1	What is an urban bird? Trait-based predictions of urban tolerance for the
2	world's birds are modulated by latitude and human population density
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# 20 Abstract

21 As human density increases, biodiversity must increasingly co-exist with urbanization or face local 22 extinction. Tolerance of urban areas has been linked to numerous functional traits, yet few globally-23 consistent patterns have emerged to explain variation in urban tolerance, which stymies attempts at a 24 generalizable predictive framework. Here, we calculate an Urban Association Index (UAI) for 3768 bird 25 species in 137 cities across all permanently inhabited continents. We then assess how UAI varies as a 26 function of ten species-specific traits and further test whether the strength of trait relationships vary as 27 a function of three city-specific variables. Of the ten species traits, nine were significantly associated 28 with urban tolerance. Urban-associated species tend to be smaller, less territorial, have greater dispersal 29 ability, broader dietary and habitat niches, larger clutch sizes, greater longevity, and have lower 30 elevational limits. Only bill shape showed no global association with urban tolerance. Additionally, the 31 strength of several trait relationships varied across cities as a function of latitude and/or human 32 population density. For example, the effects of body mass and diet breadth are more pronounced at 33 higher latitudes, while the effects of territoriality and longevity were reduced in cities with higher 34 population density. Thus, the importance of trait filters in birds varies predictably across cities, 35 indicating biogeographic variation in selection for urban tolerance that could explain prior challenges in 36 the search for global patterns. A globally-informed framework that predicts urban tolerance will be 37 integral to conservation as increasing proportions of the world's biodiversity are impacted by 38 urbanization.

# 39 Introduction

40 Urbanization is one of the greatest threats to biodiversity (Mcdonald et al. 2008; McKinney 2008; 41 McDonald et al. 2020). By 2030, an expected 5.2 billion people will live in urban areas alone (United 42 Nations 2018) and urban land cover is predicted to exceed 1.2 million km<sup>2</sup> globally (Seto et al. 2012). 43 Urbanization is accompanied by a consistent loss of biodiversity (Aronson et al. 2014; Evans et al. 2018; 44 de Camargo Barbosa et al. 2020), including reduced phylogenetic (Morelli et al. 2016; Sol et al. 2017) 45 and functional diversity (Lizée et al. 2011; Evans et al. 2018; Palacio et al. 2018), resulting in more 46 homogenized wildlife communities. Despite these overall losses, cities can still harbor substantial 47 biodiversity (Spotswood et al. 2021), including threatened species (Ives et al. 2016), with several factors 48 contributing to an increase in species richness within urban areas. For example, biodiversity can be 49 bolstered by green space (Beninde et al. 2015; Callaghan et al. 2018; Fidino et al. 2021), greater habitat 50 heterogeneity (Oliveira Hagen et al. 2017; Souza et al. 2019), higher tree cover (Threlfall et al. 2016; De 51 Castro Pena et al. 2017; Planillo et al. 2021), or reduced housing density (Fontana et al. 2011; Fidino et 52 al. 2021). Within these species pools, some species – often termed urban adapters, urban exploiters, or 53 urban-tolerant species – generally succeed in cities where others do not (Spotswood et al. 2021). The 54 relative tolerance of species to urbanization can result from shared evolutionary history (Iglesias-55 Carrasco et al. 2022) and is often linked to functional traits. For example, in Australian birds, urban 56 adapters show diet generalization, bigger brains and larger clutch sizes (Callaghan et al. 2019). Although 57 many such traits have been suggested or regionally evaluated, what remains untested is whether the 58 traits that confer urban tolerance in species differ across the cities and biogeographic contexts of the 59 world. With recently-available global data on occurrence (Sullivan et al. 2009) and species trait (e.g., 60 AVONET, Tobias *et al.* 2022), birds are an ideal system to explore this question.

Several ecological traits have been linked with urban association in birds (McClure 1989; Sol *et al.* 2014; Callaghan *et al.* 2019). For example, urban tolerance is often positively associated with niche
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63	breadth (Bonier et al. 2007; Evans et al. 2011), including dietary (Croci et al. 2008; Lizée et al. 2011;
64	Morelli et al. 2016) and habitat breadth (Ducatez et al. 2018; Callaghan et al. 2019; Sayol et al. 2020).
65	The degree of sociality also plays a role, with urban-tolerant species tending to be more social (Kark <i>et</i>
66	al. 2007; Croci et al. 2008; Sol et al. 2014). In addition, nest placement is important, with ground nesters
67	often avoiding urban areas (Conole & Kirkpatrick 2011; Evans et al. 2011; Sol et al. 2014; Dale et al.
68	2015; Guetté et al. 2017) while tree nesters tend to persist in cities (Conole & Kirkpatrick 2011; Dale et
69	al. 2015). Yet, despite some general trends, the importance of certain traits often varies between
70	studies. For example, although urban-associated species tend to have larger clutch sizes (Møller 2009;
71	Lizée <i>et al.</i> 2011; Callaghan <i>et al.</i> 2019), this pattern is not always supported (Croci <i>et al.</i> 2008;
72	Chamberlain et al. 2009), and may be mediated by other life-history traits (Sayol et al. 2020). Similarly,
73	the role of body size has also received mixed support, with urban tolerance positively associated with
74	body mass in Australia (Callaghan <i>et al.</i> 2019), but showing no relationship globally (Sol <i>et al.</i> 2017).
75	Longevity or lifespan has seldom received strong support in models (Croci et al. 2008; Guetté et al.
76	2017), while cavity nesters show mixed responses to urban areas (Conole & Kirkpatrick 2011; Lizée et al.
77	2011; Dale et al. 2015; Jokimäki et al. 2016; Evans et al. 2018). Finally, although dispersal ability has
78	been linked to urban tolerance (Møller 2009), migratory strategy is rarely associated with urban
79	tolerance (Evans <i>et al.</i> 2011, 2018; Dale <i>et al.</i> 2015; Jokimäki <i>et al.</i> 2016; Guetté <i>et al.</i> 2017; Callaghan <i>et</i>
80	<i>al.</i> 2019; Sayol <i>et al.</i> 2020).

The lack of generality in previous work may arise for multiple reasons. Many studies sample only a subset of biogeographic regions and/or species. Variation in the importance of traits may be driven by differences in species pools or by context-dependent differences in filters between different landscapes (Aronson *et al.* 2016; Oliveira Hagen *et al.* 2017). It thus seems likely that results should differ between biomes due to differences in climate and biogeographic history (Morelli *et al.* 2016; Leveau *et al.* 2017; Filloy *et al.* 2019). Yet, even studies that have taken a global perspective have been biased in their

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87	sampling towards North America, Europe, and Australia, with a distinct lack of data from the tropics
88	(Magle et al. 2012; Sol et al. 2014; Sayol et al. 2020). Moreover, the number of species in global trait
89	studies has also been limited, with the largest sample size (629 species by Sayol et al. 2020))
90	representing only ~6% of bird species found globally. Previous studies have been restricted by the lack
91	of bird occurrence data across urbanization gradients, particularly in the tropics (Magle et al. 2012), but
92	also by access to global trait datasets that have only recently become available.

93 Here, we combine global data on occurrence (>125 million records) from the citizen science 94 project eBird (Sullivan et al. 2009) with a continuous measure of urbanization (night-time lights) to 95 calculate an Urban Association Index (UAI) for 3768 bird species (~35% of extant bird species) in 137 96 cities across six continents and 11 biomes. We chose ten species-specific functional traits with globally 97 available data and hypothesized links to urban tolerance, and modeled UAI values as a function of these 98 traits. We further chose three city-specific landscape variables that we predicted would influence the 99 importance of our traits for urban tolerance (Oliveira Hagen et al. 2017), assessing whether the effects 100 of each trait varied as a function of latitude, human population density, and landscape greenness. We 101 present the first evidence that the importance of different traits for urban tolerance varies predictably 102 across the planet.

103

### 104 Methods

105 Data filtering

We downloaded the global eBird basic dataset (Sullivan *et al.* 2009) including all records up until
February, 2022 (v1.14). We restricted the dataset to the years 2002–2021 – the 20 complete years
before present. We then limited eBird protocol types to "traveling", "stationary", and "area" and
removed incomplete checklists. Following eBird best practices (Johnston *et al.* 2021), we removed
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110	checklists with >10 observers, with durations >5 hr, with distances >5 km (for "traveling" protocol), and
111	with areas >500 ha (for "area" protocol). For group checklists involving duplicate records, we randomly
112	retained one checklist per group. Finally, we removed records that were not identified to species level,
113	including all hybrids, intergrades, "slashes" (e.g., "Greater/Lesser Yellowlegs"), indefinite species (e.g.,
114	"hummingbird sp."), and domestics. Although many of the species in our dataset are introduced in some
115	cities, they are native in others, so we did not remove or classify species based on being exotic (e.g.,
116	Passer domesticus, Sturnus vulgaris). We made a single exception to these exclusions, retaining the
117	widespread and ubiquitous Feral (Rock) Pigeon (Columba livia) despite having been domesticated as it is
118	a key avian species in many cities. Initially we considered including all species found in cities but
119	restricted our dataset to exclude water birds (~15% of the species set) since they have substantially
120	different natural histories and traits compared to land birds, following Callaghan et al. (2019).
121	City selection
122	From the data repository OpenDataSoft, we downloaded the dataset "Geonames – All Cities with a
122 123	From the data repository OpenDataSoft, we downloaded the dataset "Geonames – All Cities with a population > 1000" (OpenDataSoft 2022), and reduced the dataset down to cities with a population
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123 124	population > 1000" (OpenDataSoft 2022), and reduced the dataset down to cities with a population >100,000, yielding 4643 cities. We chose this relatively low population cut-off to include smaller, remote
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123 124 125 126 127 128 129	population > 1000" (OpenDataSoft 2022), and reduced the dataset down to cities with a population >100,000, yielding 4643 cities. We chose this relatively low population cut-off to include smaller, remote cities in ecologically distinct regions – including Darwin (Australia), Port Louis (Mauritius), and Reykjavík (Iceland). We then calculated the pairwise distance between every city using the package <i>geodist</i> (Padgham & Sumner 2019). Starting with the cities with the largest populations, we sequentially removed all smaller cities within 500 km of the larger city in order to produce a set of non-overlapping, spatially-independent cities. This algorithm retained 289 cities separated by at least 500 km (Fig. S1).
123 124 125 126 127 128 129 130	population > 1000" (OpenDataSoft 2022), and reduced the dataset down to cities with a population >100,000, yielding 4643 cities. We chose this relatively low population cut-off to include smaller, remote cities in ecologically distinct regions – including Darwin (Australia), Port Louis (Mauritius), and Reykjavík (Iceland). We then calculated the pairwise distance between every city using the package <i>geodist</i> (Padgham & Sumner 2019). Starting with the cities with the largest populations, we sequentially removed all smaller cities within 500 km of the larger city in order to produce a set of non-overlapping, spatially-independent cities. This algorithm retained 289 cities separated by at least 500 km (Fig. S1). After identification of these target urban areas around the world, we filtered the eBird dataset to

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134	<0.01% of all occurrences per city. The first filter ensured a minimum data requirement while the second
135	filter was a threshold intended to filter out vagrant species while retaining scarce but expected species.
136	As some cities lacked 100 records for even one species, we removed any city with <50 species remaining
137	after restricting species to ≥100 records, such that all remaining cities had ≥5000 bird records. This 50-
138	species threshold was chosen in order to remove cities that contained only a handful of species that
139	would tend to be more urban associated (high UAIs), but to retain cities in environments with low
140	species richness (e.g., boreal regions) that would have been removed if the threshold was 100 species.
141	Our final dataset contained 16,455 UAI estimates representing 127,046,578 eBird occurrence records of
142	3768 species across 137 cities (Fig. 1a).
143	Urbanization association index (UAI)
144	To quantify species' relationships with urban areas we aimed to create a continuous metric of urban
145	association that would avoid the assumptions of using arbitrary thresholds to categorize species based
146	on urban tolerance (Callaghan <i>et al.</i> 2019; Fanelli <i>et al.</i> 2022). Following Callaghan et al. (2019), we
147	downloaded the VIIRS night-time lights imagery (Annual VNL V1; Elvidge et al. 2017), a composite global
148	image of night-time lights for the year 2016. Although 2016 is not the mid-point of the eBird data
149	(2012), the number of eBird records has increased exponentially so 2016 is close to the mean year of all
150	checklists (2017). We chose light radiance as a proxy for urbanization because it is available as a
151	continuous measure across the world, it is a close proxy for human population density (Elvidge et al.
152	2017), and, when combined with eBird data, it is correlated with other measures of urban tolerance in
153	birds (Callaghan et al. 2021). From this imagery, we extracted the radiance value for every eBird
154	checklist locality. As these radiance values start at 0 (total darkness) and increase exponentially, we
155	added 1 then log-transformed all radiance values to reduce the leverage of extremely bright buildings.
156	Then, for every species within each city, we calculated the mean radiance value of all occurrence
157	records. We chose to use the mean instead of the median because we found that many species had a

157 records. We chose to use the mean instead of the median because we found that many species had a

median radiance of 0, as they occurred predominantly in non-urban areas. Moreover, the distribution of
mean radiance values was fairly normal (Fig. S2), while the distribution of median radiance values was
heavily right skewed. We also tested our chosen metric for sensitivity to mismatches in scale between
the spatial resolution of VIIRS and checklists (Supporting Information) but decided to retain the metric
as described. Thus, our Urban Association Index (UAI) for each species is the mean of the transformed
radiance values across eBird records where the radiance value of each record is taken from a single pixel
of night-time lights.

165 Species traits

We chose species-specific functional traits that have been linked to urban tolerance in the past and/or traits that we hypothesized would predict urban tolerance that have not been tested globally. We chose traits that were available for the entire species set and, where possible, we chose numerical (rather than categorical) traits in order to reduce the number of parameters estimated. We therefore did not use traits such as residual brain size where data does not exist for all species (Sayol *et al.* 2020) and excluded categorical traits with many levels, such as primary diet. Traits for every species were then extracted from several datasets, as follows (Table 1).

173 From the publicly available AVONET (Tobias et al. 2022) we extracted body mass, four bill 174 measurements (length from culmen, length from nares, width, and depth), and hand-wing index (HWI). 175 These data were complete for all species. To reduce the four bill measurements down to a single axis, 176 we conducted a PCA on the variables and extracted the second principal component, ignoring the first 177 principal component, which is highly correlated with body size (Pigot et al. 2020). This second principal component – which we refer to as "bill shape" – represents a spectrum from long, thin, pointy bills (e.g., 178 179 *Ensifera ensifera*) to short, thick bills (e.g., *Callocephalon fimbriatum*), a spectrum associated with 180 foraging specializations (Pigot et al. 2020). As a measure of dispersal ability, HWI has not been tested as

181

a global predictor of urban tolerance but is highly correlated with several ecological factors, including 182 primary diet and habitat type (Sheard et al. 2020).

183 From the dataset BirdBase (Sekercioğlu et al. 2004; Buechley et al. 2019) we extracted diet 184 breadth, habitat breadth, lower elevational limit, clutch size, and nest shape/substrate. Diet breadth is 185 the number of major food groups (1-9) that a species eats (e.g., invertebrates, fruit, seeds) while habitat 186 breadth (1–11) is the number of major habitats where a species is found (e.g., forest, grasslands, 187 desert). Lower elevational limit was included because we hypothesized that cities – which tend to be 188 found non-randomly at lower elevations (Luck 2007) – would favor species that occur at lower 189 elevations. Nest shape and nest substrate were originally sourced as two separate data columns, but we 190 collapsed these into one. As there was no way to define these nests numerically by shape and substrate, 191 we instead defined four categories: ground (nests of any form located on the ground), cavity (nests 192 above ground in cavities or crevices), open (nests above ground with open tops such as cups, saucers, 193 and platforms), and enclosed (nests above ground with entrance holes such as spheres, pendants, and 194 domes). Clutch size data were augmented with data from an existing published dataset (Myhrvold et al. 195 2015), while further gaps in BirdBase variables were filled using the online database Birds of the World 196 (Billerman et al. 2020). Where information was lacking for a species, missing values were inferred from 197 close extant relatives with complete data. Finally, longevity (a measure of lifespan) and territoriality (a 198 scale from 1 to 3 where 3 is more territorial) were extracted from published datasets (Tobias et al. 2016; 199 Bird *et al.* 2020). Once assembled, we had complete data for ten functional traits (Table 1). 200 Trait variables were transformed, as necessary, prior to analysis. Given expected non-linear

201 relationships, we took the log of body mass, longevity, and clutch size. We then scaled and centered all 202 numerical traits to have a mean of 0 and a standard deviation of 1.

203 City variables

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Urban birds, traits, and landscapes

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204	For each 100 km-radius city circle we gathered data on three covariates that we hypothesized would
205	alter the importance of traits: latitude, greenness, and population density. We chose numerical
206	covariates in order to reduce the number of parameters, as each new city covariate adds nine
207	parameters (one for each numerical trait) to the model. However, combined, latitude and greenness
208	cover much of the variation among biomes (Fig. 1b).

209 Many factors vary with latitude including climate, species richness, and human development, so 210 there are many possible avenues through which latitude could affect urban tolerance. For example, the 211 stability of tropical climate and ecosystems (Janzen 1967) may mean stronger filters in urban areas 212 against ecological specialists in the tropics compared to temperate regions (Newbold *et al.* 2013). We 213 extracted the latitude of each city from the same Geonames dataset as the city populations.

214 The amount of greenness in a city – whether tree cover or vegetation diversity – is an important 215 predictor of bird diversity in cities (Beninde et al. 2015; Threlfall et al. 2016; De Castro Pena et al. 2017; 216 Callaghan et al. 2018; Souza et al. 2019; Planillo et al. 2021). Moreover, overall greenness of the 217 landscape depends on the primary habitat. For example, desert cities such as Phoenix (USA) and Dubai 218 (UAE) are greener than the surrounding landscape while forest cities such as Iquitos (Peru) and Nashville 219 (USA) are less green than the surroundings. We thus hypothesized that the amount of greenery would 220 also alter trait filters (Oliveira Hagen et al. 2017). For example, less green landscapes with fewer 221 resources may select for habitat generalists or more mobile species. We used NDVI as a measure of the 222 greenness of each city, derived from the MOD13A3 product (Didan 2021). This product provides 1km 223 monthly NDVI values globally, excluding water bodies. We calculated the mean NDVI values within the 224 100 km buffer of each city for each month for the year 2021, and retained the maximum NDVI value. We 225 used the maximum NDVI value as each city has a different seasonal cycle over which greenness is likely 226 to vary (i.e., greenness peaks in some cities in August, while in January in others).

227	Human population density has been linked to taxonomic and functional diversity in cities
228	(Fontana et al. 2011; Oliveira Hagen et al. 2017). We hypothesized that cities with higher population
229	densities may present strong selection pressures against species that are, for example, larger with
230	narrower diets. To obtain population density (number of people/cell), we downloaded Gridded
231	Population of the World data from the Center for International Earth Science Information Network
232	(CIESIN 2018). The data are available on 5-year intervals between 2000–2020. We used 30 arc-second
233	resolution population size for the year 2015 as the year closest to the VIIRS imagery and the mean year
234	of eBird records. We buffered city midpoints by 100 km and extracted the mean value of the gridded
235	density data within each buffer.

For the models, we calculated the absolute value of latitude and the log of population density.All three city covariates were then scaled and centered.

238 Modeling

We modeled UAI values as a function of traits and city variables in a Bayesian hierarchical framework that accounted for the random effects of city and species. We modeled the effect of the ten species traits on UAI with the following structure:

242 
$$y_{ij} \sim Gamma(\zeta_j, \frac{\zeta_j}{e^{\mu_{ij}}})$$

243 
$$\mu_{ij} = \alpha_{ij} + \delta_1 \cdot enclosed + \delta_2 \cdot cavity + \delta_3 \cdot ground + \delta_2 \cdot cavity + \delta_3 \cdot ground + \delta_3 \cdot grou$$

244 
$$\beta_{1,j} \cdot mass + \beta_{2,j} \cdot beak + \beta_{3,j} \cdot HWI + \beta_{4,j} \cdot DB + \beta_{5,j} \cdot HB + \beta_{5,j} \cdot$$

245 
$$\beta_{6,j} \cdot elevation + \beta_{7,j} \cdot territory + \beta_{8,j} \cdot longevity + \beta_{9,j} \cdot clutch,$$

where the estimated mean UAI, y<sub>ij</sub>, for species *i* in city *j* was modeled as a gamma-distributed random

variable with a city-specific shape parameter  $\zeta_j$  and a rate parameter equal to  $\frac{\zeta_j}{e^{\mu_{ij}}}$ . We chose a gamma

248	distribution to reflect the fact that our response variable was bounded by 0 on the lower end and right
249	skewed. The shape of the distribution was allowed to vary among cities to accommodate variation in the
250	data. In turn, $\mu_{ij}$ was modeled as a linear combination of an intercept for open nesters, $lpha_{ij}$ , three
251	differences in intercepts ( $\delta_1$ to $\delta_3$ ) and nine covariates with corresponding parameters ( $eta_{1,j}$ to $eta_{9,j}$ ). The
252	parameters $\delta_1$ to $\delta_3$ represent the difference in UAI for three dummy variables ( <i>enclosed, cavity,</i> and
253	<i>ground</i> ) that together encode the three other nest types, where all three covariates are binary (1 =
254	species' nest type, 0 = otherwise) and mutually exclusive. The parameters $eta_{1,j}$ to $eta_{9,j}$ represent the
255	slopes of the effects of nine numerical traits on $\mu_{ij}$ .
256	The intercept $\alpha_{ij}$ can be further decomposed,
257	$\alpha_{ij} = \gamma + \eta_i + \omega_j,$
258	into a global intercept, $\gamma$ , and the random effects of species, $\eta_i$ , and city, $\omega_j$ . The random effect of
259	species accounts for species being represented across multiple cities. The random effect of city allows
260	species in different cities to have different average UAIs based on unmodeled factors such as differences
261	in brightness, development, and birdwatching effort. The random effects of species,

262 
$$\eta_i \sim t(0, \sigma_{\eta}, \kappa),$$

were drawn from a t-distribution with a mean 0, standard deviation  $\sigma_{\eta}$ , and degrees of freedom  $\kappa$ (which controls the degree to which the distribution resembles a normal, as  $\kappa$  approaches infinity, or a Cauchy, as  $\kappa$  approaches 1). The choice of t-distribution allowed for fatter tails in the distribution of intercepts across species. The random effects of city,

267 
$$\omega_j \sim Normal(0, \sigma_{\omega}),$$

268 were drawn from normal distributions with mean 0 and standard deviation and  $\sigma_{\omega}$ .

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269 In order to integrate the city covariates with the functional traits, every  $\beta_j$  parameter for trait *k* 270 was drawn from a normal distribution

271 
$$\beta_{kj} \sim Normal(v_{kj}, \sigma_{\beta_k})$$

with a mean  $v_{kj}$  and a standard deviation  $\sigma_{\beta_k}$  such that each  $\beta_{kj}$  had its own process error, to accommodate variation in the data. The mean was then modeled as a linear combination of city covariates

275 
$$v_{kj} = \varphi_k + \theta_{1,k} \cdot latitude + \theta_{2,k} \cdot density + \theta_{3,k} \cdot greenness,$$

such that the effect of each numerical trait on UAI varied as a function of the city-level covariates.
Importantly, this allowed urban tolerance to be predicted differently by different traits in different
geographical contexts.

279 We ran this Bayesian model using the program JAGS (Plummer 2003) via the R package R2jags (Su & Yajima 2021). We used vague priors (mean of 0, standard deviation of 100) and we ran three 280 281 chains, each with 40,000 iterations, beginning with a burn-in of 10,000 followed by a thinning of 30, 282 retaining 1000 posteriors per chain. We verified that the model had successfully converged (Rhat  $\leq$  1.01 283 and n.eff > 400). We performed posterior predictive checks to ensure that data generated by the model were similar to data used to fit the model (Gelman et al. 2000). We found that 49.4% of the means of 284 285 the posteriors were less than the observed mean UAI, indicating that our model could successfully 286 reproduce the mean UAI. From the parameter posteriors we calculated the means and 95% credible 287 intervals for each parameter. Due to Bayesian shrinkage within the random effects framework, post hoc testing indicated that the model was less able to estimate suitable species-specific intercepts,  $\eta_i$ , for 288 289 species with only a few data points – i.e., those species present in only one or a few cities. We therefore 290 fit a second, identically-structured model using a dataset where species represented in <5 cities were

excluded, and checked for consistency of model results (i.e.,  $\beta$ s) as compared to the original, full dataset model.

293 Phylogenetic structure

294 In order to test for phylogenetic signal in UAI values, we aligned the eBird (Clements) taxonomy with the 295 taxonomy of BirdTree.org (Jetz et al. 2012) and downloaded 100 phylogenetic trees with the Hackett 296 backbone. We averaged UAI values across the tips of the phylogeny. For each tree, we calculated Pagel's 297  $\lambda$  as a measure of phylogenetic signal using the package *phytools* (Revell 2012). We then calculated the 298 mean  $\lambda$  across trees, with associated 95% quantiles. Directly incorporating these phylogenies into our 299 Bayesian model was not practical due to the extensive computational time required for an analysis that 300 includes so many species (over a year). Rather, we tested whether model residuals, averaged at the 301 species level, contained phylogenetic signal (Revell 2010). This test would tell us whether there was 302 unexplained variation in the model associated with phylogeny. We also tested for signal in the residuals 303 of the model with the reduced species set.

304

# 305 Results

Our analysis included 16,455 UAI estimates representing data from >125 million eBird records across
 137 cities (Fig. 1a). This list comprised cities from 62 countries including 39 in North America, 28 in South
 America, 27 in Asia, 22 in Europe, 10 in Africa, and 10 in Australasia. Together, these cities span 11 of
 the world's 14 terrestrial biomes (Olson & Dinerstein 1998). The number of avian species meeting the
 inclusion criteria in each city ranged from 56 in Naha (Japan) to 533 in Bogotá (Colombia).
 Of the 3768 species for which we calculated UAI, the five species present in the most urban

areas (see Fig. S4 for top 30) were Feral (Rock) Pigeon (*Columba livia*), House Sparrow (*Passer* 

domesticus), Barn Swallow (*Hirundo rustica*), Osprey (*Pandion haliaetus*), and Peregrine Falcon (*Falco peregrinus*). Across species, UAI values ranged from 0 (for 46 species) to 3.97 (Yellow-crested Cockatoo – *Cacatua sulphurea* – a species introduced to Hong Kong) with a mean of 1.14. Of species present in at
least ten cities, the top-five species with the highest UAI (see Fig. S5 for top 30) were Monk Parakeet
(*Myiopsitta monachus*), Rose-ringed Parakeet (*Psittacula krameri*), Yellow-chevroned Parakeet
(*Brotogeris chiriri*), Feral Pigeon, and Sayaca Tanager (*Thraupis sayaca*).

319 There was considerable phylogenetic signal in UAI across species (Fig. 2;  $\lambda$  = 0.61, CI = 0.56– 320 0.65). Notable families with high average UAI values, indicating broad urban associations, included 321 Sturnidae (starlings; 1.75 ± 0.13 SE, n<sub>cities</sub> = 40), Apodidae (swifts; 1.61 ± 0.12, n<sub>cities</sub> = 44), Hirundinidae 322 (swallows;  $1.55 \pm 0.09$ ,  $n_{cities} = 52$ ), Psittacidae (parrots;  $1.55 \pm 0.11$ ,  $n_{cities} = 86$ ), and Icteridae (New World 323 orioles and blackbirds; 1.47 ± 0.08, n<sub>cities</sub> = 80). Notable families with low average UAI included Pipridae 324 (manakins; 0.33 ± 0.07, n<sub>cities</sub> = 21), Petroicidae (Australasian robins; 0.38 ± 0.09, n<sub>cities</sub> = 20), Trogonidae (trogons; 0.45 ± 0.07, n<sub>cities</sub> = 24), Thamnophilidae (antbirds; 0.55 ± 0.06, n<sub>cities</sub> = 72), and Tinamidae 325 326 (tinamous;  $0.58 \pm 0.09$ ,  $n_{cities} = 22$ ).

Of the ten species-specific traits considered, all except bill shape were significantly associated with UAI (Figs. 3,4). Body mass (Fig. 3a), lower elevational limit (Fig. 3e), territoriality (Fig. 3f), and ground nesting (Fig. 3i) were negatively associated with UAI, while hand-wing index (HWI; Fig. 3b), diet breadth (Fig. 3c), habitat breadth (Fig. 3d), longevity (Fig. 3g), and clutch size (Fig. 3h) were positively associated with UAI. In other words, more urban-tolerant species are smaller, tree- or building-nesting species with higher dispersal ability, wider diet and habitat breadth, lower elevational limits, lower territoriality, longer lifespan, and greater clutch size.

334 Seven of the traits varied significantly as a function of city-level covariates (Fig. 4,5). In terms of 335 latitude (Fig. 4b), the negative effect of body mass on UAI (Fig. 5a) and the positive effects of diet

336	breadth (Fig. 5b) and longevity (Fig. 5d) became more pronounced in cities at higher latitudes.
337	Contrastingly, the positive effect of habitat breadth (Fig. 5e) and the negative effect of lower elevational
338	range limit (Fig. 5g) on UAI became more pronounced in tropical cities. The effect of bill shape on UAI –
339	which showed no globally consistent relationship – varied with latitude (Fig. 5h) such that species with
340	longer, pointier bills were more urban tolerant at higher latitudes while species with shorter, thicker bills
341	were more urban tolerant at lower latitudes. In terms of population density (Fig. 4d), the negative effect
342	of territoriality (Fig. 5c) and the positive effects of diet breadth (Fig. 5f) and longevity (Fig. 5i) became
343	more pronounced in cities with lower population density. Finally, none of the nine numerical traits
344	varied significantly in effect as a function of landscape greenness (Fig. 4c).
345	Model residuals contained relatively low phylogenetic signal ( $\lambda$ = 0.37, CI = 0.33–0.43). Most of
346	this signal resulted from species with few data points – i.e., those represented in 1–4 cities – as the
347	model was less able to estimate suitable species-specific intercepts due to the shrinkage of intercept
348	parameters towards the cross-species mean. Removing these 2848 species (76% of the species set) and
349	re-running the model produced qualitatively similar trait coefficients (Fig. S6) and greatly reduced the
350	phylogenetic signal in the residuals ( $\lambda$ = 0.15, CI = 0.09–0.22). Thus, we are confident that our estimates
351	of the effect of traits on UAI are robust to potential phylogenetic or sample-based biases.

352

# 353 Discussion

354 Many studies have linked species-specific functional traits to urban tolerance (Møller 2009; Sol *et al.* 

2014; Callaghan *et al.* 2019; Sayol *et al.* 2020) but none have tested for interactions between traits and

- 356 geographic factors, especially not at the global taxonomic and spatial scale we employ here. For 35% of
- the world's bird species across 137 cities and 11 biomes including regions of the world
- underrepresented in ecological studies (i.e., Asia, Africa, South America; Magle et al. 2012; Estes et al.

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359 2018) – we find that nine different functional traits are related to urban association. Furthermore, we 360 find that two geographic variables – latitude and human population density – significantly modulate the 361 effects of seven of these traits, meaning that the strength of trait-based filters in urban environments 362 varies systematically across the planet (Aronson et al. 2016; Oliveira Hagen et al. 2017). Our study is the 363 first at a global scale to demonstrate the effects of body size, hand-wing index (HWI), diet breadth, 364 lower elevational limit, territoriality, longevity, and clutch size on urban association, and confirms the 365 positive association of habitat breadth and the negative association of ground nesting (Sol et al. 2014; 366 Ducatez et al. 2018).

367 Urban associated species tended to have wider diet and habitat breadths (Fig. 3c,d), confirming 368 the role of ecological generalism in urban tolerance (Bonier et al. 2007; Ducatez et al. 2018; Callaghan et 369 al. 2019; Fidino et al. 2022). As cities erase or erode most native habitats (McDonald et al. 2020), 370 ecological specialists are less able to survive while more versatile species persist. However, we found 371 that the importance of diet and habitat breadth had opposing patterns across latitude (Fig. 5b,d). 372 Habitat breadth was more important in tropical urban areas, possibly because most tropical land birds 373 have high forest dependency (Tobias et al. 2013), and thus are more likely to experience a stronger filter 374 in urban areas (Newbold et al. 2013). But, with fewer habitats to specialize on towards the poles, habitat 375 breadth becomes less important at higher latitudes. By contrast, diet breadth was more important in 376 temperate areas. Many urban-associated tropical birds are dietary specialists, particularly nectarivores 377 and frugivores, where they take advantage of plentiful year-round fruiting and flowering ornamental 378 trees (Lim & Sodhi 2004). Temperate cities, with seasonal resource pulses and troughs, favor omnivores 379 that can make use of a wide variety of food sources (Croci et al. 2008; Lizée et al. 2011; Jokimäki et al. 380 2016; Evans et al. 2018).

381 Related to diet, the effect of beak shape on urban associations changed sign with latitude (Fig.
382 5f). In the tropics, species with short, thick bills were favored in urban areas, a result that may be
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383 explained by the abundance of specialist frugivores in fruit-plentiful tropical cities, exemplified by urban-384 tolerant parrots (Cacatuidae, Psittaculidae, Psittacidae). At more temperate latitudes, species with 385 short, stubby bills tend to be granivores and also tend to avoid urban areas where grasses are cut short. 386 While the occassional short, stubby bill does well in temperate urban environments (e.g., House 387 Sparrow or House Finch, Haemorhous mexicanus), many temperate granivores such as game birds 388 (Phasianidae), longspurs (Calcariidae), and grassland sparrows (Passerellidae) require suitable habitat far 389 from development (Croci et al. 2008; Callaghan et al. 2019). Meanwhile, the hummingbirds (Trochilidae) 390 - long-billed species with high data leverage - present an interesting outlier. In the Neotropics, where 391 their diversity peaks, only a fraction of species are found in urban areas (Maruyama et al. 2019) such as 392 Panama City, while in North America, most hummingbird species frequent urban feeders (Greig et al. 393 2017; Miller et al. 2017). Variation in the importance of bill shape is clearly complex, underscoring the 394 diverse responses of different feeding guilds to urbanization (Kark et al. 2007; Jokimäki et al. 2016; 395 Evans et al. 2018; Callaghan et al. 2019).

396 Previous studies have suggested that migratory strategy was not associated with urban 397 tolerance (Dale et al. 2015; Jokimäki et al. 2016; Guetté et al. 2017; Evans et al. 2018; Callaghan et al. 398 2019; Sayol et al. 2020), while only one study has found dispersal ability per se to be related to urban 399 associations (Møller 2009). We, however, found that species with higher HWI, i.e., longer, more pointed 400 wings associated with greater dispersal ability (Sheard et al. 2020), have higher UAI values (Fig. 3b). 401 Although dispersal ability is positively associated with migratory capacity (Sheard et al. 2020), previous 402 studies focusing on temperate cities may not have found a role for migratory capacity as migrants tend 403 to broaden their habitat use to include cities on their wintering grounds in the tropics. Additionally, this 404 pattern could be driven by a number of factors, including the sensitivity of low-dispersal species to 405 anthropogenic change (Claramunt et al. 2022), and the association between HWI and specific foraging 406 modes such as flycatching, aerial insectivory, frugivory, and nectarivory (as opposed to gleaning,

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407 terrestrial insectivory etc.) that would be favored in urban environments (Lees & Peres 2009; Sheard *et*408 *al.* 2020).

409 The role of body size in urban tolerance has mixed support, including a positive association in 410 Australia (Callaghan et al. 2019) but no effect globally across 358 species (Sol et al. 2014). Here, we 411 found urban-tolerant species are significantly smaller, an effect (Fig. 3a) that strengthens towards the 412 poles (Fig. 5a). Many families of large species, such as bustards (Otididae), tinamous, and pheasants, 413 appear to be urban avoidant (Fig 2). These species tend also to be cursorial, which could put them at 414 elevated risk of urban-associated predators (e.g., domestic cats; Loss et al. 2013) and nest predators 415 (e.g., rats; Smith et al. 2016). In the tropics, these families of large species might be balanced out by 416 urban-tolerant arboreal-nesting large hornbills (Bucerotidae), turacos (Musophagidae), parrots, and 417 cockatoos (Conole & Kirkpatrick 2011). In temperate regions, game birds are likely selected against in 418 urban areas due to habitat requirements, the history of hunting, or pressure from meso-predators 419 (Crooks & Soule 1999).

420 Body size can be associated with other life-history traits that predict urban tolerance – although 421 we found little correlation between body mass, longevity, and clutch size in this study. Supporting 422 results from other studies (Møller 2009; Lizée et al. 2011; Callaghan et al. 2019), we found that species 423 with larger clutches were more urban tolerant (Fig. 3h). Species with larger clutch sizes tend to live at 424 the faster end of the life-history continuum and may be able to adapt faster to novel environments 425 (Møller 2009). Conversely, however, we found that species with longer lifespans were also more urban 426 tolerant (Fig. 3g), corroborating the finding that urban-tolerant species also have higher annual survival 427 rates (Møller 2009). One possibility is that long-lived species are also more intelligent species (Smeele et 428 al. 2022). The role of brain size in urban tolerance appears linked to other life-history strategies, with big 429 brains important for species with high brood value (i.e., fewer broods over a lifetime) and small brains 430 important for species with low brood value (Sayol et al. 2020). While we lacked the data to test this Page **19** of **42** 

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hypothesis globally across our full species set, our results suggest a similar trade-off, that it helps to
either have large clutch sizes, or live longer in order to learn to exploit urban environments. The
importance of longevity also increases in temperate cities (Fig. 5c), suggesting that living longer and,
perhaps, being smarter are more beneficial closer to the poles.

435 Certain aspects of breeding biology were also tied to urban tolerance. While we did not test 436 sociality per se (a trait which is not available broadly), we did find a significant negative effect of 437 territoriality (Fig. 3f). Urban-tolerant species tend to be more social or gregarious (Kark et al. 2007; Croci 438 et al. 2008; Sol et al. 2014) and therefore less territorial. Being strongly territorial year-round (level 3 on 439 the scale) is usually tied to defense of resources (Tobias et al. 2016) and in resource-poor cities it makes 440 less sense to be territorial and more sense to follow resources more plastically. Where species nest also 441 matters, and we confirm the results of others that ground-nesting species tend to be less urban-tolerant 442 (Conole & Kirkpatrick 2011; Evans et al. 2011; Sol et al. 2014; Dale et al. 2015; Guetté et al. 2017). 443 Species that nest above the ground with open or enclosed nests had the highest urban tolerance (Fig. 444 3i), probably due to safety from predators (Jokimaki & Huhta 2000; Chace & Walsh 2006). Some studies 445 have suggested that cavity nesters would have higher urban tolerance (Chace & Walsh 2006; Croci et al. 446 2008; Conole & Kirkpatrick 2011) while others have suggested the opposite (Evans et al. 2018). We 447 found intermediate UAI values for cavity nesters, perhaps reflecting the contrast of relative success of 448 cavity nesters with lower availability of nest cavities in urban areas (Blewett & Marzluff 2005).

The effects of territoriality, diet breadth, and longevity were all reduced in cities with higher
population density (Fig. 5g–i). As population density is calculated across the whole 100 km radius circle,
it is possible that the most densely populated cities are more homogenous with less non-urban habitat
for urban avoiders. For example, Anchorage (USA) and Reykjavík (Iceland) are small cities surrounded by
wilderness where habitats strongly differ between urban and non-urban areas. In contrast, cities like
Bangkok (Thailand) and İstanbul (Turkey) are vast sprawling metropolises with abundant feral predators
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where there is little room for specialized, long-lived, territorial urban avoiders. Finally, we did not find
that any trait effects varied as a function of landscape greenness, indicating that ecological filters in
urban areas are similar for cities at the same latitude that differ in greenness, a measure driven in large
part by habitat (Fig. 1b).

459 In summary, we found that numerous species-specific functional traits (smaller body size, lower 460 territoriality, greater dispersal ability, broader dietary and habitat niches, larger clutch sizes, greater 461 longevity) predict urban tolerance across the planet. However, many of these trait effects are 462 modulated by landscape-level properties, most notably latitude. Where previous studies have 463 demonstrated the importance of certain traits in certain parts of the world, we demonstrate the 464 importance of geography in determining trait-based urbanization filters (Ferenc et al. 2014; Aronson et 465 al. 2016; Leveau et al. 2017; Filloy et al. 2019) at an unprecedented taxonomic and spatial scale. 466 Moreover, much of the region-specific variation in previous trait-seeking studies could be due to 467 predictable geographic variation in trait strength that varies with latitude and human population 468 density. Studying how traits filter diversity across the globe moves us toward a more predictable 469 framework that will better allow us to understand future biodiversity loss - and how we might mitigate 470 it – given the expected future expansion of urban areas.

471

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Trait	Description	Primary sources
Body mass	Log-transformed	AVONET
Bill shape	Second PC from a PCA of four bill measurements	AVONET
Hand-wing index	The ratio of Kipp's distance to wing length	AVONET
Diet breadth	Number of major food groups, 1–9	BirdBase, Birds of the World
Habitat breadth	Number of major habitats, 1–11	BirdBase, Birds of the World
Lower elevational limit	Lower limit of elevational range reported in the literature	BirdBase, Birds of the World
Territoriality	A scale from 1 (low) to 3 (high)	Tobias et al. 2016
Longevity	Log-transformed	Bird et al. 2020
Clutch size	Log-transformed	BirdBase, Myhrvold et al.
		2015, Birds of the World
Nest type	Categorical: ground, cavity, open, and enclosed	BirdBase, Birds of the World

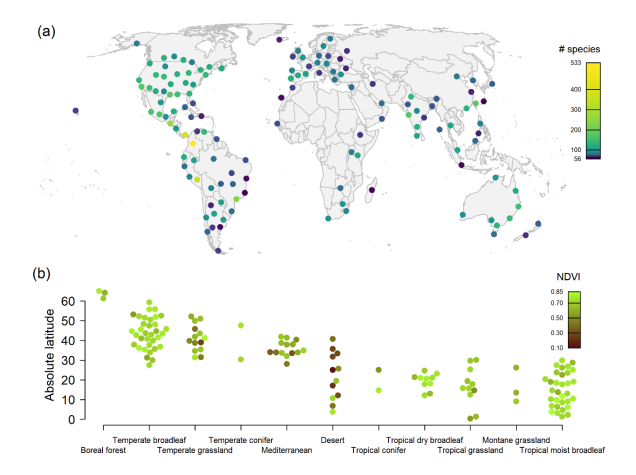
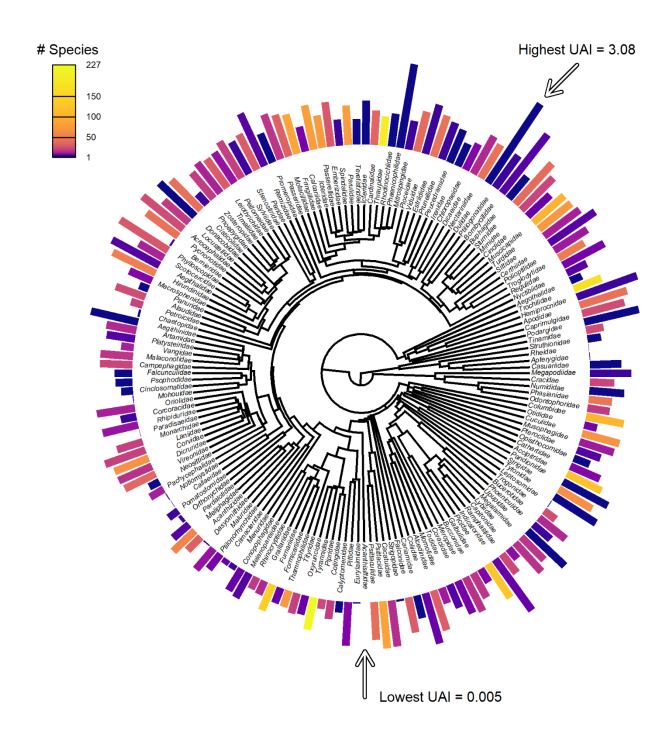
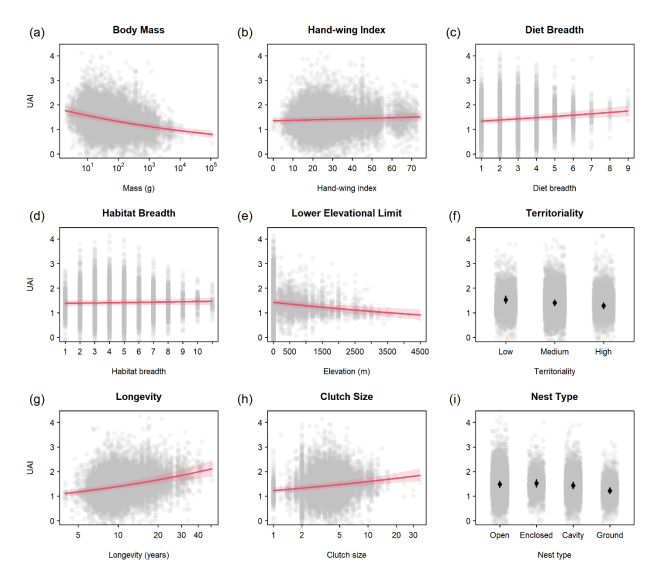


Figure 1. Geographical coverage of 137 cities included in the analysis. Cities were distributed (a) across 62 countries, including 39 in North America, 28 in South America, 27 in Asia, 22 in Europe, 10 in Africa, and 10 in Australasia. Each of these cities was initially selected to have a population of  $\geq$ 100,000 people and be  $\geq$ 500 km apart. Cities were then retained that had  $\geq$ 50 species each with  $\geq$ 100 eBird records within a 100 km radius circles over 20 years (2002–2021). Cities are colored by the (log) number of species that met the criteria from 56 (dark purple) to 533 (yellow). Cities were representative (b) of 11 of the world's biomes (Olson & Dinerstein 1998). Biomes are ordered by the mean absolute latitude of the cities included, and cities are colored by the NDVI of the greenest month, from the greenest city (lime green) to the least green city (dark brown).



# **Figure 2.** The phylogeny of Urban Association Index (UAI) across 3768 species by family. For visualization, UAI values were averaged across species and then across taxonomic families. The height of the bar indicates the UAI, with taller bars indicating higher urban tolerance. The color indicates the (log) number of species in the family from 1 (dark purple) to 231 (Tyrannidae, yellow).

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**Figure 3.** The global mean effects of nine species traits on the Urban Association Index (UAI) of 3768 bird species across 137 cities. There was a significant relationship between UAI and (a) body mass, (b) hand-wing index, (c) diet breadth, (d) habitat breadth, (e) lower elevational limit, (f) territoriality, (g) longevity, (h) clutch size, and (i) nest type. Gray points show the partial residuals of each data point. Trend lines for numerical traits are shown along with the 95% credible intervals. Territoriality is treated as numerical in the model but here we summarize the data for the three levels of territoriality (low, medium, high). Nest type is treated as categorical in the model (open, enclosed, cavity, or ground). For territoriality and nest type, black points show the mean and bars show the 95% credible intervals.

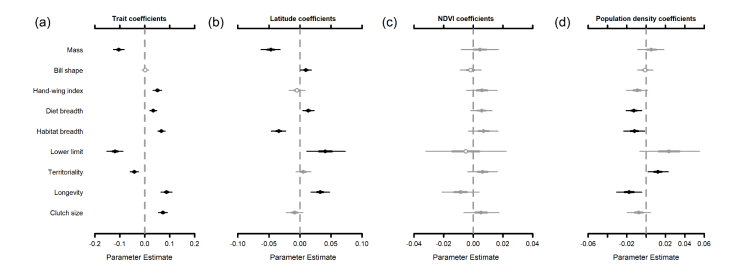


Figure 4. The relationship between Urban Association Index (UAI), species traits, and city variables for 3768 birds species across 137 cities. Covariate coefficients ( $\beta$ ) show how UAI varies as a function of nine numerical trait covariates (a). In turn, these trait coefficients vary across cities as a function of three citylevel variables with corresponding coefficients ( $\theta$ ): (b) latitude, (c) NDVI, and (d) human population density. Points show the covariate coefficient estimates with corresponding interquartile range (thick lines) and 95% credible intervals (thin lines). Points are open when the interquartile range overlaps 0. Points and lines are gray when the 95% credible intervals overlap 0 and black when they do not.

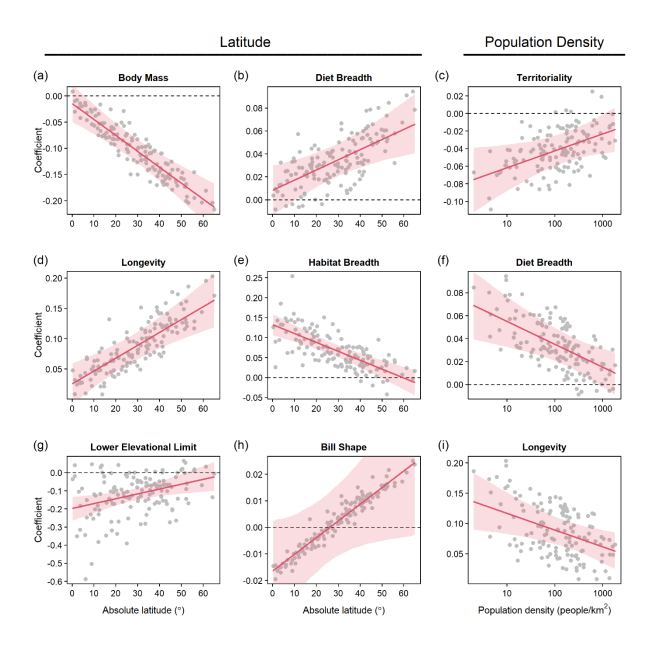


Figure 5. The effect of latitude and human population density on the city-level trait coefficients for Urban Association Index. Latitude had a significant effect on the city-level trait coefficients for (a) body mass, (b) diet breadth, (d) longevity, (e) habitat breadth, (g) lower elevational limit, and (h) bill shape. Human population density had a significant effect on the city-level trait coefficients for (c) territoriality, (f) diet breadth, and (i) longevity. Points represent the model-estimated trait coefficients for each city (n = 137). Trend lines and 95% credible intervals show how these coefficients vary as a function of the citylevel covariates.

# Supporting Information

We tested for the effects of the difference in spatial scale between VIIRS and eBird checklists. The VIIRS night-time lights imagery has a spatial resolution of ~500 m, much smaller than the 5 km filter applied to eBird checklists, and so a single point value may not be representative of the landscape sampled during a specific checklist. In order to check whether this affected our index, we experimented on six example cities, one from each continent (Los Angeles, Buenos Aires, London, Nairobi, Mumbai, Sydney). For every checklist locality within each city, we sampled 100 points from a bivariate normal distribution of latitude and longitude centered on the checklist locality, with a standard deviation of 1 km, truncating values >5 km from the locality. This sampling approach created a scatter of points around the locality, from which a mean radiance value can be calculated. From these mean estimates, we then calculated the mean radiance values for each species across localities. We found that these species-level estimates based on sampled points were highly correlated to the estimates based on single radiance values per locality (r = 0.97–0.98; Fig. S3). Thus, our Urban Association Index (UAI) for each species is the mean radiance value across records where the radiance value of each record is taken from a single pixel of radiance.

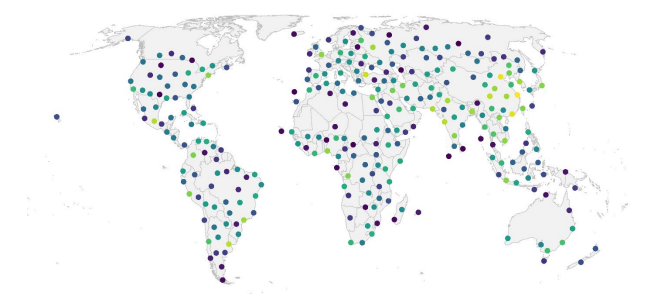
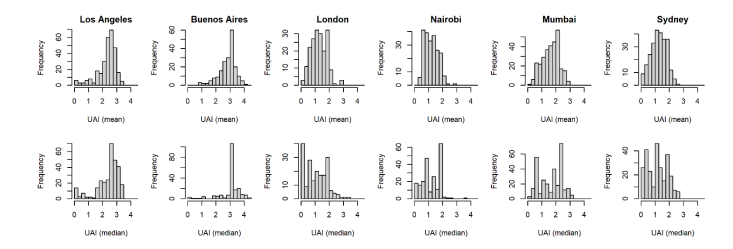
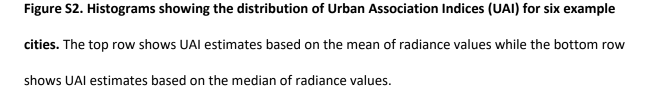
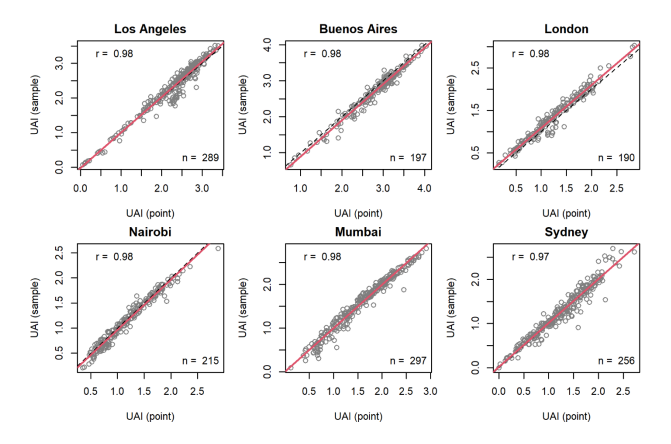


Figure S1. A map of 289 cities spaced ≥500 km apart with populations ≥100,000. Points are colored



from least (purple) to most (yellow) populous.





**Figure S3. A comparison of two Urban Association Indices (UAI) for six example cities.** UAI (point) estimates are based on radiance values from a single pixel of night time lights for each locality. UAI (sample) estimates are based on the mean of 100 points sampled randomly from around each locality. Each point represents the UAI of a species. The dashed line shows 1:1 correspondence, the red line shows the trend between the two indices, and the Pearson's correlation coefficient is given in the top-right corner.

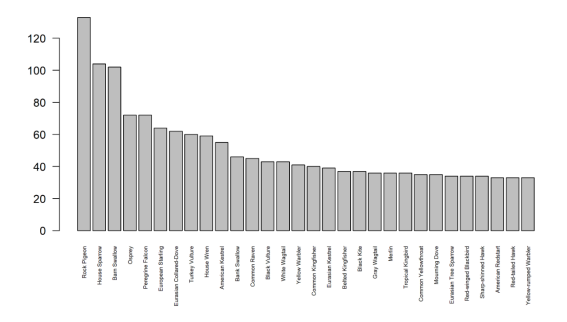


Figure S4. The 50 species represented in the most city circles.

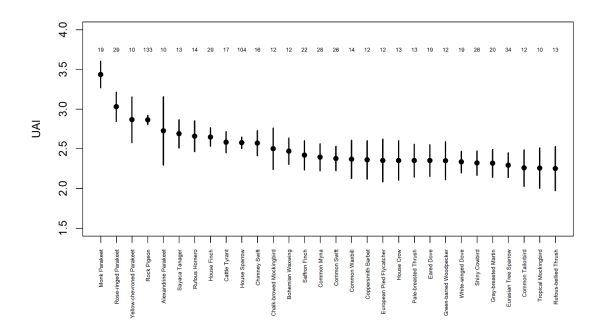


Figure S5. The 30 species with the highest Urban Association Index (UAI). For each species the mean and standard error of their UAI is shown, along with the sample size (i.e., number of cities). Only species present across  $\geq$ 10 city circles are shown.

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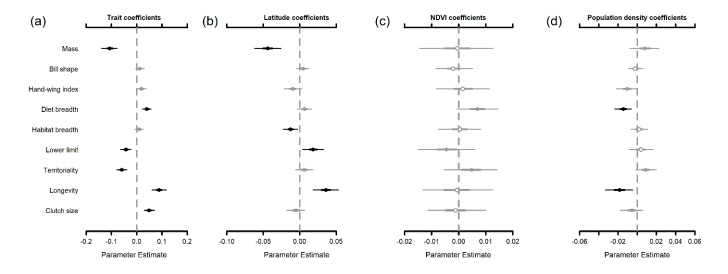


Figure S6. The relationship between Urban Association Index (UAI), species traits, and city covariates for the 920 birds species each represented in at least five cities. This figure is comparable with the results for the full species set in Fig. 4. Covariate coefficients ( $\beta$ ) show how UAI varies as a function of nine numerical trait covariates (a). In turn, these trait coefficients vary across cities as a function of three city-level variables with corresponding coefficients ( $\theta$ ): (b) latitude, (c) NDVI, and (d) human population density. Points show the covariate coefficient estimates with corresponding interquartile range (thick lines) and 95% credible intervals (thin lines). Points are open when the interquartile range overlaps 0. Points and lines are gray when the 95% credible intervals overlap 0 and black when they do not.