Increased tolerance to humans among disturbed wildlife

Diogo S.M. Samia¹, Shinichi Nakagawa²,³, Fausto Nomura¹, Thiago F. Rangel¹ & Daniel T. Blumstein⁴

Human disturbance drives the decline of many species, both directly and indirectly. Nonetheless, some species do particularly well around humans. One mechanism that may explain coexistence is the degree to which a species tolerates human disturbance. Here we provide a comprehensive meta-analysis of birds, mammals and lizards to investigate species tolerance of human disturbance and explore the drivers of this tolerance in birds. We find that, overall, disturbed populations of the three major taxa are more tolerant of human disturbance than less disturbed populations. The best predictors of the direction and magnitude of bird tolerance of human disturbance are the type of disturbed area (urbanized birds are more tolerant than rural or suburban populations) and body mass (large birds are more tolerant than small birds). By identifying specific features associated with tolerance, these results guide evidence-based conservation strategies to predict and manage the impacts of increasing human disturbance on birds.
Animals often perceive humans as predators\(^1\). Thus, even non-consumptive human activities may affect their behaviour\(^2,3\) and generate population-level consequences\(^4,5\). For instance, declining populations of European and Australian birds are less tolerant of humans, as evidenced by them fleeing at greater distances from an approaching human (that is, flight initiation distance—FID), whereas populations from more tolerant species are increasing\(^6,7\). In addition, a synergistic interaction between human tolerance and increased urbanization is expected\(^8,9\), resulting in an increased negative effect for less tolerant species. Hence, there is an urgent need to understand the factors that explain species tolerance of humans and to develop predictive models of tolerance\(^10,11\). One strategy to do so is to identify traits associated with species tolerance (that is, reduced responsiveness to people in areas with more people) or intolerance (that is, increased responsiveness to people in areas with more people) to human presence, which can be done by capitalizing on studies that compare the responses of animals in areas with varying levels of human presence.

We conducted a meta-analysis of 75 studies on birds (180 species), mammals (16 species) and lizards (16 species) to comprehensively evaluate species' tolerance of human disturbance. To address the potential non-independence among effect sizes, we incorporated study identity and species phylogeny as random factors. For each observation, we calculated the Hedges' \(g\)\(^11\), a measure of effect size, by comparing the mean FID of a population observed in areas of higher human disturbance with the mean FID of a population observed in areas of lower human disturbance. Negative effect sizes indicate tolerance of human disturbance, whereas positive effect sizes indicate intolerance of human disturbance. For the best-studied taxa—birds—we used a multi-model inference approach\(^12\) to identify the most important predictors of tolerance of human disturbance. Specifically, we tested for the effects of body mass, clutch size, habitat openness, group size, foraging habit, diet, migration behaviour and the type of habitats contrasted in populations under low and high human disturbance (for example, rural versus urban, natural versus urban; hereafter referred to as 'habitat contrast') on bird's tolerance of human disturbance. Because it is well known that urbanization is a correlate of avian extinction\(^13\), we conducted a separate meta-analysis focusing on bird populations from the rural–urban habitat contrast (the best studied contrast) to investigate if traits were similarly likely to explain differential behavioural responses\(^13,14\).

Here we find that, overall, disturbed populations of animals of the three major taxa are more tolerant of human disturbance than undisturbed (or less disturbed) populations. We also find that habitat contrast, body mass, clutch size and diet are the best predictors of direction and magnitude of tolerance of human disturbance. When we focused only on bird populations from the rural–urban habitat contrast, however, we find that body mass, diet and habitat openness are the best predictors of direction and magnitude of tolerance of these populations. Our results help to identify species that are potentially more vulnerable to increasing human disturbance and provide evidence-based guidance to future conservation actions.

Results

Overall results. In general, lizards, mammals and birds became equally more tolerant of human approaches with increasing human presence (\(Q_h = 0.09\), \(df = 2\), \(P = 0.96\); Fig. 1 and Supplementary Data 1). Importantly, a regression between the mean and standard deviation of raw FID values showed that our effect size estimates genuinely reflect the magnitude of the mean FID differences of populations in areas of low versus high human disturbance (that is, they were not biased by any potential difference in variance of FIDs as a function of human disturbance level; Supplementary Fig. 1). Using \(I^2\), an index of heterogeneity among effect sizes\(^15\), we found substantial variation both in the meta-analysis of the three major taxa (\(I^2_{\text{total}} = 91.44\%\), \(I^2_{\text{between-study}} = 46.09\%\), \(I^2_{\text{phylogeny}} = 10.15\%\), \(I^2_{\text{within-study(residuals)}} = 35.20\%\)) and in the birds-only meta-analysis (\(I^2_{\text{total}} = 89.51\%\), \(I^2_{\text{study}} = 36.87\%\), \(I^2_{\text{phylogeny}} = 13.90\%\), \(I^2_{\text{residual}} = 38.74\%\)). This amount of heterogeneity justified our further exploration of the covariates. There was little evidence that our conclusions were affected by publication bias, where studies with low sample sizes are more prone to rejection because their higher probability of finding non-significant effects\(^16\) (Egger's regression; all taxa: intercept = -0.22, \(P = 0.243\); birds-only: intercept = -0.17, \(P = 0.397\); Supplementary Fig. 2).

Using \(H^2\), an index for phylogenetic signal\(^17\), we found relatively small phylogenetic signal in the species tolerance of human disturbance (all taxa: \(H^2 = 11.10\%\); birds-only: \(H^2 = 15.53\%\)). In fact, a separate set of analyses with meta-analytic models using studies and species as random factors, but not controlling for the phylogeny, yielded the same conclusions (Supplementary Figs 3–6, Supplementary Tables 1 and 2 and Supplementary Data 2 and 3).

Model selection using all bird species. The best predictors of bird response to human disturbance were habitat contrast, body mass, clutch size and diet (Table 1 and Supplementary Data 4). Magnitude and direction of birds responses varied according to the habitat contrast (\(Q_h = 20.34\), \(df = 8\), \(P = 0.009\); Fig. 2). Birds in urbanized environments (that is, rural versus urban populations, and suburban versus urban populations) showed the greatest tolerance of humans (Table 1 and Fig. 2). Overall, the magnitude of the FID difference (that is, the effect sizes) of populations under low versus high human disturbance was lower, albeit marginally significant, in contrasts made ‘within’ a same habitat type (for example, low versus high human disturbance in islands) than in contrasts made ‘between’ habitat types (for example, rural versus urban areas; \(Q_h = 3.70\), \(df = 1\), \(P = 0.054\)). Larger birds were more tolerant of human disturbance than smaller birds (Table 1 and Fig. 2). Species that produced more eggs were less tolerant of human disturbance than those that produced fewer eggs (Table 1 and Fig. 2). Herbivorous and
Table 1 | Summary of the multi-model inference conducted to explain avian responses to human disturbance.

| Predictor | Levels                              | Estimate | s.e.  | Importance |
|-----------|-------------------------------------|----------|-------|-----------|
| All birds (180 species, 457 effect sizes) | Intercept | 0.736   | 0.599 |           |
| Habitat contrast | Natural versus urban | −0.941 | 0.632 | 1         |
| Rural versus suburban | −1.438 | 0.762 | —    |           |
| Rural versus urban | −1.689 | 0.589 | —    |           |
| Suburban versus urban | −1.690 | 0.740 | —    |           |
| Low versus high in urbanized areas | −1.274 | 0.635 | —    |           |
| Low versus high in recreational nature | −0.684 | 0.593 | —    |           |
| Low versus high in islands | −0.894 | 0.717 | —    |           |
| Low versus high in reserve | −1.622 | 0.723 | —    |           |
| Body mass | −0.315 | 0.075 | 0.98      |
| Clutch size | 0.052 | 0.021 | 0.93      |
| Diet | Herbivorous | −0.232 | 0.131 | 0.91      |
| Omnivorous | −0.260 | 0.105 | —    |           |
| Habitat openness | 0.182 | 0.091 | 0.83      |
| Migration | 0.127 | 0.080 | 0.70      |
| Group size | 5–50 Individuals | −0.060 | 0.088 | 0.51      |
| >100 Individuals | 0.097 | 0.138 | —    |           |
| Foraging habit | −0.007 | 0.109 | 0.41      |

Rural versus urban birds (103 species, 324 effect sizes)

| Predictor | Levels                              | Estimate | s.e.  | Importance |
|-----------|-------------------------------------|----------|-------|-----------|
| Intercept | −0.662 | 0.309 |           |
| Body mass | −0.403 | 0.088 | 0.99      |
| Diet | Herbivorous | −0.376 | 0.164 | 0.88      |
| Omnivorous | −0.244 | 0.130 | —    |           |
| Habitat openness | 0.215 | 0.105 | 0.85      |
| Migration | 0.035 | 0.025 | 0.64      |
| Group size | 0.123 | 0.095 | 0.62      |
| 5–50 Individuals | −0.100 | 0.108 | 0.56      |
| >100 Individuals | 0.091 | 0.176 | —    |           |
| Foraging habit | 0.079 | 0.130 | 0.46      |

Results are shown from both a meta-analysis using the full data set (all birds) and a meta-analysis focusing on the contrast between rural and urban populations. Values are average coefficients of models estimated, their associated standard error (s.e.), and the importance of each factor in explaining species responses to human disturbance (the closer than 1, the more important the factor). Habitat contrasts presented as 'low versus high in' refer to contrast between populations experiencing low and high human disturbance within a given habitat type.

Omnivorous species were more tolerant than carnivorous species (Table 1 and Fig. 2).

Model selection using birds from the rural–urban contrast. When we focused only on bird populations from the rural–urban habitat contrast, we found a slightly different rank order of importance of the best predictors (Table 1 and Supplementary Data 5). Once again, body mass was the most important species trait that explained variation in tolerance of human disturbance. However, both the larger regression coefficient and greater importance in relation to the other traits suggest that the effect of body mass on tolerance was even greater in the rural–urban habitat contrast (Fig. 3 and Table 1). Diet was the second most important trait explaining species tolerance in the rural–urban habitat contrast (Fig. 3 and Table 1). Habitat openness was more important trait explaining species tolerance in the rural–urban habitat contrast (Fig. 3 and Table 1). Diet was the second most important trait explaining species tolerance in the rural–urban habitat contrast (Fig. 3 and Table 1). Habitat openness was more important trait explaining species tolerance in the rural–urban habitat contrast (Fig. 3 and Table 1). Diet was the second most important trait explaining species tolerance in the rural–urban habitat contrast (Fig. 3 and Table 1).

Discussion

We found the habitat type contrasted in populations under low and high human disturbance (that is, the habitat contrasts) was one of the main drivers of the degree to which birds tolerate human disturbance. Rural versus urban and suburban versus urban populations had the largest effect sizes among the habitat contrasts, indicating that urbanized birds are those with the highest degree of tolerance of human disturbance. A previous study showed that time since urbanization was positively associated with the degree of tolerance in birds, which could help to explain further variation in human tolerance within urban places. However, the lack of additional data prevents further tests of the effect of time since urbanization as a predictor of tolerance of human disturbance. Although sample sizes were admittedly smaller for the other habitat contrasts, the effect sizes of all contrasts (except inside versus outside reserves) were in the same direction, suggesting that different types of human exposure act generally increasing tolerance of populations. It is essential to realize that some species might be absent from certain habitat comparison because they are unable to survive in habitats with high human presence (for example, in urban areas) because of their low tolerance for human activity. Therefore, a more detailed examination of variation in human tolerance among habitats types would require more sampling in most habitats, a historical record of human or bird invasions, and a systematic
measurement of human presence (as conducted by few studies of our data set\textsuperscript{19,20}). Nevertheless, our finding that FID differences were lower in contrasts within habitat types than between habitat types seems to support previous studies that showed a positive relationship between human exposure and degree of human tolerance\textsuperscript{20–22}.

Based on a robust body of evidence\textsuperscript{23–27}, we expected body mass to be one of the most important traits affecting how species respond to increased human disturbance. However, we did not initially expect the observed direction of the response (see Covariates in Methods). It is well established that large animals are more intolerant to human presence than small animals (that is, they have larger FID\textsuperscript{23–27}), but, surprisingly, we found that large birds were those that had the greatest reduction in FID as human disturbance increased. Optimal escape theory states that animals must counterbalance both costs and benefits when making escape decisions\textsuperscript{27,28}. Larger animals may suffer higher costs of flight either because they are less agile because of their size\textsuperscript{29,30} or because their energetic costs of not foraging are greater (opportunity cost\textsuperscript{26,31,32}). Either way, we suggest that larger birds (and potentially, larger species of other taxa too) might be under intense pressure to be more tolerant of non-lethal human approaches. This finding may indicate that energetic costs of unnecessary escape are important variables driving tolerance.

There are at least two other reasons why body size may be associated with increased tolerance of humans in birds. First, larger birds may also be less likely to be killed by predators because of their body size\textsuperscript{33} and this reduced risk may select for increased tolerance of non-threatening humans in areas where humans are commonly encountered. Second, larger birds with relatively larger brains may have greater cognitive abilities and might be able to better assess risk\textsuperscript{34}.

At a proximate level, our finding that carnivorous birds are less tolerant of humans may reflect a carry-over effect from their increased sensitivity to movement and thus their general degree of responsiveness\textsuperscript{23}, or it may reflect the reduced foraging efficiency of carnivorous birds foraging around humans if their prey are similarly disturbed by humans. Carnivorous birds may also be persecuted because they compete with humans for prey, and, if so, may have been selected to be less tolerant of humans. Regardless of the mechanism, the evidence accumulated to date, and summarized in our meta-analysis, suggests that carnivorous birds are likely to be particularly vulnerable to human disturbance.

Animals in open habitats should be better able to monitor predator behaviour once they are detected, and to track risk dynamically\textsuperscript{34}. Based on the ability to better track predation risk, we expect that animals in open habitats to be better at learning not fear encounters with non-threatening humans than animals living in visually closed environments where sightings are either brief or interrupted\textsuperscript{34}. Human-altered habitats, such as urban places, are often characterized by having reduced vegetative cover. Species naturally living in open habitats might suffer less in urbanized habitats once vegetation cover is reduced in cities, and consequently they might better tolerate humans. This may explain why habitat use (specifically, regarding to habitat openness) was a trait particularly important in explaining variation in tolerance in urbanized birds. As an extension of

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**Figure 2 | Effects of eight predictors on the effect sizes (Hedges’ g) of bird’s tolerance of human disturbance using the full data set (180 species, 457 effect sizes).** Negative effect sizes show that species tolerate human disturbance, whereas positive effect sizes show that species are intolerant of human disturbance. Predictors are presented in their order of importance in explaining bird’s tolerance (or intolerance) to human disturbance. (a) Type of habitats contrasted (contrasts are presented as habitat with ‘low versus high’ human disturbance). Habitat contrasts presented as ‘low versus high in’ refer to contrast between populations experiencing low and high human disturbance within a given habitat type. (b) Body mass (g). (c) Mean clutch size per reproductive period. (d) Diet. (e) Habitat openness. (f) Migration. (g) Group size. (h) Foraging habit. Horizontal dashed line indicates zero effect size. Different sizes of symbols in b and c reflect differences in sample size. The error bars illustrate 95% confidence intervals. The number of effect sizes in each categorical level is shown in the bottom of figures.
Our meta-analysis shows that bird species that produce smaller clutches are more tolerant of human disturbance than those that produce larger clutches. Life history theory predicts that species that invest more in their offspring should be more likely to protect that investment\(^{36,37}\) (but see ref. 38). If we assume that small clutch-sized species generally have larger parental investment per offspring, then adults should accept more risk to guarantee their reproductive success. Abandoning a profitable patch, in addition to the energetic cost of flight, should thus be particularly costly to these species. Therefore, there should be a pressure for small clutch-sized birds to become more tolerant of non-threatening encounters. This hypothesis is consistent with the energetic hypothesis proposed by optimal escape theory\(^{27,28}\).

At least three mechanisms can explain variation in tolerance as a function of human disturbance. First, recent studies have suggested that differential selection among personality types could account for increased tolerance of humans in some species, whereby bold individuals settle in areas with higher human disturbance and shy individuals settle in areas with reduced disturbance\(^{4,39}\). Second, increased tolerance could result from local adaptation\(^{40}\). Third, increased tolerance could result from habituation\(^{41,42}\). We suggest that habituation-like processes are likely to be common and important because we know that individuals are exquisitely sensitive to unexpected behaviour from humans and potential predators\(^{27}\). Indeed, we expect location-specific habituation where even a slight deviation in a predator’s (or human’s) routine behaviour can re-elicit fearful responses\(^{43}\) (for example, when predator appears on a different side of lake\(^{44}\) or a human steps off a well-travelled hiking trail or crosses a fence\(^{45}\)). Such a pattern is entirely consistent with habituation.

Discriminating between the three competing mechanistic hypotheses to explain tolerance of human disturbance would require a set of long-term studies on marked individuals (for example, because habituation is a learning process that occurs in individuals over time\(^{42}\), in addition to population-level genetic information to investigate selection and adaptation. These data are currently unavailable for the vast majority of the species studied. However, our results show that regardless of the mechanism leading to increasing tolerance of human disturbance, its magnitude and direction, at least in birds, is influenced by a species’ traits and environmental differences. The present study thus helps address an urgent need to identify species potentially more vulnerable to the threats of human expansion\(^{10,46}\).

The results of our systematic review and meta-analysis have implications for wildlife conservation. For example, our findings suggest that small birds may benefit more from protection that reduces human disturbance than large birds, specifically because they are less likely to become tolerant of human disturbance. In practical terms, it is possible that set-back distance zones established within protected areas\(^{10}\) are more effective for small bird species. In addition, and contra previous suggestions based only on the relationship between body size and FID\(^{23}\), our findings suggest that ecotourism may not be as deleterious for larger birds (and perhaps larger animals) because they are better able to tolerate human disturbance and consequently may suffer smaller population-level consequences\(^{2,4,5}\).

Despite the conservation implications discussed above, at least two points related to our results require caution. First, large birds might become more vulnerable to illegal hunting if they become tolerant of humans as a function of exposure\(^{47}\). Thus, exposure to benign human presence (for example, ecotourists) could create an ecological trap if tolerant birds later encounter hunters. Therefore, wildlife managers interested in using our results to design a management programme should also consider actions aimed to curb poaching.

Figure 3 | Effects of seven predictors on the effect sizes (Hedges’ g) of bird’s tolerance of human disturbance from populations of rural-urban habitat contrast (103 species, 324 effect sizes). Negative effect sizes show that species tolerate human disturbance, whereas positive effect sizes show that species are intolerant of human disturbance. Predictors are presented in their order of importance in explaining bird’s response to human disturbance. (a) Body mass (g). (b) Diet. (c) Habitat openness. (d) Mean clutch size per reproductive period. (e) Migration. (f) Group size. (g) Foraging habit. Horizontal dashed line indicates zero effect size. Different sizes of symbols in a and d reflect differences in sample size. The error bars illustrate 95% confidence intervals. The number of effect sizes in each categorical level is shown in the bottom of figures.
Second, the bird traits correlated with greater tolerance of human disturbance are the same found among many endangered species (that is, endangered species are typically large body sized and have low reproductive rates46,48,49). This apparently contradictory finding can be explained by considering the source of human disturbance. Large animals with low reproductive rates are in decline mainly because of habitat loss, overexploitation by humans and by climate change58,59. Large animals require larger home ranges, have reduced recruitment and are often hunted by humans for food and recreation46,48,49. Our results focus on behavioural responses to non-lethal human disturbance (that is, tolerance), which may influence a species' ability to live successfully with humans24–7,42. Therefore, although our findings may be used to predict bird species potentially most vulnerable to human disturbance, and to manage the impacts of increasing human presence on wildlife, other conservation actions must be adopted to decelerate the alarmingly rapid loss of biodiversity51.

The present study represents the first effort to develop an evidence-based model of wildlife tolerance of human disturbance. We hope the results of our meta-analysis serve as a starting point to predict and manage the impacts of increasing human disturbance on wildlife and guide conservation strategies. Additional studies must be conducted to understand the mechanisms that promote tolerance of human disturbance, because this mechanistic understanding should help us design more effective management strategies10,52. Also, studies are necessary to determine visitation rates that maximise the development of tolerance of humans in wildlife. Ultimately, it is evidence-based actions that will help reduce biodiversity loss in a rapidly urbanizing world.

Methods

Literature search and selection criteria. We first compiled all studies that cited or were cited by two key reviews28,31. We searched for studies published before 31 July 2014 on Web of Knowledge, Scopus and Google Scholar databases with the key words ‘flight initiation distance’, ‘flight distance’, ‘escape distance’, ‘approach distance’, ‘flushing distance’ and ‘response distance’. The references cited in these studies were also examined. Our criteria to include a study were that FID of a given species (measured sensor27) had to be collected in areas of low and high human presence (= human disturbance). We followed the criteria used by authors to categorize each area as a function of its degree of disturbance. Hunted populations were not included in our data set. A PRISMA diagram describing our literature search is available in Supplementary Fig. 8 and Supplementary Table 3, respectively. Our final data set consisted of 504 effect sizes from 75 studies across 212 species distributed in three major taxa: birds, mammals and lizards (data available in Supplementary Data 1).

Estimating effect sizes. To quantify species response to disturbance, we used the effect size metric Hedges’ g11; a bias-corrected measure of standardized mean differences, which does not overestimate the magnitude of effect when sample size is small51. For each species, we compared the mean FID of populations in areas of higher human disturbance with those in areas of lower human disturbance. These FID comparisons were restricted to populations of species studied in a same study. In our data set (Supplementary Data 1), positive effect sizes indicate sensitization, whereas negative values indicate tolerance of human disturbance. When mean, variance and sample size of FIDs were not provided in a paper, we estimated Hedges’g from the statistical results (t, F, χ2, Z and P)52. We directly contacted several authors for missing data (see Acknowledgements for details). Importantly, recent studies have shown that the starting distance of an approaching person (that is, animal–human distance when the approach begins) can affect FID7,52. However, the potential effects of starting distance on FID were controlled in all effect sizes either because studies used a fixed starting distance among experiments (for example, starting distance fixed in 30 m) or because starting distance was controlled analytically by using it as a covariate in statistical models, or because we used the marginal means for those studies that provided them or those studies in which we were able to obtain the raw data (see Supplementary Data 1 for details).

Meta-analysis. We used multi-level mixed-effects meta-analysis37 to test for both mean effect sizes and the importance of our predictors. We controlled for phylogenetic and study non-independence by including phylogeny (Supplementary Fig. 9) and study identity as random-factors in our models17,56. Although we also have multiple estimates per species in our data set (Supplementary Data 1), a model selection approach showed that the inclusion of ‘species identity’ as an additional random-effect did not improve our models (Supplementary Table 4). Phylogenies of birds, mammals and lizards and how they were combined to test for difference among these taxa is described in Supplementary Methods 1. The mean effect sizes (that is, mean of the effect sizes weighted by the inverse of their variance) were considered significant if their 95% confidence intervals did not include zero53. We used the bootstrap zero-group heterogeneity statistic (Q0) to test for significant difference between mean effect sizes54,55. We used I2 as a measure of heterogeneity in the effect sizes in which the value represents the proportion of total variation in data that is not sampling error; 100% sampling error; 100% sampling variation (that is, Q0 = 0). We calculated the degree of phylogenetic signal in our effect size estimates using the phylogenetic heritability index15, H2, which is the variance attributable to phylogeny in relation to the total variance expected in the data. When the unit of analysis is species, H2 is equivalent to Pagel’s λ (ref. 55), in which higher values are associated with stronger phylogenetic signals. Primary studies can suffer from publication bias, where studies with low sample size are more prone to be rejected because of their higher probability of not finding significant effect sizes66,63. We checked for publication bias using Egger’s regression66 in which intercepts significantly different from zero suggest potential publication bias. To overcome the non-independent nature of our data, we applied the Egger’s regression test on the meta-analytic residuals67. Analyses were conducted using the metafor54 R package v1.9–4.

Covariates. The large number of observations in birds permitted us to investigate whether certain variables were potentially important predictors of tolerance of human disturbance. Based on previous findings in the literature and our own hypotheses, we collected information on eight variables. Seven were associated with a species’ morphology, life-history and natural history traits. These data were obtained from Del Royo et al.36 (see Supplementary Data 1 for details). The eighth variable, the habitat contrast, describe the type of habitats contrasted in populations under low and high human disturbance. These data were extracted from each surveyed paper. Importantly, for these covariates, multi-collinearity was not an issue (variance inflation factor < 1.50, below the suggested threshold of 3; see also correlation matrix in Supplementary Table 5). Below we justify the use of each variable and our predictions concerning their effect on bird response.

Body mass. A substantial amount of empirical evidence shows that large animals are less tolerant of human approaches—they generally flush at a greater distance from humans—7–27. There are two non-exclusive hypotheses to explain this response: (i) large animals are generally less maneuverable29,30 or (ii) large animals suffer a greater opportunity cost of not foraging because of their greater absolute metabolic needs7,25,32. Thus, we hypothesized that large animals would tolerate less human disturbance by showing larger positive effect sizes than small animals. Body mass was measured in grams but transformed into log10.

Group size. Three models of predation risk assessment predict a declining risk of predation as group size increases58–60. If predation risk decreases with group size, individuals in larger groups might thus tolerate closer approach. Moreover, if tolerance of non-lethal human disturbance is a socially transmitted behaviour, one could test the effect of social transmission by examining group size61. We used a mixed-effects model as a measure of heterogeneity in the effect sizes in which the value represents the proportion of total variation in data that is not sampling error; 100% sampling variation (that is, Q0 = 0). We calculated the degree of phylogenetic signal in our effect size estimates using the phylogenetic heritability index15, H2, which is the variance attributable to phylogeny in relation to the total variance expected in the data. When the unit of analysis is species, H2 is equivalent to Pagel’s λ (ref. 55), in which higher values are associated with stronger phylogenetic signals. Primary studies can suffer from publication bias, where studies with low sample size are more prone to be rejected because of their higher probability of not finding significant effect sizes66,63. We checked for publication bias using Egger’s regression66 in which intercepts significantly different from zero suggest potential publication bias. To overcome the non-independent nature of our data, we applied the Egger’s regression test on the meta-analytic residuals67. Analyses were conducted using the metafor54 R package v1.9–4.

Habitat openness. Animals that live in habitats can simultaneously detect predators at a greater distance and might have to travel a longer distance to reach protective cover. A recent study showed that birds originally from open habitats tend to delay the flight when compared with birds from closed habitats24. Moreover, highly altered habitats, such as urban and suburban places, usually have reduced vegetation cover. We thus hypothesized that species living in large open habitats would better tolerate humans because they would suffer less as vegetation cover was reduced. Either way, habitat openness might influence FID and must be accounted for if we are to isolate human disturbance effects. We categorized species as being originally from open habitats (for example, uplands and grassland) or closed habitats (for example, dense forests and woodlands).

Diet. Birds that eat live prey, particularly carnivorous raptors, have especially good visual acuity and motion sensitivity66. Given this sensitivity, birds that eat live...
prey are disturbed at greater distances. This increased sensitivity to disturbance could select for tolerance if they are better to learn that humans are not a threat. Alternatively, it could also reflect the possibility that foraging efficiency is reduced because their prey are similarly disturbed by humans. In the latter case, we might expect that these species would be less likely to tolerate the on-going disturbance. We categorized species as carnivorous, herbivorous, or omnivorous.

Migration. Birds that migrate are exposed to a greater variety of habitats and might be selected to rapidly learn to assess predation risk in new habitats. Under this scenario, we expected that these species would be more likely to tolerate increased disturbance. We ceded species as resident or migratory.

Clutch size. Like body size, clutch size is a life history trait that reflects energetic investment, and therefore need. In general, we might expect that birds that produce fewer eggs per reproductive period might be more energetically stressed than those that produce more eggs because they have larger parental investment per offspring. In this case, small-clutch-sized species would tolerate closer human presence and thus a differential human tolerance is expected as a function of frequency of exposure to humans. Studies have shown that even subtle temporal or spatial change in human disturbance within a given habitat type may trigger changes in animal’s tolerance of humans. Justifying our exploration of the contrasts between low versus high human disturbance in islands and (ix) low versus high human disturbance in urbanized areas, (vii) low versus high human disturbance in suburban areas, (vi) rural versus urban area, and (v) inside versus outside reserve, (v) low versus high human disturbance in urbanized areas, (vii) low versus high human disturbance in recreational nature (for example, beaches, ski areas and other tourist locations), (viii) low versus high human disturbance in islands and (ix) low versus high human disturbance in reserve. We used these habitat contrasts either because of their difference in human disturbance degree or because a particular characteristic of habitat is expected to influence animal’s tolerance of humans. Specifically, contrasts between natural, rural, suburban and urban populations were tested because they represented increasing human presence and thus a differential human tolerance is expected as a function of frequency of exposure to humans. Studies have shown that even subtle temporal or spatial change in human disturbance within a given habitat type may trigger changes in animal’s tolerance of humans. Justifying our exploration of the contrasts between low versus high human disturbance within a same habitat type (levels vi–ix). Overall, we expect a lower FID difference that is, tolerance in comparisons within than between habitat types. Recreational areas, islands and protected areas (reserves) were tested separately because their marked difference in the pattern of human disturbance. Because island populations often have reduced predation risk compared with mainland populations, we expected either none or a small difference in FIDs of different populations found on islands. Animals living in natural areas with tourism may be more responsive to humans because they commonly experience seasonality in human disturbance (for example, visitation only in summer or winter). Beyond temporal variation, populations inside and outside protected areas may suffer marked spatial variation in human disturbance, as well as potential lethality associated with human presence (for example, individuals in protected areas might be safer). Human spatial variation in human disturbance may also occur in populations ‘within’ protected areas if the frequency of human visitation varies in the protected area. Importantly, our habitat contrasts were restricted to comparisons between populations of species tested in the same study. As explained in the Meta-analysis section, the variation among studies was controlled for by using study identity as a random factor in our models.

Multi-model inference. We used a multi-model inference approach based on Akaike’s criteria corrected for small sample size to estimate the relative importance of the predictor variables. To calculate the importance of each predictor, we first assessed the relative strengths of each candidate model by calculating its Akaike weight; analogous to the probability of that model is the best model. A constant importance of the predictor variables. To calculate the importance of each predictor, we first estimated the importance of each predictor by summing the Akaike weights of all models in which that candidate variable appeared, which can be interpreted as the probability that a particular predictor is a component of the best model, which allowed us to rank predictors in order of importance. We used a model averaging approach to estimate model parameters. Multi-model analyses were conducted using the MuMIn R package v. 1.14.0.

References
1. Frid, A. & Dill, L. Human-caused disturbance stimulus as a form of predation risk. Conserv. Ecol. 6, 1–11 (2002).
2. Preiser, E. L., Bolnick, D. I. & Bernard, M. F. Scared to death? The effects of intimidation and consumption in predator-prey interactions. Ecology 86, 1583–1592 (2005).
3. Peckarsky, B. L. et al. Revisiting the classics: considering nonconsumptive effects in textbook examples of predator-prey interactions. Ecology 89, 2416–2425 (2008).
4. Ellenberg, U., Mattern, T., Seddon, P. J. & Jordua, G. L. Physiological and reproductive consequences of human disturbance in Humboldt penguins: the need for species-specific visitor management. Biol. Conserv. 133, 95–106 (2007).
5. Baudains, T. & Lloyd, P. Habitatiation and habitat changes can moderate the impacts of human disturbance on shorebird breeding performance. Anim. Conserv. 10, 400–407 (2007).
6. Møller, A. P. Flight distance and population trends in European breeding birds. Behav. Ecol. 19, 1095–1102 (2008).
7. Møller, A. P., Sama, D. S. M., Weston, M. A., Guay, P. J. & Blumstein, D. T. American exceptionalism: population trends and flight initiation distances in birds from three continents. PLoS ONE 9, e107883 (2014).
8. Shochat, E., Warren, P. S., Faeth, S. H., McIntyre, N. E. & Hope, D. Patterns emerging processes in mechanistic urban ecology. Trends Ecol. Evol. 21, 186–191 (2006).
9. Markovich-Nicholls, L. et al. Relationships between human disturbance and wildlife land use in urban habitat fragments. Conserv. Biol. 22, 99–109 (2008).
10. Blumstein, D. T. & Fernández-Juricic, E. A Primer of Conservation Behavior (Sinauer, 2010).
11. Hedges, L. V. Distribution theory for Glass’s estimator of effect size and related estimators. J. Educ. Stat. 6, 107–128 (1981).
12. Burnham, K. P. & Anderson, D. R. Model Selection and Multimodel Inference: a Practical Information-Theoretic Approach (Springer, 2002).
13. McKinney, M. L. Urbanization, biodiversity, and conservation. Bioscience 52, 883–890 (2002).
14. Miranda, A. C., Schieflitz, H., Sonntag, T. & Partecke, J. Urbanization and its effects on species personality traits: a result of microevolution or phenotypic plasticity? Glob. Chang. Biol. 19, 2634–2644 (2013).
15. Higgins, J. P. T., Thompson, S. G., Deeks, J. J. & Altman, D. G. Measuring inconsistency in meta-analyses. BMJ 327, 557–560 (2003).
16. Egger, M., Smith, G. D., Schneider, M. & Minder, C. Bias in meta-analysis detected by a simple, graphical test. BMJ 315, 629–634 (1997).
17. Nakagawa, S. & Santos, E. S. A. Methodological issues and advances in biological meta-analysis. Evol. Ecol. 26, 1253–1274 (2012).
18. Møller, A. P. Interspecific variation in fear responses predicts urbanization in birds. Behav. Ecol. 21, 365–371 (2010).
19. Li, C., Monclús, R., Maul, T. L., Jiang, Z. & Blumstein, D. T. Quantifying human disturbance on antipredator behavior and flush initiation distance in yellow-bellied marmots. Appl. Anim. Behav. Sci. 129, 146–152 (2011).
20. Levey, D. J. et al. Urban mockingbirds quickly learn to identify individual humans. Proc. Natl Acad. Sci. USA 106, 8959–8962 (2009).
21. Engelhardt, S. C. & Weladji, R. B. Effects of levels of human exposure on flight initiation distance and distance to refuge in foraging eastern gray squirrels (Sciurus carolinensis). Can. J. Zool. 89, 823–830 (2011).
22. McGiffin, A., Lill, A., Beckman, J. & Johnstone, C. P. Tolerance of human approaches by Common Mynas along an urban-rural gradient. Emu 113, 154–160 (2013).
23. Blumstein, D. T. Developing an evolutionary ecology of fear: how life history and natural history traits affect disturbance tolerance in birds. Anim. Behav. 71, 269–269 (2006).
24. Fernández-Juricic, E. et al. Relationships of anti-predator escape and post-escape responses with body mass and morphology: a comparative avian study. Evol. Ecol. Res. 8, 731–752 (2006).
25. Gotanda, K. M., Turgeon, K. & Kramer, D. L. Body size and reserve affect flight initiation distance in parrotfishes. Behav. Ecol. Sociobiol. 63, 1563–1572 (2009).
26. Sama, D. S. M., Blumstein, D. T., Stankovich, T. & Cooper, Jr W. E. Fifty years of chasing lizards: new insights advance optimal escape theory. Biol. Rev. doi:10.1111/brv.12173 (2015).
27. Cooper, Jr W. E. & Blumstein, D. T. (eds) Escaping from Predators: an Integrative View of Escape Decisions (Cambridge Univ., 2015).
28. Vedengberg, R. C. & Dill, L. M. The economics of fleeing from predators. Adv. Study Behav. 16, 229–249 (1986).
29. Norberg, U. M. Vertebrate Flight: Mechanics, Physiology, Morphology, Ecology and Evolution (Springer Science & Business Media, 2012).
30. Witter, M. S., Cuthill, I. C. & Bonser, R. H. C. Experimental investigations of mass-dependent predation risk in the European starling. Starnus vulgaris. Anim. Behav. 48, 201–222 (1994).
31. Stankovich, T. & Blumstein, D. T. Fear in animals: a meta-analysis and review of risk assessment. Proc. R. Soc. B 272, 2627–2634 (2005).
32. Bennett, P. M. & Harvey, P. H. Active and resting metabolic rate in birds: allometry, phylogeny and ecology. J. Zool. 213, 327–344 (1987).
33. Werner, E. E., Gilliam, J. F., Hall, D. J. & Mittelbach, G. G. An experimental test of the effects of predation risk on habitat use in fish. Ecology 64, 1540–1548 (1983).
34. Sama, D. S. M., Møller, A. P. & Blumstein, D. T. Brain size as a driver of avian escape strategy. Sci. Rep. 5, 11193 (2015).
35. Fernández-Juricic, E., Venier, M. P., Renison, D. & Blumstein, D. T. Sensitivity of wildlife to spatial patterns of recreationist behavior: a critical assessment of minimum approaching distances and buffer areas for grassland birds. Biol. Conserv. 125, 225–235 (2005).
36. Trivers, R. L. in Sexual Selection & the Descent of Man (ed. Campbell, B.) 136–179 (Aldine Publishing Company, 1972).
37. Yampolsky, L. Y. Life history theory. Environ. Life Sci. 1–6, doi: 10.1038/npg.els.0003219. (2002).
38. Ghalambor, C. K. & Martin, T. E. Fecundity-survival trade-offs and parental risk-taking in birds. Science 292, 494–497 (2001).
39. Carrete, M. & Tella, J. L. High individual consistency in fear of humans throughout the adult lifespan of rural and urban burrowing owls. Sci. Rep. 3, 3524 (2013).
40. Möller, A. P. Flight distance of urban birds, predation, and selection for urban life. Behav. Ecol. Sociobiol. 63, 63–75 (2008).
41. Blumstein, D. T. in Avian Urban Ecology: Behavioural and Physiological Adaptations (eds Gil, D. & Brumm, H.) 41–53 (Oxford Univ., 2014).
42. Herrero, S., Smith, T., DelBruyn, T. D., Grunther, K. & Matt, C. A. From the field: Brown bear habituation to people—safety, risks, and benefits. Wildl. Soc. Bull. 33, 362–373 (2005).
43. Shalter, M. D. in Habituation, Sensitization, and Behavior (ed. Pecke, H.) 349–391 (Elsevier, 1984).
44. Curio, E. Probleme des feinderkennens bei vogeln. Proc. Int. Ornithol. Congr. 13, 206–239 (1963).
45. Ikuta, L. A. & Blumstein, D. T. Do fences protect birds from human disturbance? Biol. Conserv. 112, 467–482 (2003).
46. Dixo, R. et al. Defaunation in the Anthropocene. Science 345, 401–406 (2014).
47. Kasereka, B., Muhigwa, J.-B. B., Shalukoma, C. & Kahekwa, J. M. Vulnerability of habituated Grauer’s gorilla to poaching in the Kahuzi-Biega National Park, DRC. Afr. Study Monogr. 27, 15–26 (2006).
48. Colinvaux, P. A. Why Big Fierce Animals are Rare: an Ecologist’s Perspective (Princeton Univ., 1978).
49. Primack, R. B. Essentials of Conservation Biology (Sinnauer Associates, 2006).
50. Urban, M. C. Accelerating extinction risk from climate change. Science 348, 571–573 (2015).
51. Ceballos, G. et al. Accelerated modern human-induced species losses: Entering the sixth mass extinction. Sci. Adv. 1, e1400253 (2015).
52. Blumstein, D. T. & Berger-Tal, O. Understanding sensory mechanisms to develop effective conservation and management tools. Curr. Opin. Behav. Sci. 6, 13–18 (2015).
53. Koricheva, J., Gurevitch, J. & Mengersen, K. Handbook of Meta-Analysis in Ecology and Evolution (Princeton Univ., 2013).
54. Viechlbauer, W. Conducting meta-analyses in R with the metafor package J. Stat. Softw. 36, 1–48 (2010).
55. Pagel, M. Inferring the historical patterns of biological evolution. Nature 401, 877–898 (1999).
56. Del Hoyo, J., Elliott, A., Sargatal, J., Christie, D. A. & de Juana, E. Handbook of the Birds of the World Alive. Available at http://www.hbw.com/ (Accessed on 14 November 2014).
57. Zuur, A. F., Ieno, E. N. & Elphick, C. S. A protocol for data exploration to avoid common statistical problems. Methods Ecol. Evol. 1, 3–14 (2010).
58. Alexander, R. D. The evolution of social behavior. Annu. Rev. Ecol. Syst. 5, 325–383 (1974).
59. Hamilton, W. D. Geometry for the selfish herd. J. Theor. Biol. 31, 295–311 (1971).
60. Pulliam, H. On the advantages of flocking. J. Theor. Biol. 38, 419–422 (1973).
61. Griffin, A. S. Social learning about predators: a review and prospectus. Learn. Behav. 32, 131–140 (2004).

62. Burish, M. J., Kueh, H. Y. & Wang, S. S.-H. Brain architecture and social complexity in modern and ancient birds. Brain. Behav. Evol. 65, 107–124 (2004).
63. Knight, R. L. & Fitzner, R. E. Human disturbance and nest site placement in black-billed magpies. J. F. Ornithol. 56, 153–157 (1985).
64. Datta, T. & Pal, B. C. The effect of human interference on the nesting of the openbill stork Anastomus oscitans at the Raiganj Wildlife Sanctuary, India. Biol. Conserv. 64, 149–154 (1993).
65. Blumstein, D. T. et al. Avian risk assessment: effects of perching height and detectability. Ethology 110, 273–285 (2004).
66. Tyrrell, L. P. & Fernández-Juricic, E. in Escaping from Predators: an Integrative View of Escape Decision (eds Cooper Jr, W. E. & Blumstein, D. T.) (Cambridge Univ., 2015).
67. Blumstein, D. T. Flush early and avoid the rush: a general rule of antipredator behavior? Behav. Ecol. 21, 440–442 (2010).
68. Cooper, Jr W. E., Pyron, R. A. & Garland, T. Island tameness: living on islands reduces flight initiation distance. Proc. R. Soc. B 281, 20133019 (2014).
69. Barton, K. MulMfn: Multi-model inference. R package version 3.1. 2 (2014).

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D.S.M.S. and D.T.B. conceived the ideas for the paper and its structure; D.S.M.S. compiled published data and conducted meta-analysis with technical help from S.N.; D.S.M.S., S.N., F.N., T.F.R. and D.T.B. discussed results; D.S.M.S. and D.T.B. wrote the first draft of the manuscript; D.S.M.S., S.N., F.N., T.F.R. and D.T.B. edited the manuscript.

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