor was not included in the best model (determined by AICc). %DE, per cent of null deviance explained; DF, degrees of freedom; $\Delta AICc$, difference in AICc between full model and the best-fitting model, which omits non-significant predictors. Significance codes are indicated by asterisk. Italic text represents significant predictors at p < 0.05.

	predictors of intere	est			covariates				
	urbanization	proximity to international port	proportional richness of exotic plants	proportional bloom cover of exotic	total plant				
response variable	$(m{eta}\pm {f s}.{f e}.)$	of entry ($m{eta} \pm$ s.e.)	$(m{eta} \pm {\sf s.e.})$	plants ($meta \pm$ s.e.)	richness ($m{eta}\pm$ s.e.)	year ($meta \pm {\sf s.e.}$)	%DE	Df	ΔAICc
(a) GLMs									
abundance									
non-honeybee exotic	0.80 ± 0.21***	−0.11 ± 0.21	I	I	Ι	$1.03 \pm 0.28^{***}$	52.2	37	2.35
proportion non-honeybee exotic	0.98 ± 0.09***	$-0.34 \pm 0.09^{***}$	1	Ι	Ι	1	63.6	38	0.00
honeybee	0.10 ± 0.42	−0.31 ± 0.41	1	Ι	Ι	1	0.0	38	3.52
Hylaeus hyalinatus	$0.70 \pm 0.29^{*}$	0.53 ± 0.32	1	Ι	Ι	1	46.8	38	0.34
H. leptocephalus	2.07 ± 0.52***	-0.84 ± 0.51	1	I	1	1	55.6	38	0.35
Anthidium manicatum	$1.19 \pm 0.59^{*}$	−0.07 ± 0.62	1	I		1	24.8	38	2.45
Megachile rotundata	1.51 ± 0.45***	−0.61 ± 0.45	I	I	1		38.6	38	0.69
exotic cavity-nesting	0.81 ± 0.21***	0.05 ± 0.21	1	Ι	I	$1.66 \pm 0.27^{***}$	68.4	37	2.55
native overall	-0.35 ± 0.25	1	1	I	I	0.37 ± 0.14**	17.6	38	0.58
native cavity-nesting	0.27 ± 0.14	I	1	Ι	Ι	2.92 ± 0.33***	73.7	38	0.35
native non-cavity-nesting	-0.01 ± 0.07	I	1	I		0.30 ± 0.14*	9.7	38	2.42
richness									
exotic	0.42 ± 0.16*	<i>−</i> 0.11 ± 0.16	1	I	1	<i>0.78</i> ± <i>0.21</i> ***	42.4	37	2.01
proportion exotic	0.49 ± 0.16**	0.19 ± 0.16	1	I		1	30.5	38	0.95
exotic cavity-nesting	$0.59 \pm 0.22^{**}$	−0.10 ± 0.21	1	I		1.26 ± 0.27***	55.4	37	2.24
native overall	-0.18 ± 0.16	I	1	Ι	I	$0.34 \pm 0.09^{***}$	27.6	38	1.24
native cavity-nesting	0.11 ± 0.11	1			1	$2.24 \pm 0.30^{***}$	70.0	38	1.30
native non-cavity-nesting	-0.02 ± 0.04	1	1	I		$0.26 \pm 0.09^{**}$	17.6	38	2.20
total	-0.04 ± 0.16	1	1	Γ	I	0.40 ± 0.09	33.8	38	2.39
(b) GLMMs									
non-honeybee exotic abundance	$0.66 \pm 0.13^{***}$	1	0.11 ± 0.14	0.01 ± 0.11	$0.39 \pm 0.08^{***}$	$0.76 \pm 0.29^{**}$	10.4	106	3.59
H. hyalinatus abundance	$1.02 \pm 0.20^{***}$	-	-0.23 ± 0.24	0.09 ± 0.21	$0.59 \pm 0.13^{***}$	1	15.5	107	3.48
H. leptocephalus abundance	$1.59 \pm 0.40^{***}$		0.13 土 0.40	-0.02 ± 0.31	1		9.8	108	4.22
A. manicatum abundance	0.84 ± 0.42*		−0.27 ± 0.50	0.34 ± 0.39	$0.75 \pm 0.28^{**}$		10.8	107	3.60
<i>M. rotundata</i> abundance	1.01 ± 0.32**		0.08 ± 0.41	0.15 ± 0.35	1		7.4	108	3.89
non-honeybee exotic richness	0.39 ± 0.09***		0.11 ± 0.15	−0.23 ± 0.13	$0.25 \pm 0.10^{*}$	0.71 ± 0.21***	11.6	106	1.30
$p < 0.05; \ ^{**}p < 0.01; \ ^{***}p < 0.001.$									

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about the bee communities supported by urban environments. While urbanization can increase bee beta-diversity by supporting a different suite of species from those found outside cities, this may come at a cost to native species. Our research suggests that, globally, urbanization may homogenize bee communities by increasing the dominance of a small number of cosmopolitan, synanthropic species.

Ethics. No permits or ethics committee approvals were required for this research. We received permission from all landowners and managers to conduct research.

Data accessibility. Data and R scripts are available from the Dryad Digital Repository: https://dx.doi.org/10.5061/dryad.9kf08rj [41].

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Authors' contributions. All authors jointly conceived of this paper and participated in data collection. G.F. conducted statistical analyses; C.J.W. conducted geographical analyses; G.F. and C.J.W. drafted the manuscript. All authors revised and approved the manuscript and are accountable for all aspects of the work.

Competing interests. We declare we have no competing interests.

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