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# Global loss of avian evolutionary uniqueness in urban areas

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# Abstract

Urbanization, one of the most important anthropogenic impacts on Earth, is rapidly expanding worldwide. This expansion of urban land-covered areas is known to significantly reduce different components of biodiversity. However, the global evidence for this effect is mainly focused on a single diversity measure (species richness) with a few local or regional studies also supporting reductions in functional diversity. We have used birds, an important ecological group that has been used as surrogate for other animals, to investigate the hypothesis that urbanization reduces the global taxonomical and/or evolutionary diversity. We have also explored whether there is evidence supporting that urban bird communities are evolutionarily homogenized worldwide in comparison with nonurban ones by means of using evolutionary distinctiveness (how unique are the species) of bird communities. To our knowledge, this is the first attempt to quantify the effect of urbanization in more than one single diversity measure as well as the first time to look for associations between urbanization and phylogenetic diversity at a large spatial scale. Our findings show a strong and globally consistent reduction in taxonomic diversity in urban areas, which is also synchronized with the evolutionary homogenization of urban bird communities. Despite our general patterns, we found some regional differences in the intensity of the effect of cities on bird species richness or evolutionary distinctiveness, suggesting that conservation efforts should be adapted locally. Our findings might be useful for conservationists and policymakers to minimize the impact of urban development on Earth's biodiversity and help design more realistic conservation strategies.

*Keywords:* biodiversity, biotic homogenization, birds, evolutionary distinctiveness, human-induced environmental change, urban ecology

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# Introduction

The expansion of urban areas is considered one of the most important anthropogenic impacts on the Earth's ecosystem (Foley *et al.*, 2005; Groom *et al.*, 2006), dramatically changing natural habitat structure, ecosystem functioning, and life's biodiversity (Grimm *et al.*, 2008a; Gaston, 2010; Forman, 2014; Gil & Brumm, 2014). Urban areas are growing proportionally faster than any other form of land cover (United Nations, 2012) as urbandwelling human population is increasing by 1 million per week (Seress & Liker, 2015). By 2030, there is an expected 200% increase in global urban land cover in comparison with the year 2000 (Fragkias *et al.*, 2013). It is not surprising therefore that the process of urbanization and its environmental impact is now recognized as

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a major global challenge (United Nations, 2016) and has attracted increased attention in the past decades (McDonnell *et al.*, 2016).

Organisms can exhibit different responses to urbanization (McDonnell & Hahs, 2008), and while a few species are beneficiated by cities (i.e., urban exploiters), many more are negatively affected by this landscape change (i.e., urban avoiders; Blair, 1996; Sol et al., 2014). In fact, one of the most important ecological effect of urbanization is the overall decrease in taxonomic diversity (Marzluff, 2001; Chace & Walsh, 2006; McKinney, 2008; Newbold et al., 2015). This loss of biodiversity is caused by both changes in the abiotic (i.e., habitat loss Clergeau et al., 1998; Kühn & Klotz, 2006) and biotic component of the environment (i.e., invasive species Grimm et al., 2008b; Luck & Smallbone, 2010; Garcillán et al., 2014) which lead to the extirpation of native organisms and can drive the homogenization of biotas in cities (McKinney, 2006; Devictor et al., 2007; Morelli

© 2016 The Authors. *Global Change Biology* Published by John Wiley & Sons Ltd. This is an open access article under the terms of the Creative Commons Attribution-NonCommercial License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited and is not used for commercial purposes. *et al.*, 2016). This issue is even more relevant considering that this negative impact of urban land-cover expansion on biodiversity is expected to increase globally in the future due to the selective expansion of towns and cities in key biodiversity hot spots (Seto *et al.*, 2012). This scenario has led to consider urbanization as a major topic in conservation biology (Miller & Hobbs, 2002; McKinney, 2006) and has also attracted the attention of politics and policymakers (Secretariat of the Convention on Biological Diversity, 2012; Elands *et al.*, 2015). The recently published World Cities Report (United Nations, 2016) states that one of the main challenges regarding urbanization is related to 'minimizing the negative impacts of land transformations in the use of resources, biodiversity, and ecosystems'.

However, previous knowledge on the effect of the urban land-cover expansion on biodiversity is mainly based on a single measure of biodiversity, species richness (Marzluff, 2001; Chace & Walsh, 2006; McKinney, 2008), while very little attention has been paid to other diversity measures (Devictor et al., 2008; Knapp et al., 2008; Morelli et al., 2016). Biodiversity refers to variation at all levels of biological organization and includes three main elements (genetic, ecological, and organismal diversity), and although species richness is the most commonly used measure of biodiversity, it has important restrictions to capture certain components of biodiversity (Gaston & Spicer, 2004). Another important issue directly related to the biodiversity loss driven by urbanization is the process of biotic homogenization (Clergeau et al., 2006; McKinney, 2006). Irrespective of the biome, urbanization creates habitats with particular physical characteristics across locations (Groffman et al., 2014) that favor some synurbic species (those particularly associated to urban areas) while reducing local species (Francis & Chadwick, 2012). And, at the same time, it favors transportation of non-native species around the world (McKinney, 2008). This process tends to create similar genetic, taxonomic, and/or functional biotas all over the world (Olden et al., 2004; Devictor et al., 2008; Knop, 2016) and acts mainly through the extinction of native species and introduction of alien organisms (Olden & Poff, 2003; Klotz & Kühn, 2010; Shochat et al., 2010; Čeplová et al., 2015).

The evolutionary distinctiveness or uniqueness is a measure of evolutionary history, more related to the genetic element of biodiversity. Basically, it measures how isolated a species is in a phylogenetic tree and represents uniquely divergent organisms (Isaac *et al.*, 2007; Jetz *et al.*, 2014). Consequently, this measure of phylogenetic diversity can play an important role to establish conservation priorities (Isaac *et al.*, 2007;

EDGE of Existence, 2015). For instance, recent recommendations in nature conservation optimization encourage the use of measures of phylogenetic diversity alongside species richness (Winter et al., 2012; Monnet et al., 2014). In relation to urbanization, two recent regional-scale studies have found that urban areas hold communities with low levels of phylogenetic diversity. Čeplová et al. (2015) showed that the phylogenetic diversity of plant communities in Central Europe was lower than random. Similarly, Morelli et al. (2016) has found a significant reduction in evolutionary distinctiveness of urban bird assemblages in comparison with rural ones (Morelli et al., 2016). This study did not detect significant differences in bird species richness between the two habitats, which strongly supports the need for exploring other measures of biodiversity, to gain a more complete picture of the ecological effects of urbanization. However, even if these two studies suggest the phylogenetic homogenization of European bird and plant assemblages due to urbanization (Čeplová et al., 2015; Morelli et al., 2016), there has not been a general attempt to explore these patterns at a larger-scale yet. The global-scale perspective is particularly relevant because even though urbanization is a worldwide process (Grimm et al., 2008a), the majority of papers that explore how urbanization affects biodiversity are geographically restricted preventing generalization of many observed patterns (Pautasso et al., 2011; Müller et al., 2013; Seress & Liker, 2015). Moreover, urban land-cover change can also have different impacts depending on the particular characteristics of the region (McKinney, 2002; Grimm et al., 2008b). In fact, some particular areas are expected to be affected more deeply than others by urbanization (Seto et al., 2012) and, therefore, the comparison among regions will help us to establish conservation priorities more accurately.

Here, using birds and a large spatial scale approach, we (i) investigate whether urban areas from different continents are associated with lower biodiversity measures (species richness and evolutionary distinctiveness) than nonurban areas and (ii) look for evidence supporting that urban landscapes are associated with phylogenetically homogenized communities worldwide. We have used birds as our model group because they are key components of ecosystems (Sekercioğlu, 2006), good bioindicator for other taxa (Rodrigues et al., 2007), with a widely validated phylogeny (Jetz et al., 2012; Prum et al., 2015) and well studied within the urban ecology context (Marzluff et al., 2001; Lepczyk & Warren, 2012; Gil & Brumm, 2014), thus offering the opportunity to compile a geographically wide database while testing our hypotheses in relation to an ecological relevant group.

## Materials and methods

#### Bibliographic research and selection of studies

We did an exhaustive literature search in Web of Science, Google Scholar, and SmartCat (www.rug.nl/bibliotheek/smartca t/) using different combinations of the following terms: 'bird\*', 'assemblages' and 'urban\*'. We look for studies presenting information on the avian community of both urban and nonurban habitats from the same area. If possible, we collected data from the wildland habitat sensu Marzluff et al. (2001) as the nonurban habitat, although it was not always available and in some cases it represented an agricultural (rural) land cover, according to definitions provided by Marzluff et al. (2001). Given that agricultural habitats usually present lower avian biodiversity values than native ones (e.g., Sinclair et al., 2002; Billeter et al., 2008), we consider our procedure to be conservative. We discarded studies not showing the complete bird community (i.e., only most common species) or those presenting information from only one of the two habitats (i.e., only urban species). This study selection criterion allowed us to collect data on urban and nonurban bird communities that were obtained following the same field method, during the same time period and by the same field observer, thus standardizing for many potential confounding factors in this kind of comparative study. Some papers presented information from several locations within an urban gradient. In such cases, we extracted the data of those two locations that better represented the urban and wildland/rural habitats sensu Marzluff et al. (2001).

From each study, we extracted the following information: (i) urban bird assemblage, (ii) nonurban bird assemblage, (iii) continent, (iv) country, (v) site, (vi) latitude, (vii) longitude, and (viii) whether the data correspond to the breeding, nonbreeding season, or both. Coordinates were obtained directly from the paper or from Google Maps if not available in the study.

#### Biodiversity and evolutionary distinctiveness

From each study, we calculated two diversity values (species richness and evolutionary distinctiveness) for each habitat. The bird species richness was calculated as the highest number of species observed in each location as it is a basic surrogate for taxonomic diversity (Magurran, 2004). The evolutionary distinctiveness (ED) score is a measure of richness in phylogenetic diversity (Tucker et al., 2016) based on the uniqueness of a species (Redding et al., 2010; Jetz et al., 2014). The ED score is calculated considering the sum of all phylogenetic branch lengths connecting species (Cadotte et al., 2008, 2012) and by applying a value to each branch equal to its length divided by the number of species subtending the branch (Isaac et al., 2007). Using the ED score for each bird species present in a community, we estimated the community evolutionary distinctiveness (CED) as the average ED considering all species belonging to the community. The CED is a measure of divergence, related to the phylogenetic isolation (Tucker et al., 2016). This procedure has also the advantage to make this variable independent from the number of species present in each community, optimizing the comparison among many studies. Furthermore, the highest ED score of a bird species present in the assemblage was labeled as the maximum ED for this bird community (max ED).

#### Statistical analyses

We used generalized linear mixed models (GLMMs) to study the patterns of bird richness, CED and max ED in relation to urban and nonurban habitats, latitude and longitude modeled as fixed effects. We decided to include these geographical predictors as they can be related to large-scale changes in biodiversity (Mannion *et al.*, 2014; Morelli *et al.*, 2016). Country (groups = 17) and season (group = 3) were included as random effects to account for possible consistent differences among countries or in relation to breeding or nonbreeding period. The use of random effects permits statistical tests when some observations can be correlated. In the particular study case, it is useful in order to remove any potential differences related to methods or sampling effort among studies in each country.

Models using CED and max ED as response variable were fitted by maximum likelihood, using the package 'lme4' in R (Bates et al., 2014), while models using bird species richness as response variable were fitted assuming a Poisson distribution after having explored the variable distribution as suggested in Box & Cox (1964) using the package 'MASS' (Venables & Ripley, 2002) and 'glmmADMB' in R (Fournier et al., 2012; Skaug et al., 2013). In this study, the Akaike information criterion (AIC) was calculated to determine the models that 'best' explained variation in the data (Burnham & Anderson, 2002). The model selection and multimodel inference were performed using the package 'AICcmodavg' in R (Mazerolle, 2016). The best model was selected considering both lowest AIC and large Akaike weights, because this model had the strongest support for data (Mazerolle, 2016). The confidence intervals for the significant variables selected in the best model were calculated by the Wald method from the package 'MASS' (Venables & Ripley, 2002). All statistical tests were performed with R software (R Core Team, 2016).

#### Results

Our literature search provided us with 26 studies that matched our strict search criteria. We identified a total of 665 bird species from 28 paired urban vs. nonurban locations from 17 countries and four continents (Fig. 1, Table S1). The list of all bird species used in this study in both urban and nonurban sites, with their ED score, is provided in the Table S2.

All three diversity measures (bird richness, CED, and max ED) showed significantly lower values in urban than in nonurban habitats for the global analyses (Table 1; Fig. 2). The confidence intervals for the predictor 'habitat' always ranged across negative values for all three response variables (Table 1). Based on our

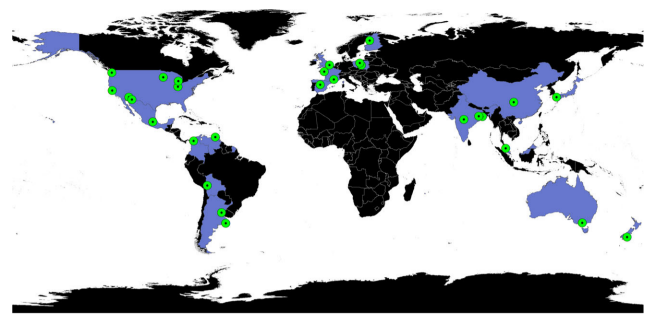


Fig. 1 Location of the 28 cities (and their corresponding nonurban areas) used in our study (green dots) distributed in 17 countries (blue) from four different continents. For coordinates and city names involved in each study, see Table S1. [Colour figure can be viewed at wileyonlinelibrary.com]

**Table 1** Results of fixed-effect parameters in the generalizedlinear mixed models (GLMMs), accounting for variation inbird richness, community evolutionary distinctiveness (CED),and max ED in relation to urban or nonurban habitat

Source of variation	Estimate	CI	SE	t/z	Р
Bird richne Habitat: urban	ess -0.55	-0.65, -0.45	0.05	-10.6	2E-16
CED Habitat: urban	-0.56	-1.05, -0.08	0.23	-2.3	0.002
max ED Habitat: urban	-4.75	-9.29, -0.26	2.3	-2.1	0.038

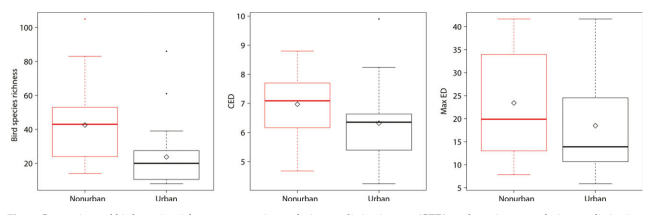
CI, confidence interval (lower/higher); SE, standard error. Only significant variables are shown in the table.

statistical procedures, these differences were independent of the country or season as we included them as random factors. This general pattern is also maintained after grouping the results of all studies by continent for both bird species richness (Fig. 3) and CED (Fig. 4). Then, the urban biodiversity was lower than the nonurban biodiversity in all continents. However, we found some interesting results with South American bird communities showing the largest difference in species richness but the smallest difference for CED (Figs 3 and 4). Oceania also presented remarkable results with a similar difference for bird species richness between habitats as that observed in other continents, but the largest difference in CED (Figs 3 and 4).

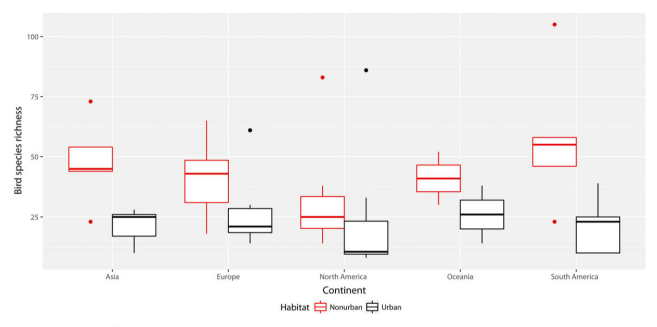
Longitude and latitude did not significantly improve the proportion of species richness, CED, or max ED variance explained by the model including only habitat as a predictive variable (Table S3) suggesting that those geographical predictors are of little importance regarding our data. This result supports the generality of our findings and highlights that the main factor explaining variation in avian CED is the urban vs. nonurban comparison. The list of evaluated models, with the number of predictors considered, delta AIC values, and AIC weights for each one is provided in Table S3.

# Discussion

Our findings highlighted the fact that urbanization significantly reduced both species richness and evolutionary distinctiveness worldwide (Fig. 2), supporting the assumption that the expansion of urban areas reduces biodiversity. Considering that urbanization is a similar process acting worldwide (McKinney, 2006; Grimm *et al.*, 2008a), the large-scale geographical distribution of our database (including cities in four continents) and the nonsignificant effect of geographical variables (latitude and longitude) in any of our biodiversity



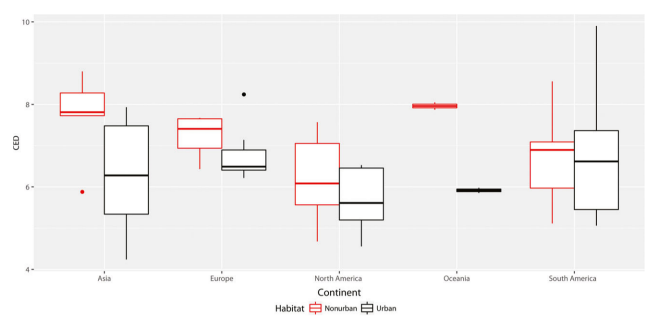
**Fig. 2** Comparison of bird species richness, community evolutionary distinctiveness (CED), and maximum evolutionary distinctiveness (max ED) between urban and nonurban habitats. The *y*-axis represents the estimated variable. The box plots show the median (bar in the middle of rectangles), mean (rhombus), upper and lower quartiles, maximum and minimum values (vertical dashed lines), and outliers. [Colour figure can be viewed at wileyonlinelibrary.com]



**Fig. 3** Comparison of bird species richness between habitats in each continent. The *y*-axis represents the estimated variable. The boxplots show the median (bar in the middle of rectangles), upper and lower quartiles, maximum and minimum values (vertical lines), and outliers. [Colour figure can be viewed at wileyonlinelibrary.com]

measures, we believe that our results could be reasonably considered representative of a global pattern. The result on species richness matches with previous findings on birds (Marzluff, 2001; Chace & Walsh, 2006; Sol *et al.*, 2014) and other organisms (McKinney, 2008; Luck & Smallbone, 2010; Knop, 2016). However, it contrasts with another previous global-scale study on avian assemblages which did not find support for this effect (Pautasso *et al.*, 2011). The reason for this difference might be our selection of papers presenting simultaneously urban and nonurban assemblages, while Pautasso *et al.* (2011) compared independent studies for each habitat. In general, our study presents another piece of evidence indicating that the number of bird species is reduced in urban habitats in comparison with nonurban habitats.

Our results on global evolutionary distinctiveness support those obtained by Morelli *et al.* (2016) for European bird communities, clearly showing that urbanization also affects other components of avian biodiversity (phylogenetic diversity) other than species richness. The reduced avian phylogenetic diversity associated



**Fig. 4** Comparison of community evolutionary distinctiveness between habitats in each continent. The *y*-axis represents the estimated variable. The boxplots show the median (bar in the middle of rectangles), upper and lower quartiles, maximum and minimum values (vertical lines), and outliers. [Colour figure can be viewed at wileyonlinelibrary.com]

with urban landscapes could have important conservation implications, such as reorienting our conservation efforts to prevent the loss in evolutionary information (Mace et al., 2003; Winter et al., 2012). Phylogenetic diversity is a good surrogate for species rarity and can be used as complementary information for conservation plans by identifying and prioritizing those species in need of protection (Winter et al., 2012) or the best targets to retain a major proportion of evolutionary history in urban bird communities. It will also be very interesting to test whether other organisms are impacted in a similar way. Birds are good predictors of abundance of other vertebrates (Rodrigues et al., 2007), and even if invertebrates follow a similar pattern of reduction in species richness due to urbanization (McKinney, 2008; Knop, 2016), they might differ regarding other components of biodiversity. Plants are particularly interesting organisms in this respect because of the structural role that they play in cities (Groffman et al., 2014) and the different patterns (in comparison with animals) that manifest due to urbanization (increase in overall species richness, but reduction in native vegetation), particularly because of the introduction and growth of non-native species in urban areas (Palomino & Carrascal, 2006; McKinney, 2008; Luck & Smallbone, 2010; Čeplová et al., 2015). Furthermore, it has been shown that the high species richness of urban flora in Germany was not reflected in high phylogenetic distinctness suggesting that these two measures of biodiversity are not directly related (Knapp et al., 2008).

Interestingly, a recent study has shown that urbanization seems to reduce phylogenetic diversity of urban plant communities in Central Europe, suggesting that the pattern observed in our study might apply to plants too (Čeplová et al., 2015). Worth to mention is that habitat type (urban vs. nonurban) and not geographical predictors (latitude or longitude) explained our CED results which suggests that this effect is global and independent of the location. This is also supported by the shared pattern found for this biodiversity proxy in all continents (Fig. 4). However, these results contrast with those found by Morelli et al. (2016) showing a negative significant correlation between CED and latitude for European birds. It is possible that this pattern is restricted to Europe or that it only arises when cities from very high latitudes are included in the analyses. Only 4% of our cities were located higher than 55°N, compared with 50% of the locations from Morelli's European study. In addition to this lack of high altitude studies on urbanization, we found other geographical areas in which we should investigate this issue further (Fig. 1). Probably the clearest example is Africa, which in addition is going to be a great center of urban development in the forthcoming years (Seto et al., 2012; McPhearson et al., 2016).

Despite the general effect found for biodiversity, the expansion of urban landscapes is not completely homogeneous in all continents (Figs 3 and 4). Bird communities in South America seem to be the most affected in terms of taxonomic diversity, probably because the native habitats in this continent hold the largest avian species richness in the world (Newton, 2003). Nevertheless, South American CED is not reduced so strongly, which suggests that some species with high evolutionary uniqueness are being retained in urban areas in this continent. Oceania shows just the opposite pattern being highly affected in terms of CED, much more than any other region (Fig. 4). This might be due to the small sample size of urban studies for this particular region included in our database, but could also indicate that cities in Australia and New Zealand are selectively removing evolutionary unique species. The latter alternative is of special concern considering that Australia and New Zealand are among the most important hot spots in evolutionary diversity of birds (Jetz et al., 2014). These differences between regions provide valuable information in order to design more efficient conservation plans and point out that conservation strategies should be adapted to the region to minimize the impact of urban land-cover expansion.

Another clear conclusion that could be extracted from our results is about biotic homogenization. We found strong evidence supporting the global taxonomical and evolutionary homogenization of avian communities (Fig. 2). This is in line with previous findings showing that urban birds are taxonomical, functional, and phylogenetically homogenized (e.g., Devictor et al., 2008; Reif et al., 2013; Godet et al., 2015; Morelli et al., 2016). This evolutionary homogenization of bird assemblages could happen because of exclusion of unique species, which will not be suitable to inhabit in urban areas, or due to many recent speciation events in cities (Morelli et al., 2016). Nevertheless, the latter option seems unlikely given that urbanization is a relative recent process in Earth's history (5000-6000 years, Gaston, 2010) and the majority of birds have evolved before humans gathered in settlements (>2.5 MYr, Jetz et al., 2012). Our results could suggest an association between recently diversified lineages and some traits that favor urban life (i.e., generalism, Müller et al., 2013), although the link between phylogenetic and functional diversity is not completely clear yet (Winter et al., 2012). For instance, Sol et al. (2014) showed that sensitivity to urbanization is not randomly distributed across the avian phylogeny, suggesting that there are some lineages of birds more prone to survive in urban areas than others. Furthermore, not only cities contain species with lower average evolutionary distinctiveness, but they also seem to impose an upper limit on evolutionary diversity of urban bird assemblages similarly to what happen with the number of bird species. The max ED is significantly lower in urbanized landscapes than in close nonurbanized areas (Table 1, Fig. 2) which support the biotic homogenization hypothesis. However, this result is difficult to explain in ecological terms and will need further research. One potential explanation could be the extirpation of lineages with extremely high ED values, which would be associated at the same time to those species with very particular requirements/adaptations (i.e., particular food or inability to fly). Alternatively, these high ED birds could be representative of locally restricted species (Webb & Gaston, 2000) which are most likely to not become urbanized (Bonier et al., 2007; Møller, 2014). However, correlations between breeding range size and avian ED are not straightforward and seem to be complex and multifactorial (Jetz et al., 2014). Our study investigated patterns at the community level; however, these patterns should have arisen as a consequence of differences among individuals (i.e., those less afraid to humans (Møller, 2008; Díaz et al., 2013)) and subsequently among species (i.e., those presenting certain traits that facilitate living in urban areas (Luck & Smallbone, 2010; Møller, 2014; Sol et al., 2014). Future work should aim to integrate these three levels (individuals, species, and communities) in order to completely understand the effect of urbanization on biodiversity.

Recent studies have highlighted that cities can host an important level of biodiversity (Müller et al., 2013; Aronson et al., 2014) and can also act as key reservoirs for threatened species (Ives et al., 2016). However, our results raise important conservation concerns given the significant and worldwide reduction in avian biodiversity associated with urbanization. Furthermore, our findings suggest that urbanization, like agriculture, is another human landscape change responsible for the global loss of evolutionary information (Frishkoff et al., 2014) and a key player in the homogenization of biotas. Our conservation efforts should aim at retaining a major proportion of biodiversity in cities (McPhearson et al., 2016; United Nations, 2016), in the case of birds, for example by increasing the native vegetation in urban areas (Aronson et al., 2014; Threlfall et al., 2016) or with particular conservation plans to retain species with high ED values (Winter *et al.*, 2012). The different trends found for some regions between species richness and ED also suggest that more studies using different and complementary biodiversity measures are needed to fully understand how urbanization impacts biodiversity. We hope that our findings can be valuable for conservationists and policymakers when establishing conservation priorities and regulating urban expansion.

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## References

Aronson MFJ, La Sorte FA, Nilon CH et al. (2014) A global analysis of the impacts of urbanization on bird and plant diversity reveals key anthropogenic drivers. Proceedings of the Royal Society of London B – Biological Sciences, 281, 20133330.

- Bates D, Maechler M, Bolker B, Walker S (2014) lme4: linear mixed-effects models using Eigen and S4 R Package.
- Billeter R, Liira J, Bailey D et al. (2008) Indicators for biodiversity in agricultural landscapes: a pan-European study. Journal of Applied Ecology, 45, 141–150.
- Blair RB (1996) Land use and avian species diversity along an urban gradient. Ecological Applications, 6, 506–519.
- Bonier F, Martin PR, Wingfield JC (2007) Urban birds have broader environmental tolerance. *Biology Letters*, 3, 670–673.
- Box GEP, Cox DR (1964) An analysis of transformations. Journal of the Royal Statistical Society B, 26, 211–252.
- Burnham KP, Anderson DR (2002) Model Selection and Multimodel Inference: A Practical Information-Theoretic Approach (2nd edn). Springer-Verlag, New York, NY.
- Cadotte MW, Cardinale BJ, Oakley TH (2008) Evolutionary history and the effect of biodiversity on plant productivity. Proceedings of the National Academy of Sciences of the United States of America, 105, 17012–17017.
- Cadotte MW, Dinnage R, Tilman D (2012) Phylogenetic diversity promotes ecosystem stability. *Ecology*, **93**, 223–233.
- Čeplová N, Lososová Z, Zelený D et al. (2015) Phylogenetic diversity of central-European urban plant communities: effects of alien species and habitat types. Preslia, 87, 1–16.
- Chace JF, Walsh JJ (2006) Urban effects on native avifauna: a review. Landscape and Urban Planning, 74, 46–69.
- Clergeau P, Savard J-PL, Mennechez G, Falardeau G (1998) Bird abundance and diversity along an urban-rural gradient: a comparative study between two cities on different continents. *Condor*, **100**, 413–425.
- Clergeau P, Croci S, Jokimäki J, Kaisanlahti-Jokimäki ML, Dinetti M (2006) Avifauna homogenisation by urbanisation: analysis at different European latitudes. *Biologi*cal Conservation, 127, 336–344.
- Devictor V, Julliard R, Couvet D, Lee A, Jiguet F (2007) Functional homogenization effect of urbanization on bird communities. *Conservation Biology*, **21**, 741–751.
- Devictor V, Julliard R, Clavel J, Jiguet F, Lee A, Couvet D (2008) Functional biotic homogenization of bird communities in disturbed landscapes. *Global Ecology and Biogeography*, 17, 252–261.
- Díaz M, Møller AP, Flensted-Jensen E et al. (2013) The geography of fear: a latitudinal gradient in anti-predator escape distances of birds across Europe. PLoS One, 8, e64634.

EDGE of Existence (2015) www.edgeofexistence.org.

- Elands BHM, Wiersum KF, Buijs AE, Vierikko K (2015) Policy interpretations and manifestation of biocultural diversity in urbanized Europe: conservation of lived biodiversity. *Biodiversity and Conservation*, 24, 3347–3366.
- Foley JA, Defries R, Asner GP et al. (2005) Global consequences of land use. Science, 309, 570–474.
- Forman R (2014) Urban Ecology: Science of Cities. Cambridge University Press, Cambridge.
- Fournier DA, Skaug HJ, Ancheta J et al. (2012) AD Model Builder: using automatic differentiation for statistical inference of highly parameterized complex nonlinear models. Optimization Methods and Software, 27, 233–249.
- Fragkias M, Güneralp B, Seto KC, Goodness J (2013) A synthesis of global urbanization projections. In: Urbanization, Biodiversity and Ecosystem Services: Challenges and Opportunities (eds Elmqvist T, Fragkias M, Goodness J, Güneralp B, Marcotullio PJ, McDonald RI, Parnell S, Schewenius M, Sendstad M, Seto KC, Wilkinson C), pp. 409–435. Springer, Dordrecht.
- Francis RA, Chadwick MA (2012) What makes a species synurbic? Applied Geography, 32, 514–521.
- Frishkoff LO, Karp DS, M'Gonigle LK et al. (2014) Loss of avian phylogenetic diversity in neotropical agricultural systems. Science, 345, 1343–1346.
- Garcillán PP, Dana ED, Rebman JP, Peñas J (2014) Effects of alien species on homogenization of urban floras across continents: a tale of two mediterranean cities on two different continents. *Plant Ecology and Evolution*, **147**, 3–9.

Gaston K (2010) Urban Ecology. Cambridge University Press, Cambridge.

- Gaston KJ, Spicer JI (2004) *Biodiversity: An Introduction*. Blackwell Publishing Ltd, Cornwall.
- Gil D, Brumm H (2014) Avian Urban Ecology: Behavioural and Physiological Adaptations. Oxford University Press, Oxford.
- Godet L, Gaüzere P, Jiguet F, Devictor V (2015) Dissociating several forms of commonness in birds sheds new light on biotic homogenization. *Global Ecology and Bio*geography, 24, 416–426.
- Grimm NB, Faeth SH, Golubiewski NE et al. (2008a) Global change and the ecology of cities. *Science*, **319**, 756–760.
- Grimm NB, Foster D, Groffman P et al. (2008b) The changing landscape: ecosystem responses to urbanization and pollution across climatic and societal gradients. *Frontiers in Ecology and the Environment*, 6, 264–272.
- Groffman PM, Cavender-Bares J, Bettez ND et al. (2014) Ecological homogenization of urban USA. Frontiers in Ecology and the Environment, 12, 74–81.
- Groom MJ, Meffe GK, Carroll CR (2006) Principles of Conservation Biology. Sinauer Associates, Sunderland, MA.
- Isaac NJB, Turvey ST, Collen B, Waterman C, Baillie JEM (2007) Mammals on the EDGE: conservation priorities based on threat and phylogenv. PLoS One, 2, e296.
- Ives CD, Lentini PE, Threlfall CG et al. (2016) Cities are hotspots for threatened species. Global Ecology and Biogeography, 25, 117–126.
- Jetz W, Thomas GH, Joy JB, Hartmann K, Mooers AO (2012) The global diversity of birds in space and time. Nature, 491, 444–448.
- Jetz W, Thomas GH, Joy JB, Redding DW, Hartmann K, Mooers AO (2014) Global distribution and conservation of evolutionary distinctness in birds. *Current Biology*, 24, 919–930.
- Klotz S, Kühn I (2010) Urbanisation and alien invasion. In: Urban Ecology (ed. Gaston KJ), pp. 120–133. Cambridge University Press, Cornwall.
- Knapp S, Kühn I, Schweiger O, Klotz S (2008) Challenging urban species diversity: contrasting phylogenetic patterns across plant functional groups in Germany. *Ecology Letters*, **11**, 1054–1064.
- Knop E (2016) Biotic homogenization of three insect groups due to urbanization. Global Change Biology, 22, 228–236.
- Kühn I, Klotz S (2006) Urbanization and homogenization comparing the floras of urban and rural areas in Germany. *Biological Conservation*, **127**, 292–300.
- Lepczyk C, Warren P (2012) Urban Bird Ecology and Conservation. University of California Press, London.
- Luck G, Smallbone L (2010) Species diversity and urbanisation: patterns, drivers and implications. In: Urban Ecology (ed. Gaston KJ), pp. 88–119. Cambridge University Press, Cornwall.
- Mace GM, Gittleman JL, Purvis A (2003) Preserving the tree of life. *Science*, **300**, 1707–1709. Magurran A (2004) *Measuring Biological Diversity*. Blackwell Science, Oxford.
- Mannion PD, Upchurch P, Benson RBJ, Goswami A (2014) The latitudinal biodiversity gradient through deep time. *Trends in Ecology and Evolution*, 29, 42–50.
- Marzluff JM (2001) Worldwide urbanization and its effects on birds. Avian Ecology and Conservation in an Urbanizing World (eds Marzluff JM, Bowman R, Donnelly R), pp. 19–47. Springer US, Boston, MA.
- Marzluff J, Bowman R, Donnelly R (2001) Avian Ecology and Conservation in an Urbanizing World. Springer Science, New York, NY.
- Mazerolle MJ (2016) AICcmodavg: model selection and multimodel inference based on (Q)AIC(c). R package.
- McDonnell MJ, Hahs AK (2008) The use of gradient analysis studies in advancing our understanding of the ecology of urbanizing landscapes: current status and future directions. *Landscape Ecology*, 23, 1143–1155.
- McDonnell MJ, MacGregor-Fors I, von der Lippe M et al. (2016) The ecological future of cities. Science, 352, 936–938.
- McKinney ML (2002) Urbanization, biodiversity, and conservation. *BioScience*, **52**, 883.
- McKinney ML (2006) Urbanization as a major cause of biotic homogenization. Biological Conservation, 127, 247–260.
- McKinney ML (2008) Effects of urbanization on species richness: a review of plants and animals. Urban Ecosystems, 11, 161–176.
- McPhearson T, Pickett STA, Grimm NB *et al.* (2016) Advancing urban ecology toward a science of cities. *BioScience*, **66**, 198–212.
- Miller JR, Hobbs RJ (2002) Conservation where people live and work. Conservation Biology, 16, 330–337.
- Møller AP (2008) Flight distance of urban birds, predation, and selection for urban life. Behavioral Ecology and Sociobiology, 63, 63–75.
- Møller A (2014) Behavioural and ecological predictors of urbanization. In: Avian Urban Ecology (eds Gil D, Brumm H), pp. 54–68. Oxford University Press, Oxford.

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- Monnet AC, Jiguet F, Meynard CN, Mouillot D, Mouquet N, Thuiller W, Devictor V (2014) Asynchrony of taxonomic, functional and phylogenetic diversity in birds. *Global Ecology and Biogeography*, 23, 780–788.
- Morelli F, Benedetti Y, Ibáñez-Álamo JD, Jokimaki J, Mänd R, Tryjanowski P, Møller AP (2016) Evidence of evolutionary homogenization of bird communities in urban environments across Europe. *Global Ecology and Biogeography*, 25, 1284–1293.
- Müller N, Ignatieva M, Nilon C, Werner P (2013) Patterns and trends in urban biodiversity and landscape design. In: Urbanization, Biodiversity and Ecosystem Services: Challenges and Opportunities: A Global Assessment (eds Elmqvist T, Fragkias M, Goodness J, Güneralp B, Marcotullio PJ, McDonald RI, Parnell S, Schewenius M, Sendstad M, Seto KC, Wilkinson C), pp. 123–174. Springer Netherlands, Amsterdam.
- Newbold T, Hudson LN, Hill SLL et al. (2015) Global effects of land use on local terrestrial biodiversity. Nature, 520, 45–50.
- Newton I (2003) Speciation and Biogeography of Birds. Academic Press, London.
- Olden JD, Poff NL (2003) Toward a mechanistic understanding and prediction of biotic homogenization. *The American Naturalist*, **162**, 442–460.
- Olden JD, Poff NL, Douglas MR, Douglas ME, Fausch KD (2004) Ecological and evolutionary consequences of biotic homogenization. *Trends in Ecology and Evolution*, 19, 18–24.
- Palomino D, Carrascal LM (2006) Urban influence on birds at a regional scale: a case study with the avifauna of northern Madrid province. *Landscape and Urban Plan*ning, 77, 276–290.
- Pautasso M, Böhning-Gaese K, Clergeau P et al. (2011) Global macroecology of bird assemblages in urbanized and semi-natural ecosystems. Global Ecology and Biogeography, 20, 426–436.
- Prum RO, Berv JS, Dornburg A, Field DJ, Townsend JP, Moriarty Lemmon E, Lemmon AR (2015) A comprehensive phylogeny of birds (Aves) using targeted nextgeneration DNA sequencing. *Nature*, **526**, 569–573.
- R Core Team (2016) R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna.
- Redding DW, Dewolff CV, Mooers AØ (2010) Evolutionary distinctiveness, threat status, and ecological oddity in primates. Conservation Biology, 24, 1052–1058.
- Reif J, Prylová K, Šizling AL, Vermouzek Z, Štastný K, Bejček V (2013) Changes in bird community composition in the Czech Republic from 1982 to 2004: increasing biotic homogenization, impacts of warming climate, but no trend in species richness. *Journal of Ornithology*, **154**, 359–370.
- Rodrigues ASL, Brooks TM, Rodrigues ASL, Brooks TM (2007) Shortcuts for biodiversity conservation planning: the effectiveness of surrogates. *Annual Review of Ecol*ogy, Evolution and Systematics, 38, 713–737.
- Secretariat of the Convention on Biological Diversity (2012) Cities and Biodiversity Outlook: A Global Assessment of the Links between Urbanization, Biodiversity, and Ecosystem Services. Secretariat of the Convention on Biological Diversity, Montreal.
- Şekercioğlu CH (2006) Increasing awareness of avian ecological function. Trends in Ecology and Evolution, 21, 464–471.

- Seress G, Liker A (2015) Habitat urbanization and its effects on birds. Acta Zoologica Academiae Scientiarum Hungaricae, 61, 373–408.
- Seto KC, Güneralp B, Hutyra LR (2012) Global forecasts of urban expansion to 2030 and direct impacts on biodiversity and carbon pools. *Proceedings of the National Academy of Sciences of the United States of America*, **109**, 16083–16088.
- Shochat E, Lerman SB, Anderies JM, Warren PS, Faeth SH, Nilon CH (2010) Invasion, competition, and biodiversity loss in urban ecosystems. *BioScience*, 60, 199–208.
- Sinclair ARE, Mduma SAR, Arcese P (2002) Protected areas as biodiversity benchmarks for human impact: agriculture and the Serengeti avifauna. Proceedings of the Royal Society of London B: Biological Sciences, 269, 2401–2405.
- Skaug H, Fournier D, Nielsen A (2013) glmmADMB: generalized linear mixed models using AD Model Builder – R Package.
- Sol D, González-Lagos C, Moreira D, Maspons J, Lapiedra O (2014) Urbanisation tolerance and the loss of avian diversity. *Ecology Letters*, 17, 942–950.
- Threlfall CG, Williams NSG, Hahs AK, Livesley SJ (2016) Approaches to urban vegetation management and the impacts on urban bird and bat assemblages. *Landscape* and Urban Planning, 153, 28–39.
- Tucker CM, Cadotte MW, Carvalho SB et al. (2016) A guide to phylogenetic metrics for conservation, community ecology and macroecology. *Biological Reviews*, in press. doi: 10.1111/brv.12252
- United Nations (2012) World Urbanization Prospects: The 2011 Revision. United Nations, New York, NY.
- United Nations (2016) Urbanization and Development: Emerging Futures. World Cities Report 2016. United Nations, Nairobi.
- Venables WN, Ripley BD (2002) Modern Applied Statistics with S (4th edn). Springer, New York, NY.
- Webb TJ, Gaston KJ (2000) Geographic range size and evolutionary age in birds. Proceedings of the Royal Society of London B: Biological sciences, 267, 1843–1850.
- Winter M, Devictor V, Schweiger O (2012) Phylogenetic diversity and nature conservation: where are we? *Trends in Ecology and Evolution*, 28, 199–204.

# **Supporting Information**

Additional Supporting Information may be found in the online version of this article:

Table S1. List of sites used and associated information. Table S2. List of species used and associated information. Table S3. List of GLMMs performed in the study.