Effects of urbanization on Neotropical wasp and bee assemblages in a Brazilian metropolis

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Abstract

The rapid growth of cities in many parts of the world has stimulated an increasing number of ecological studies of urban environments. Here, 12 study sites in the city of Belo Horizonte (MG), southeastern Brazil, were compared to analyze the effects of habitat changes related to urbanization on the species richness and abundance of native wasps and bees. Two spatial scales were considered: large portions of the urban landscape (entire sites) and small areas within these sites (public squares). Overall, the abundance of advanced eusocial bees (i.e. stingless bees), which were the dominant species, was directly affected by the loss of vegetation cover and the increase of buildings associated with urbanization. The magnitude of this effect varied according to the nesting habits of each species. The loss of vegetation cover associated with urbanization also had a negative effect on the abundance and species richness of advanced eusocial wasps. Generalist species of bees and wasps, such as Trigona spinipes and Polybia occidentalis, were very abundant and not sensitive to the habitat changes related to urbanization. Advanced eusocial bees also responded to small-scale habitat changes (size and vegetation cover of public squares). No relation was found between the solitary and the primitive eusocial wasps and bees and the habitat changes considered here. Conservation strategies in urban environments need to consider different spatial scales in order to maintain or enhance the local diversity of wasps and bees.

Keywords: Aculeata; Stingless bees; Habitat structure; Public squares

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1. Introduction

Urban environments are generally characterized as areas with a high density of buildings exposed to constant and intense human activity (McIntyre et al., 2001). Urbanization, the process by which these environments are generated, can be summarized as a local increase in the density of inhabitants coupled with increased per capita energy consumption and extensive modifications of the environment (including the microclimate) (Gilbert, 1991; Vitousek et al., 1997). This process generates unstable ecosystems that depend on large inputs of energy, and where great amounts of waste materials are accumulated (Stearns, 1970; McDonnell and Pickett, 1990). Despite covering a small fraction of the Earth’s surface (<5%), urban environments have a widespread influence...
on surrounding ecosystems (Vitousek et al., 1997; Goudie, 2000). Half of the world’s population currently lives in urban areas. This proportion is expected to increase to 60% in the year 2030 (United Nations, 1999). In Brazil, approximately 55% of the population was living in cities in 1970. Today, almost 80% of the population is living in urban areas (IBGE, 2001). This fast growth of cities is promoting an increasing interest in urban ecosystems and in the impact of urbanization on distinct biotas (Gilbert, 1991; Rebele, 1994; Grimm, 1997; Vitousek et al., 1997). Ecological studies of such ecosystems are essential to reduce local and regional impacts of urbanization (Gilbert, 1991; Bedê et al., 1997; Niemelä, 1999).

Different approaches have been used to study the fauna of urban environments (Moore, 1979; McDonnell and Pickett, 1990; Niemelä, 1999). The theory of island biogeography, for instance, has been the foundation for many studies of insect and bird richness in “urban islands” (e.g. parks and public squares) (reviews in Davis and Glick, 1978; McIntyre, 2000). Despite being spatially heterogeneous, urban environments are structurally well defined (Turner, 1989). Generally, each part (or structural component) of the urban landscape presents clear limits (Wiens et al., 1993; Grimm, 1997). Public squares, for example, are well-defined habitat portions surrounded by a matrix of buildings equally well defined. Although “urban islands” may be easily perceived, this biogeographic approach has some limitations since the permeability of the urban matrix may be highly variable (Niemelä, 1999). Species with great dispersal ability, for example, may pass through the matrix; consequently they can use a portion of the urban habitat larger than a specific “island”. For these reasons different spatial scales have to be analyzed to study the effects of urbanization on a particular group of species (Hostetler, 1999). Additionally, it has to be considered that urbanization is a scale-dependent process. Urban planning and development are done in an hierarchical way (“from the mayor to the gardener”), and consequently, the effects of urbanization can not be fully appreciated by considering only one spatial scale (Savard et al., 2000).

Despite the ubiquity of insects, there are few studies especially concerned with the effects of urbanization on non-pest species, particularly in the Neotropical region (Laroca et al., 1982; Davis, 1982; Ruszczyk 1986a,b,c,d, 1996; Diefenbach and Becker, 1992; McIntyre, 2000). Earlier studies found a relatively high insect diversity in some cities (e.g. London and Rome), suggesting that these habitats may not be as restrictive as expected (Owen and Owen, 1975; Zapparoli, 1997). In addition, it has been observed that species richness and abundance can greatly vary within each city, indicating that different levels of urbanization have distinct effects on the local insect fauna (Owen and Owen, 1975; Frankie and Koehler, 1978; Zapparoli, 1997; McIntyre, 2000; McIntyre et al., 2001).

Information about the responses of wasps and bees to urbanization is important for a number of reasons. Many wasp species, especially the eusocial ones, are key predators in tropical ecosystems (LaSalle and Gauld, 1993; Raw, 1998b). Bees are the most important pollinators of angiosperms and consequently they are also key species in many terrestrial ecosystems (Neff and Simpson, 1993). Wasps and bees are sensitive to variations in abiotic conditions (e.g. temperature, luminosity and moisture), which may be related to changes in the urbanization level (Genise, 1981; Roubik, 1992; Morgan and Jeanne, 1992). In addition, wasps and bees are frequently found in urban environments, efficiently occupying different microhabitats (e.g. walls, roofs, ceilings, etc.) (Nogueira-Neto, 1970; Fowler, 1983; Martins and Pimenta, 1993; Knoll et al., 1994; West-Eberhard et al., 1995; Saure, 1996; Raw, 1998a). Despite their ecological importance and their potential as indicators of environmental conditions (Brown, 1991), few authors have analyzed the effects of urbanization on wasp and bee assemblages. The literature suggests considerable variation in the responses of these insects to urbanization (Skibinska, 1986; Gayubo et al., 1987; Gayubo and Torres, 1989, 1990, 1991; Torres and Gayubo, 1989; Saure, 1996). In Brazil, previous studies restricted to bees have found a great variation in species sensitivity to urbanization and a considerable reduction of diversity in highly urbanized areas (Laroca et al., 1982; Bortoli and Laroca, 1997; Knoll et al., 1994).

It is generally assumed that urbanization (not considering deserts), reduces habitat complexity, mostly by reducing natural vegetation cover. Consequently, it may be predicted that less urbanized areas are structurally more complex. For these reasons, it can also be
predicted that less urbanized areas support a greater abundance and species richness of wasps and bees. Our main objective in this paper is to examine these predictions. Additionally, we have attempted to analyze the effects of urbanization at a smaller spatial scale, focusing on the microhabitats of public squares. Based on island biogeography theory, it was assumed that public squares surrounded by an urbanized matrix could be considered islands with different levels of isolation according to the structural complexity of their surroundings. Hence, the following predictions were also examined: (1) larger squares support a higher abundance and richness of wasps and bees, and (2) squares with more complex surroundings support a higher abundance and richness of wasps and bees.

2. Material and methods

2.1. Study area

Two regions of the city of Belo Horizonte (19°52’S, 43°58’W) in the state of Minas Gerais, southeastern Brazil were considered in the present study. Founded in 1897, Belo Horizonte was especially planned to be the capital of the state. Today the metropolitan area covers an area of approximately 330.9 km² and has 6746.79 inhabitants per km² (IBGE, 2001). In the 1980s, Belo Horizonte was considered to be the fastest growing capital in Brazil (da Silva, 1998). The two regions selected within the city have distinct urbanization histories. The Centro-Sul region is the oldest part of Belo Horizonte and has been intensively urbanized since the foundation of the capital. Although buildings predominate in the region, mature plant communities can still be found in intensively managed gardens of some old houses and institutional buildings. The urban development of the Pampulha region began in the 1940s, with the creation of the Oscar Niemeyer’s Pampulha Complex by the side of an artificial lake, and since then has never stopped (da Silva, 1998). Several patches of native vegetation can still be found in the region, particularly in vacant lots and some large private gardens subject to infrequent management.

A combination of demographic and socio-economic data were used to select three areas with different levels of urbanization within the two regions of Belo Horizonte (see Hostetler, 1999; McIntyre et al., 2000):

- L (West Pampulha): low level of urbanization (9.6 inhabitants per hectare and less than one non-residential building per hectare);
- M (East Pampulha): medium level of urbanization (48.7 inhabitants per hectare and eight non-residential buildings per hectare);
- H (Centro): high level of urbanization (98 inhabitants per hectare and 48 non-residential buildings per hectare).

Twelve sites were randomly selected for insect sampling: four in the L area, three in the M area and five in the H area. In each site one public square and two adjacent streets were sampled. These public squares are intensively managed public gardens of exotic and native ornamental plants. Since each square has a particular structure they provide an ideal situation to analyze the effects of small-scale habitat changes.

2.2. Characterization of the sites

Aerial photographs of the city of Belo Horizonte (scale 1:1000; source Prodabel, 2001) were used to quantify the habitat features of the twelve study sites. For each site, twelve photos were analyzed, each photo corresponding to an area of approximately 15,000 m². Each photo was analyzed using a grid of 560 square cells (each cell corresponding to 26 m²). The number of cells in each class was used to estimate the percentages of vegetation cover, impervious surfaces and bare soil (Michener et al., 1958; Brockmann, 1979). The coefficient of variation (CV) of the number of cells in each class (i.e. the variation within each set of 12 photos) was used as a simplified measure of spatial variation for the three land cover types.

Considering that large vegetated areas such as public parks and urban forest fragments may be sources of colonizing species, the distance between these large green areas and each study site was also included in the analysis. The minor distance between each site and the nearest vegetated area of more than 100 ha was...
measured directly from a map of the city of Belo Horizonte (source: Prodabel, 1997; scale 1:5000).

2.3. Characterization of the public squares

The plant species found in each square were identified in loco, with the supervision of botanists of the Belo Horizonte County Public Parks and Gardens Department, and thus the total richness of plant species in each public square was obtained.

Plant volume of herbs and shrubs (excluding grasses) was used as an estimate of live non-arboreal plant biomass. Volume was estimated using the height and the perimeter of each plant, which were measured using a metric strip and a compass. Perimeter measures were used to estimate the area occupied by each plant species using the program AutoCAD14.0. Considering that most plants in the squares were constantly managed, their forms were mostly regular (generally cubic or cylindrical); therefore, they could be easily measured. The coefficient of variation for the plant volumes was used as an estimate of the spatial heterogeneity of the non-arboreal plants in each square. The same method used to estimate the area occupied by each plant species was used to estimate the area of bare soil in each square.

An estimation of trunk volume was used as a measure of arboreal biomass for each tree species in the squares. Trunk volume was calculated by multiplying the tree height (visually estimated) by the trunk diam-
eter at chest height (Bonham, 1989). The coefficient of variation of the trunk volumes was used as an indirect measure of the spatial heterogeneity of the trees in each square.

The total area and the percentage of each ground cover type in each square were also measured with the program AutoCAD 14.0, using digitized maps of each square (source Prodabel, 2000; scale 1:2000) (Fig. 1).

2.4. Sampling of wasps and bees

Samples were collected intensively within a short period of time to minimize possible effects of climatic variations. Samples were taken on sunny days between 2 April 2000 and 23 July 2000, from 9 a.m. to 4 p.m., the period of more intense activity for most wasp and bee species (Sakagami et al., 1967; Roubik, 1992; Macedo and Martins, 1999; Williams et al., 2001). During each sampling day, three sites were sampled (i.e. three squares and six streets) with an interval of 15–20 min between each place. In total, all sites were sampled 12 times (four times in each of the following periods: 9 a.m.–11 a.m.; 11 a.m.–2 p.m.; 2 p.m.–4 p.m.). The sampling schedule for each day was planned using a table of random numbers.

Native wasps and bees seen flying, or on plants, up to a maximum height of 4 m were collected using an entomological hand net. The total area of each square was covered once. In the surrounding streets, the sidewalks were sampled once within a 300 m radius from the center of the square. To avoid over sampling a particular place, the maximum time spent at each point (e.g. flowering plant) was 3 min (see Sakagami et al., 1967; Heithaus, 1979). Collected wasps and bees were identified and placed in the entomological reference collection of the Laboratorio de Ecologia e Comportamento de Insetos of the Universidade Federal de Minas Gerais.

2.5. Data analysis

The occurrence of forager recruitment on the advanced eusocial species of bees and the large number of individuals found in the colonies of advanced eusocial wasps and bees may greatly increase the chances of capturing these insects using the method described above (Laroca et al., 1982). For this reason, the subsequent analyses were done considering two basic groups: advanced eusocial species and non-advanced eusocial species.

A Jackknife procedure was needed to estimate the species richness, as the number of sampled insects was not very large (Heltshle and Forrester, 1985; Palmer, 1990). The number of species collected each day in each site was used as a value for the Jackknife procedure. Confidence limits (at 95%) were calculated for each mean value of richness estimated to allow further comparisons between sites (Zar, 1999).

A Canonical Discriminant Analysis (CDA) was used to test whether the 12 study sites previously separated in groups according to their urbanization level, could also be differentiated using the measured habitat features for each site (Huberty, 1994; Jongman et al., 1995). CDA was also used to test whether these groups could be separated using the abundance of the species collected at each site. In the latter CDA, locally rare species, which occurred in less than 20% of the sampled places, were not considered (90% of these species were found three times or less).

A simple linear regression between the significant canonical roots extracted in the previously described CDAs was used to test if the abundance of wasps and bees was affected by the habitat changes related to urbanization. Another simple linear regression between the estimated values of species richness and the significant discriminant root obtained with the habitat features, was used to test if the richness of wasps and bees was affected by the habitat changes related to urbanization.

The variables that characterized the public squares and their surrounding were used to generate eigenvectors in a principal component analysis (PCA). The resulting main factors were used as explanatory variables in simple linear regression models, which tested if insect richness and abundance per square responded to the most important habitat features of each square (e.g. August, 1983; Therriault and Kolasa, 2000). Since the areas M and H had no significant structural differences, which would have interfered in the small-scale analyses (PCAs), only the squares in these two areas were used in these analyses. The small number of squares considered precluded a more direct analysis such as Canonical Correspondence Analysis. However, to avoid rejecting important variables (a common mistake in indirect analysis) the first five factors obtained in the PCAs.
were considered (together representing 90% or more of the total variance in the data) (Valentin, 2000).

In all the analyses percent land-cover values were arcsine transformed and area, distance and abundance values were log transformed (Krebs, 1999).

3. Results

3.1. Abundance and species richness of wasps and bees

Overall, 6268 individuals from 110 species, belonging to eleven families in the order Hymenoptera were captured (Figs. 2 and 3). The most abundant families were Apidae \( (n = 5150) \) and Vespidae \( (n = 794) \). Sixty-nine species were found in the less urbanized area (L), 47 in the area with an intermediate level of urbanization (M) and 56 in the highly urbanized area (H). Only three species were found in all sites, and 58 species were collected in only one site.

A total of 5368 bees from 69 species were captured. Advanced eusocial species were clearly dominant. Approximately 96% of all captured bees \( (n = 5150) \) were from the family Apidae, 93% \( (n = 4809) \) of these belonging to the sub-tribe Meliponina (stingless bees). The most abundant species were *Nannotrigona testaceicornis* (Lepeletier) and *Trigona spinipes* (Fabricius) \( (n = 1469 \text{ and } 1336, 28 \text{ and } 26\% \text{ of all Apidae respectively}) \). Only *Trigona spinipes* and *Tetragonisca angustula* (Latreille) were found in all twelve sites, 34 species were captured in only one site. Forty-one species were collected in the area L; 34 in the area M and 38 in the area H. The observed richness ranged from 9 (site L1) to 26 (site L3). When considering only the advanced eusocial species, observed richness ranged from 3 (site L1) to 9 (sites M2 and M3) (Fig. 4a). For the non-advanced eusocial bees, richness values ranged from 4 (site L1) to 19 (site L3) (Fig. 4b).

Similarly to the bees, the advanced eusocial wasp species dominated the urban habitats studied. A total of 900 wasps from 41 species were captured. Almost 90% of these \( (n = 794) \) were species from the family Vespidae. *Polybia occidentalis* (Olivier) was the most abundant species \( (n = 646; 82\% \text{ of all Vespidae}) \) and the only one captured in all sites. Thirteen species of Vespidae were collected in only one site. Seventeen species were found in the less urbanized area (L), seven in the area with an intermediate level of urbanization and eight in the highly urbanized area. The observed values of species richness varied from 2 (sites H4 and H5) to 11 (site L4) (Fig. 5a).

![Fig. 2. Species richness observed for each Hymenoptera family collected.](image-url)
Additionally, 106 solitary wasps from 20 species were captured. Sixty-three belonged to the family Crabronidae, 29 to Pemphredonidae, 12 to Nyssonidae, 1 to Astatidae and 1 to Philantidae. *Liris* sp. was the most abundant species (*n* = 41; 39% of all solitary wasps), although this group may represent more than one species. No solitary species was captured in all sites and thirteen species were found in only one site. Eleven species were found in the less urbanized area (L), 6 in the area with an intermediate level of urbanization and 10 in the highly urbanized area. The observed richness values varied from 1 (sites L1, M2 and H3) to 6 (sites L4 and H1) (Fig. 5b).

### 3.2. Effects of urbanization

The first canonical root obtained in the CDA of the measured habitat features was significant (eigenvalue = 15.573, $\chi^2_{12} = 25.303$, $P < 0.013$). The habitat features (variables) most strongly correlated with the first root were the percentage of vegetation cover ($r = 0.618$) and the percentage of pavement/building cover ($r = -0.587$) (Table 1). The ordination of the twelve study sites using the first two canonical roots shows an evident separation between the sites in the less urbanized area (L) and the remaining sites (in M and H) (Fig. 6).

No significant roots were obtained when using the abundance data of the non-advanced eusocial species. When considering only the abundances of advanced eusocial bee species, one significant discriminant root was obtained (eigenvalue = 16.639, $\chi^2_{14} = 23.746$, $P < 0.049$). The abundance of *Nannotrigona testaceicornis* (Lepeletier) was the variable most strongly correlated with that root ($r = 0.846$) (Table 2). The ordination of the 12 sites using the first two discriminant axes clearly showed a separation between the sites in the less urbanized area and the others when considering their relative position in the first axis (Fig. 7).

A significant negative relationship was found between the first discriminant root obtained with the habitat features data and the first root obtained with the

### Table 1

<table>
<thead>
<tr>
<th>Habitat features</th>
<th>Root 1</th>
<th>Root 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percentage of pavements/buildings</td>
<td>-0.587</td>
<td>0.587</td>
</tr>
<tr>
<td>Percentage of vegetation cover</td>
<td>0.618</td>
<td>-0.559</td>
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<td>Distance to large green areas</td>
<td>-0.326</td>
<td>-0.003</td>
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<td>Variation in the percentage of</td>
<td>0.296</td>
<td>-0.206</td>
</tr>
<tr>
<td>pavements/buildings</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Variation in the percentage of</td>
<td>-0.260</td>
<td>0.291</td>
</tr>
<tr>
<td>vegetation cover</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Variation in the percentage of</td>
<td>-0.145</td>
<td>0.570</td>
</tr>
<tr>
<td>bare soil</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*The percentage of bare soil was removed from this analysis because it is completely dependent on the first two variables.*
Fig. 4. Plots of estimated species richness of bees (means and 95% confidence limits). Black dots indicate the observed number of species. (a) Advanced eusocial species (i.e. stingless bees). (b) Non-advanced eusocial species. Capital letters indicate the urbanization level of each area (low, medium or high).

abundances of advanced eusocial bees ($\beta = -0.938$, $r^2 = 0.881$, $P < 0.000$) (Fig. 8).

To avoid possible distortions of the results of the CDAs caused by spatial autocorrelation (as defined by Legendre and Legendre, 1998) between the sites in each urbanization group, each CDA was performed several times, subsequently removing one of the sites in each group. The results were considered valid only when the significance of the roots and the order of magnitude and signs of the correlations between variables and discriminant roots were preserved (Manly, 1994).

The first two factors extracted in the PCA based on the habitat features of each square and its surroundings, accounted for 39 and 23% of the variance in the model. The most important variables contributing to these factors were the total area of the square, the percentages of the surrounding area covered by pavements/buildings, and vegetation, shrub biomass within

<table>
<thead>
<tr>
<th>Species</th>
<th>Root 1</th>
<th>Root 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Geotrigona subterranea</td>
<td>-0.150</td>
<td>-0.437</td>
</tr>
<tr>
<td>Nannotrigona testaceicornis</td>
<td>0.846</td>
<td>0.160</td>
</tr>
<tr>
<td>Paratrigona lineata</td>
<td>-0.100</td>
<td>0.208</td>
</tr>
<tr>
<td>Plebeia droryana</td>
<td>0.221</td>
<td>-0.204</td>
</tr>
<tr>
<td>Tetragonisca angustula</td>
<td>0.015</td>
<td>0.195</td>
</tr>
<tr>
<td>Trigona hyalinata</td>
<td>-0.043</td>
<td>-0.551</td>
</tr>
<tr>
<td>Trigona spinipes</td>
<td>0.043</td>
<td>-0.272</td>
</tr>
</tbody>
</table>
the squares and tree structural heterogeneity (Table 3). The most important variables for the second factor were the percentage of bare soil, pavements and grass within the square. The herbal/shrub structural heterogeneity in the square and vegetation heterogeneity in the surrounding area of the square may also be considered relevant (Table 3).

Fig. 5. Plots of estimated species richness of wasps (mean values and 95% confidence limits). Black dots indicate the observed number of species. (a) Advanced eusocial species. (b) Non-advanced eusocial species. Capital letters indicate the urbanization level of each area (low, medium or high).

The first two factors obtained from a principal component analysis of the abundances of advanced eusocial bees represented 35 and 26% of the variance in the model. *Nannotrigona testaceicornis* and *Tetragonisca angustula* (Latreille) were the species (variables) more relevant for the first factor and *Paratrigona lineata* (Lepeletier) and *Geotrigona subterranea* (Friese) for the second factor (Table 4). The first factor obtained with the habitat features explained positively the variation in the first factor obtained with the abundances of advanced eusocial bees ($\beta = 0.876$, $r^2 = 0.767$, $P < 0.004$) (Fig. 9). No significant relationship was found between the PCA factors based on the habitat features and the PCA factors obtained with the abundance data of other insect groups considered here.

A significant negative relation was found between the second factor obtained with the habitat features of
Fig. 6. Ordination plot of the study sites in relation to the two roots (axes) obtained from a canonical discriminant analysis of the habitat features measured at each site. Only the first root was significant (eigenvalue = 15.573, $\chi^2_{12} = 25.303$, $P < 0.013$; see Table 1).

Fig. 7. Ordination plot of the study sites in relation to the two roots (axes) obtained from a canonical discriminant analysis of the abundances of advanced eusocial bees (i.e. stingless bees). Only the first root was significant (eigenvalue = 16.639, $\chi^2_{14} = 23.746$, $P < 0.049$; see Table 2).

the squares and the estimated values (averages) of advanced eusocial bees richness for each square ($\beta = -0.784, r^2 = 0.615, P < 0.021$) (Fig. 10). No significant relationship was found between the factors obtained with the habitat features and the estimated richness of other insect groups.

4. Discussion

The advanced eusocial bee species, particularly the stingless Meliponini, dominated the urban system studied. Previous surveys of bee assemblages in other Brazilian cities (e.g. Curitiba and São Paulo),
Fig. 8. Linear regression plot of the relationship between the significant discriminant root related to habitat features and the significant discriminant root related to the abundances of advanced eusocial bees (i.e., stingless bees). Dotted lines indicate 95% confidence limits ($\beta = -0.938$, $r^2 = 0.881$, $P < 0.000$).

Fig. 9. Linear regression plot of the relationship between the first PCA factor obtained with the habitat characteristics of the squares (Table 3) and the first PCA factor obtained with the abundances of advanced eusocial bees (Table 4). Dotted lines indicate 95% confidence limits ($\beta = 0.876$, $r^2 = 0.767$, $P < 0.004$).
also have found a superior abundance of Meliponini 
(Laroca et al., 1982; Knoll et al., 1994). However, 
this pattern is not restricted to urban environments. 
Studies of the bee fauna of Minas Gerais in non-urban 
areas have also shown the dominance of stingless bees 
(Silveira et al., 1993; Silveira and Campos, 1995). Ac-
cording to Heithaus (1979), this is a general tendency 
in bee faunas of the tropics. Roubik (1992) suggested 
that the dominance of advanced eusocial species is

Fig. 10. Linear regression plot of the relationship between the second PCA factor obtained with the habitat features measured at the squares 
(Table 3) and the estimated richness (mean values) of advanced eusocial bees. Dotted lines indicate 95% confidence limits ($\beta = -0.784$, 
r^2 = 0.615, P < 0.021).

Table 3
Factor loadings for the habitat features measured in public squares 
and their surroundings for the first two factors obtained in the 
principal component analysis

<table>
<thead>
<tr>
<th>Habitat features in public squares</th>
<th>Factor 1</th>
<th>Factor 2</th>
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</thead>
<tbody>
<tr>
<td>Total area of the square</td>
<td>0.901</td>
<td>-0.345</td>
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<tr>
<td>Grass percent cover</td>
<td>0.259</td>
<td>-0.647</td>
</tr>
<tr>
<td>Bare soil percent cover</td>
<td>0.231</td>
<td>0.810</td>
</tr>
<tr>
<td>Pavements percent cover</td>
<td>-0.272</td>
<td>0.768</td>
</tr>
<tr>
<td>Tree biomass</td>
<td>0.570</td>
<td>-0.385</td>
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<tr>
<td>Tree structural heterogeneity</td>
<td>-0.831</td>
<td>-0.310</td>
</tr>
<tr>
<td>Tree species richness</td>
<td>0.475</td>
<td>-0.555</td>
</tr>
<tr>
<td>Shrub species richness</td>
<td>0.513</td>
<td>-0.169</td>
</tr>
<tr>
<td>Shrub biomass</td>
<td>0.848</td>
<td>-0.185</td>
</tr>
<tr>
<td>Shrub structural heterogeneity</td>
<td>-0.012</td>
<td>0.611</td>
</tr>
<tr>
<td>Percentage of pavements/buildings</td>
<td>0.860</td>
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<td>Percentage of vegetation cover</td>
<td>-0.854</td>
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<td>Variation in the percentage of vegetation cover</td>
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<td>0.607</td>
</tr>
<tr>
<td>Variation in the percentage of bare soil</td>
<td>0.847</td>
<td>-0.287</td>
</tr>
<tr>
<td>Variance explained (%)</td>
<td>39</td>
<td>23</td>
</tr>
</tbody>
</table>

In the surrounding area of the squares.

Table 4
Factor loadings for the abundances of advanced eusocial bee 
species for the first two factors obtained in the principal component analysis

<table>
<thead>
<tr>
<th>Species</th>
<th>Factor 1</th>
<th>Factor 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Friesella schrotky</td>
<td>0.405</td>
<td>-0.011</td>
</tr>
<tr>
<td>Geotrigona subterranea</td>
<td>-0.383</td>
<td>0.811</td>
</tr>
<tr>
<td>Nannotrigona testaceicornis</td>
<td>0.931</td>
<td>-0.086</td>
</tr>
<tr>
<td>Paratrigna lineata</td>
<td>0.112</td>
<td>0.821</td>
</tr>
<tr>
<td>Partamona helleri</td>
<td>0.683</td>
<td>0.077</td>
</tr>
<tr>
<td>Plebeia droryana</td>
<td>0.719</td>
<td>-0.134</td>
</tr>
<tr>
<td>Tetragonisca angustula</td>
<td>0.820</td>
<td>0.217</td>
</tr>
<tr>
<td>Trigona spinipes</td>
<td>0.214</td>
<td>0.783</td>
</tr>
</tbody>
</table>

| Variance explained (%)           | 36       | 25       |
related to their great capacity to find and store food, essential in tropical environments where the resource distribution is highly heterogeneous. Such ability may be fundamental in urban environments where floral resources are sometimes scarce. In addition, it must be considered that several of the dominant stingless bees are generalist species, which can use many of the exotic ornamental plants found in the streets and public squares. Studies with other insect groups that use exotic ornamental plants found in the streets and public squares indicate that several of the dominant stingless bees are generalist species, which can use many of the exotic ornamental plants favored in urban environments (Ruszczyk, 1986a,b; Blair and Launer, 1997).

The prediction that more urbanized areas are less complex and for this reason support a smaller abundance of wasps and bees was partially confirmed. The less urbanized area (L) could be clearly differentiated from the others by its landscape structure, mostly by the percentage of vegetation cover and the smaller percentage of impervious surfaces. However, the CDA results have showed that the area with an intermediate level of urbanization (M) and the highly urbanized area (H) can not be differentiated by the measured habitat features (Fig. 6), even if these areas clearly have different levels of urbanization.

The less urbanized area (L) could also be discriminated from the others by the abundances of advanced eusocial bees (i.e. stingless bees), but no discrimination was possible between the other areas (M and H; Fig. 7). The faunistic and structural similarities between these two areas may be a consequence of the historical differences in their urbanization. The highly urbanized area (H) has been urbanized since the foundation of Belo Horizonte (1897) in a highly planned way, and has a large number of public green areas like parks, squares and gardens but a reduced number of private green areas (Horta, 1994). The area with an intermediate level of urbanization (M) has been urbanized in a less planned way during the last three decades, and consequently has only a few public green areas but has many private green areas and vacant lots (Pedersoli et al., 1996). Therefore, both areas (H and M) may be similar in terms of available habitats and resources for the insect fauna.

The significant relationship observed between the abundance of advanced eusocial bees and the percentages of pavements/buildings and vegetation clearly indicates that habitat changes associated with urbanization can affect these bees (Fig. 8). However, it also indicates that the magnitude of this effect could vary according to the nesting habits of each species (Table 2). Species that nest in pre-existent cavities, like Nannotrigona testaceicornis and Plebeia drovana, were highly abundant in the more urbanized areas where the high density of buildings provides an extensive range of nesting sites (e.g. walls, eaves and roofs). In contrast, this high density of buildings reduces the availability of nesting sites for ground nesting species, which could explain the reduced abundance of Geotrigona subterranea and Paratrignona lineata in the two more urbanized areas. Laroca et al. (1982) suggested that the availability of nesting sites was a limiting factor for several Neotropical bee species found in a highly urbanized area in the city of Curitiba (Brazil), Torres and Gayubo (1989), also have suggested that ground-nesting bees and wasps were more abundant in the less or non-urbanized regions of Salamanca (Spain) in response to bare soil availability.

Although no significant relation was found between the abundance of advanced eusocial wasps and the habitat changes related to urbanization, the small abundances observed for almost all species suggest that urban environments are more restrictive to advanced eusocial wasps than bees. The reduction in prey availability caused by the loss of vegetation may be one of the factors responsible for these results (Skibinska, 1986; Gayubo and Torres, 1990; Raw, 1998a). The high values of species richness found in the sites with the highest percentage of vegetation cover, in the less urbanized area, reinforce this hypothesis. In addition, the increase of pavements and buildings coupled with the reduction of vegetation cover, increases the exposure of wasp colonies to possible predators and particularly to humans, which are the major cause of colony mortality in urban environments (Fowler, 1983, Skibinska, 1986; Gunnarsson and Hake, 1999). Despite all these limitations, the great abundance of Polybia occidentalis (Olivier) in highly urbanized sites indicates that advanced eusocial wasp species that can exploit many food resources and use less-exposed nesting sites (i.e. pre-existent cavities) may be less affected by urbanization. Skibinska (1986) also observed that the most abundant wasp species in the highly urbanized areas of Varsovia (Poland) had these characteristics. Similarly, the only three advanced eusocial wasp species...
found in urbanized areas of Salamanca (Spain) were generalists (Gayubo and Torres, 1989, 1990).

The prediction that larger public squares can attract a greater number of insects was confirmed only for the advanced eusocial bees. However, the effect of size was directly related to the biomass of herbs and shrubs in the squares (Table 3). Since most of the floral resources available in the squares are concentrated in these plants, large squares offered more resources and consequently could attract more bees. In addition, the analyses with the squares also indicated that the local availability of floral resources (consequently the size of the squares) could be more important than the local availability of nesting sites for species like *Tetragononisa angustula*, *Frieseella schrottky* and *Partamona helleri*, independently of their nesting habits.

Contrary to what was predicted, the reduction of natural structural complexity in the surrounding area of the squares can have a positive effect on the abundance of advanced eusocial bees (e.g. *Nannotrigona testaceicornis*) (Fig. 9). These results reinforce the hypothesis that species that nest in pre-existing cavities could be favored by the increased availability of nesting sites associated with higher levels of urbanization.

The prediction that species richness could be related to the size of the squares or the structure of their surroundings was not confirmed. However, the negative correlation between the richness of advanced eusocial bee species and the percentage of bare ground and pavements in the squares (Fig. 10), suggests that the total percentage of vegetation in the squares may be affecting local species richness.

Overall, the responses of non-advanced eusocial species to urbanization or to the structural variation in the public squares were not as clear as the responses of advanced eusocial species. One possible explanation is that their responses may occur in spatial scales not considered here. Solitary species, for example, explore a smaller portion of their habitats when compared to a whole colony (considering insects with similar size) (Bohart and Menke, 1976; Roubik, 1992; von Nieuwstadt and Iraheta, 1996). Consequently, their responses to habitat changes are more rapid (Wiens, 1976; Didham et al., 1996). In addition, the spatial scales considered here could be too small to encompass the responses of species with great flying capacity. Species like *Xylocopa* (*Neoxyllocopa*) *frontalis* and *Bombus* (*Fervidobombus*) *morio*, for example, can fly more than 10 km from their nest searching for floral resources (Roubik, 1992). For such species the distances between the study sites (even between the areas) may be not relevant. Therefore, studies of larger portions of urban landscapes are necessary to analyze their responses to urbanization (Lord and Norton, 1990; Wiens et al., 1993; Hostetler, 1999).

4.1. Implications for conservation

The results presented here suggest that some species of advanced eusocial bees (e.g. *Geotrigona subterranea*), which are directly affected by the habitat changes related to urbanization, could be used as indicators of habitat conditions (as defined by Brown, 1991). These potential indicators, which are easily identified, could be linked to estimates of habitat or life quality for the local human population (Hostetler, 1999). Such association may be very useful for landscape planners (and public authorities) interested in monitoring the development of urban environments. In addition, it may provide conservationists with an extra argument when defending a particular wasp or bee species (Hostetler, 1999). Considering that the occurrence of most advanced eusocial wasps was related to the percentage of vegetation cover, they could also be considered as potential habitat quality indicators. However, further studies are necessary to test which species are the best indicators.

The different responses to urbanization presented by wasps and bees at different spatial scales indicate that any conservation initiative in urban environments has to consider distinct spatial scales to be effective (Theobald et al., 1997; Cane, 2001). Considering that the structure and functioning of a city are based on an hierarchical system of decisions, conservation actions may be ‘inserted’ in each level of decisions (Savard et al., 2000). Changes in the vegetation of a private garden or a public square (related to local authorities and lot owners), for example, may attract a greater number of stingless bees. Additionally, large scale changes in the urban landscape, like the creation of a public park (related to city authorities), may be essential to maintain or increase the diversity of advanced eusocial wasps.

Finally, one of the major objectives of nature conservation in cities is to promote direct interaction between local inhabitants and native species of plants.
The public squares studied present many opportunities for direct contact with wasp and bee species, especially and most desirably with the abundant stingless bees. The almost harmless stingless bees (Meliponini) are the most charismatic and well known group of tropical bees (Nogueira-Neto, 1970). Adding floral resources to the squares could easily increase the frequency of contacts with these bees. This notion of “faunistic landscaping” is fundamental to develop nature conservation in cities (Ruszczyk, 1986c).

Although overlooked by many conservationists and ecologists, the rapid expansion of cities is a global phenomena and a pressing problem that must be seriously confronted. Despite the fact that some species may benefit from increased urbanization, overall the results presented here indicate that this process has strong negative effects on key predator and pollinator species. Further research must be done urgently in order to find less impacting forms of urbanization.

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