The value of coarse species range maps to inform local biodiversity conservation in a global context

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Range maps of thousands of species, compiled and made freely available by the International Union for Conservation of Nature, are being increasingly applied to support spatial conservation planning. However, their coarse nature makes them prone to commission and omission errors, and they lack information on the variations in abundance within species’ distributions, calling into question their value to inform decisions at the fine scales at which conservation often takes place. Here, we tested if species ranges can reliably be used to estimate the responsibility of sites for the global conservation of species. We defined ‘specific responsibility’ as the fraction of a species’ population within a given site, considering it useful for prioritizing species within sites; and defined ‘overall responsibility’ as the sum of specific responsibility across species within a site, assuming it informative of priorities among sites. Taking advantage of an exceptionally detailed dataset on the distribution and abundance of bird species at a near-continental scale – a level of information rarely available to local decision-makers – we created a benchmark against which we tested estimates of responsibility derived from range maps. We investigated approaches for improving these estimates by complementing range maps with plausibly available local data. We found that despite their coarse nature, range maps provided good estimates of sites’ overall responsibility, but relatively poor estimates of specific responsibility. Estimates were improved by combining range maps with local species lists or local abundance data, easily available through local surveys on the sites of interest, or simulated expert knowledge. Our results suggest that combining range maps with local data is a promising route for improving the effectiveness of local conservation decisions at contributing to reducing global biodiversity losses. This is all the more urgent in hyper-diverse poorly-known regions where conservation-relevant decisions must proceed despite a paucity of biodiversity data.
well-known as birds, such as amphibians or insects. For comparison, > 75% of the world’s protected areas are smaller than 100 km² (IUCN and UNEP-WCMC 2012).

Furthermore, given that range maps only represent the limit of species’ distributions, they contain no information on the variation in abundances across species’ ranges. Their application to conservation planning therefore also assumes (implicitly or explicitly) that all sites where a species occurs are equally important to the species’ conservation. Indeed, area targets commonly applied in conservation planning (i.e. that a given species should be represented in protected areas by at least X% of its range; Rodrigues et al. 2004b, Pompa et al. 2011, Pouzols et al. 2014), make sense under the assumption that the fraction of the species’ range is a good surrogate for the fraction of the total population protected. In practice, however, this is not necessarily true: abundance often varies substantially across species’ ranges, with most species being rare at most sites where present and abundant at relatively few (Gaston 2003). Information on the fraction of each species’ global population at a given site is used to identify sites of global importance for conservation, including Ramsar Wetlands of International Importance (> 1%; Convention on Wetlands 1971) and Alliance for Zero Extinction sites (> 95%; Ricketts et al. 2005). However, such information is seldom available to local conservation planners, as (other than for species with very small ranges or that aggregate conspicuously, which can be censused relatively easily) obtaining it requires much human and monetary investment.

Given these limitations, the application of range maps to conservation planning may lead both to wasted investments to protect species in places of little or no value to their conservation, and conversely to missed opportunities to invest strategically in high-value sites. In order to quantify these effects, we focus here on the percentage of each species’ population within a site as a measure of the site’s responsibility for the global conservation of the species. We consider both this measure of ‘specific responsibility’ (defined for each species in each site) and an aggregated measure of ‘overall responsibility’ (defined for each site, as the sum of the specific responsibility values for all species in the site). We evaluate the effectiveness of coarse range maps for informing conservation decisions at the local scale by comparing estimates of specific and overall responsibility derived from IUCN range maps with a high quality benchmark obtained from an exceptionally detailed dataset on the distribution and abundance of bird species at a near-continental scale (Sauer et al. 2007). This allows us to go further than previous studies that analysed the effectiveness of range maps in representing species’ occupancy (Graham and Hijmans 2006, Hurlbert and Jetz 2007, Cantú-Salazar and Gaston 2013) by also evaluating the performance of range maps in representing the abundance structure of real distributions.

Furthermore, we test practical approaches for improving responsibility estimates by complementing range maps with other data plausibly available to local managers. We start by testing an approach for overcoming the limitations of range maps in representing occupancy, and then test two approaches for also taking into account the abundance structure of species’ distributions. From these results, we derive practical recommendations for making the best use of available data to improve estimates of site responsibility for the conservation of species, as a contribution to better inform local-scale decision-making within a global biodiversity conservation context.

Material and methods

All calculations and analyses were done in R (R Development Core Team), particularly the ‘maptools’ R package (Bivand and Lewin-Koh 2010). All maps were done in ArcGIS 10.1, using the North America Albers Equal Area Conic projection (ESRI 2012).

Species distribution data

We obtained from the IUCN Red List of threatened species range maps representing the global breeding distribution of 316 land birds in the United States and Canada (BirdLife International and NatureServe 2012). We considered only parts of the species’ range where they are classified as extant or probably extant, native or reintroduced, and where they occur as breeding migrants or as residents.

We obtained high-quality distribution and abundance data from the North American Breeding Bird Survey (BBS; Sauer et al. 2007) for the same 316 species. The BBS is a long-term (since 1966) annual avian monitoring program, in which volunteers count the numbers of birds observed along roadsides survey routes. There are currently 3700 active BBS routes, of which nearly 2900 are surveyed annually, following a standardized protocol: each survey route is randomly located and 24.5 miles long, with a 3-min point count conducted at 0.5-mile intervals, during which every bird seen or heard within a 0.25-mile radius is recorded. In our analyses, we used a derived dataset consisting of a layer of estimated relative abundance per species across an orthogonal grid (cell size ~ 462 km²; hereinafter referred to as ‘sites’, and corresponding to a ‘local scale’). This was obtained by combining the 2006–2010 raw annual count data, first by calculating the average number of individuals counted per route, and then by smoothing these average counts across sites, cutting values below a threshold of minimal relative abundance of 0.1. For many of sites, particularly in regions with sparser BBS coverage, abundance values are therefore estimates derived from count data in neighbouring areas. This averaging and smoothing likely reduce the noise created by natural spatial and temporal variabilities (e.g. reducing the impact of records of vagrant individuals), making the BBS data more comparable to the IUCN range maps. The final relative abundance layer predicts for each site the average number of birds of the species that could be seen in about 2.5 h of birdwatching along roadsides, by very good birders (for details see Sauer et al. 2007).

Our study region covers the contiguous United States and southern Canada (up to latitude 55°N), corresponding to 23 608 sites (each with an area of ca 462 km²). For each site, we had therefore: species presence/absence derived from the IUCN range maps; and species’ relative abundances (and hence also their presence/absence) derived from the high-quality BBS dataset. The latter is not perfect – as in other data obtained from field surveys, it is biased towards some
species and habitats (Keller and Scallan 1999, Bibby et al. 2000) – but it is the best available fine-scale dataset on the distribution and abundance of a taxonomic group at a near-continental scale. For the purposes of these analyses, we treated it as if it perfectly represented species’ distributions.

Quantifying commission and omission errors

For each species, we calculated (Fig. 1): 1) percentage of commission errors: percentage of the total number of sites overlapping the range map from where the species is absent according to the BBS data. 2) Percentage of omission errors: number of sites not overlapping the species range map but where the species is present according to the BBS data, in relation to the total number of sites overlapping the species range.

We investigated the distribution of both types of errors across species through histograms splitting species into 10%-width classes of error. In addition we investigated how commission and omission errors related to species’ range sizes by analysing the Spearman rank-order correlation between the mid-interval error value and the median range size of the species in each class.

Calculating reference values of ‘specific responsibility’ and ‘overall responsibility’ for each site

We use the term ‘specific responsibility’ to refer to the percentage of the population of a given species that is found at a given site. We consider this a measure of the extent to which the site affects the global conservation options for the species, while recognising that in real life this may not be a perfect measure (e.g. it may under or over-estimate site value because of source-sink population dynamics [Heinrichs et al. 2010]).

We calculate reference values of ‘specific responsibility’ from the BBS data. For species whose global range falls completely within the study region, this was calculated as:

\[
\frac{\text{species' local abundance in the site}}{\text{sum of species' local abundance across study region}} \times 100
\]

For any species whose range falls partially outside the study region, values were corrected by taking into account the fraction of the species’ global range outside the region:

\[
\frac{\text{species' local abundance in the site}}{\text{sum of species' local abundance across study region}} \times \frac{\text{IUCN global range } \cap \text{ study region}}{\text{IUCN global range}} \times 100
\]

where ‘IUCN global range \cap study region’ represents the area of overlap between the species’ global range and the study region. This correction assumes that average abundance is similar inside and outside the study region (such that, for example, if the study region covers 50% of the range, it is assumed that it includes 50% of the global population). Failure to meet this assumption may affect the veracity of the calculations of specific responsibility for these species, which may thus underestimate or overestimate the real fraction of the species’ total population in each site. However, this is unlikely to affect the conclusions of this study, whose purpose is not to identify real-life conservation priorities in North-America, but to compare approaches for better estimating reference values of responsibility assumed as perfect.

For any given site, we then calculated ‘overall responsibility’, as the sum of specific responsibility values for all the species present at the site.

Metrics for measuring the value of range maps to estimate site responsibility

We estimated specific responsibility of each site for each species as the percentage of the species’ range overlapping the site. Overall responsibility was then estimated as the sum of specific responsibility values for all species whose ranges overlap the site. We evaluated the quality of these estimates by comparing them with reference values of responsibility, using a diversity of metrics (Table 1) that compare values either in terms of relative order (to investigate if estimates are good indicators of relative priorities) or in terms of absolute values (to understand if estimates can be used as direct measures of absolute responsibility).

Figure 1. Examples of the comparison between species range maps (from the IUCN data) and areas of occupancy (from the BBS data) for two of the bird species analyzed: correct presences are sites where the two types of distribution data match; commission errors are sites where the species is present according to the range maps but absent according to the BBS data; omission errors are sites where the species is absent according to the range maps but present according to the BBS data.
Table 1. Effectiveness of estimates of sites’ specific responsibility and of overall responsibility, obtained either from range maps alone, or by combining those maps with plausibly available data through different approaches: Approach 1 (range maps + local species lists); Approach 2a (range maps + local species lists + continental expert knowledge); Approach 2b (range maps + local species lists + regional expert knowledge); Approach 3 (range maps + local species lists + local abundance data). Effectiveness was assessed in each case through metrics that contrast estimated values with the corresponding reference values, either in terms of order or in terms of absolute values. For Approaches 2 and 3, results are presented for respectively $m = 8$ and $v = 1$.

<table>
<thead>
<tr>
<th>Type</th>
<th>Metric</th>
<th>Range maps only</th>
<th>Approaches combining range maps + local/contextual data</th>
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<tr>
<td>Metrics for comparing specific responsibility values (for comparing among species within each site)</td>
<td></td>
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<td>1</td>
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<tr>
<td>Metrics comparing order</td>
<td>$J_{\text{median}}$ (first, third quartiles): median Jaccard’s coefficient; $J$ of a site is the degree of overlap between the top (25%) species in terms of estimated vs reference values of specific responsibility; varies between zero (no overlap) and 1 (perfect coincidence).</td>
<td>0.26</td>
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<td>Metrics comparing absolute values</td>
<td>$S_{\text{median}}$ (first, third quartiles): median Spearman’s rank coefficient; $S$ of a site is the correlation between the ranks of estimated and reference values of specific responsibility; varies between −1 (order of species exactly reversed) and 1 (same exact order).</td>
<td>0.26</td>
<td>0.39</td>
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<td>[0.11, 0.40]</td>
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<td>Metrics for comparing overall responsibility values (for comparing sites)</td>
<td>$R^2_{\text{median}}$ (first, third quartiles): median coefficient of determination; $R^2$ of a site is the proportion of variability explained by the linear regression model with estimated specific responsibility as predictor and reference values as the response (forced through the origin); varies between 0 (no explanatory power) and 1 (response perfectly explained by the predictor).</td>
<td>0.11</td>
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| Metrics comparing order | $S$: Spearman’s rank correlation coefficient between the ranks of estimated and reference site values of overall responsibility; varies between −1 (order of sites exactly reversed) and 1 (same exact order). | 0.47 | 0.69 | 0.84 | 0.68 | 0.90 |
| | $P$: proportion of pairs of sites for which the order based on estimated overall responsibility agrees with the order based on reference values; varies between 0 (perfect disagreement) and 1 (perfect agreement). | 0.67 | 0.75 | 0.83 | 0.75 | 0.87 |
| | $R^2$: coefficient of determination measuring the proportion of variability explained by the linear regression model where estimated overall responsibility is the predictor and reference responsibility is the response variable (forced through the origin); varies between 0 (no explanatory power) and 1 (response perfectly explained by the predictor). | 0.83 | 0.87 | 0.94 | 0.87 | 0.95 |
| | slope: slope of the line obtained from the linear regression model as obtained for $R^2$ above; slope < 1 when estimated responsibility generally overestimate reference values; slope > 1 corresponds to a tendency for underestimation. | 0.66 | 1.01 | 0.80 | 0.84 | 2.29 |
Improving the information value of range maps by combining these with plausibly available data

We explored three approaches for improving the estimates of responsibility derived from coarse range maps, by integrating local and/or contextual data plausibly available to local decision-makers into the calculations of specific responsibility: Approach 1 focuses on correcting the errors of range maps in representing occupancy, whereas Approaches 2 and 3 also take into account the abundance structure of species' distributions. Calculations of specific responsibility vary between approaches, but in all cases, overall responsibility was calculated the sum of specific responsibility values within a site.

We used the BBS occurrence data to simulate for each site the local species list that a decision-maker would have been able to obtain through local surveys in his site(s) of interest. In Approach 1 (range maps + local species lists) we then estimated specific responsibility values as above – from the percentage of species' IUCN global range overlapping each site – but considering only, and all, those species effectively present at each site according to the BBS data. Hence, in each site we considered that specific responsibility was zero for species present according to the range maps but absent according to the BBS data (thus correcting commission errors) whereas for species present according to the BBS data and absent according to the range maps specific responsibility was calculated as:

\[
\frac{1}{\text{number of sites overlapped by IUCN global range within study region}} \times 100
\]

(thus correcting omission errors and assuming the incremental correction in species range size by the focal site as negligible).

Correcting the fact that range maps have no information on the variation in species' abundance within their ranges is more complex. Estimating the degree to which a site is responsible for the global conservation of a species ideally requires data on the species' abundance not only in the site of interest but also throughout its whole range, seldom obtainable by local managers. But even in the absence of such quantitative data, it may be useful to distinguish qualitatively whether a site is particularly valuable for a species or not. We tested two approaches for doing so based on data plausibly available to a local manager: Approach 2 highlights species' peaks of abundance, building from the knowledge that most species are relatively abundant at just a few sites within their range (Gaston 2003); Approach 3 disregards local records of low relative abundance, given that species are relatively rare in most of the sites where they are present (Gaston 2003).

Approach 2 assumes that qualitative data on the relative abundance of species across their ranges already exists in the heads of experts with field experience across a broad region, who would be able to recognise if a site is exceptionally important for a particular species. We simulated this using the BBS data, by considering a species as 'exceptionally abundant' at a site if its local abundance is >10 times median abundance. We then calculated specific responsibility for each species in each site as in Approach 1, but weighting exceptionally abundant species by a factor \(m\). We explored \(m\) values from 2 to 10 (higher values led to smaller slope values, resulting in bigger overestimation errors, and reduced the overall benefit of this approach). We considered two spatial extents of expert knowledge: continental or regional. In Approach 2a (range maps + local species lists + continental expert knowledge), experts were presumed able to identify records of exceptional local abundance across North America. However this scenario is arguably unrealistic, since it assumes experts with field knowledge across the entire ranges of all species in the site(s) of interest. In Approach 2b (range maps + local species lists + regional expert knowledge), experts would only have a regional perspective of species relative abundances, simulated by identifying separately exceptional abundance records within each US State or Canadian Province.

Finally, in Approach 3 (range maps + local species lists + local relative abundance), we tested the effect of dismissing species with low local abundance as a way of increasing emphasis on the remaining species. We used BBS data on the relative abundance of species in each site to simulate the type of data a local manager could potentially obtain through field surveys. We then calculated specific and overall responsibility as in Approach 1, but assuming zero specific responsibility for all species with local abundance < \(v\). We explored \(v\) values from 1 to 50.

We evaluated the performance of the estimates of specific and overall responsibility obtained for each of these approaches by comparing them with the corresponding reference values, using the same metrics as described above (Table 1).

Results and discussion

Commission and omission errors in range maps

Consistent with previous studies (Rodrigues et al. 2004a, Hurlbert and Jetz 2007, Cantú-Salazar and Gaston 2013), we found that range maps are highly prone to commission errors: >10% for more than half of all species, and >30% for nearly a tenth (Fig. 2a). Commission errors are expected in range maps, which generally represent species’ broad extent of occurrence and often include areas of unsuitable habitat (Gaston and Fuller 2009). Some of these errors may also reflect false absences in the BBS data, although we expect this effect to be small given the sampling effort that went into the BBS data. Species with lower levels of commission errors tend to have smaller range sizes (\(S = 0.7, p = 0.04\); Fig. 2a; although there was wide variability across species), possibly reflecting a higher precision in the mapping of the boundaries of smaller ranges.

We found even higher rates of omission errors: >10% for nearly 80% of all species and >30% for one-third of species (Fig. 2b). This was unexpected since range maps are considered much more prone to range overestimation than to underestimation (Rondinini et al. 2006), even though other previous studies had also found non-trivial levels of omission errors in range maps (Rodrigues et al. 2004a, Beresford et al. 2011, Cantú-Salazar and Gaston 2013; see also Rodrigues 2011). Omission errors may result from an underestimation of species’ range extension, but may also arise at the edge of ranges from the coarse way in which range maps are drawn.
(Cantú-Salazar and Gaston 2013). In our particular analysis, high levels of omission errors are possibly an artifact of the smoothing used to construct the BBS abundance layers resulting in a ‘spillage’ of predicted presence into cells where the species is absent (Sauer et al. 2007). This effect is likely higher in species with larger perimeter-to-area ratios, and so it may explain our finding that omission errors were more proportionately frequent in species with small ranges ($S = -0.90$, $p < 0.001$, Fig. 2b; see also example in Fig. 1).

Overall, the high rates of commission and omission errors we found reinforce calls for caution in the uncritical use of range maps as sources of data on the presence or absence of species at fine spatial scales (Hurlbert and Jetz 2007). Omission errors can affect the efficiency of conservation planning by biasing results towards known species occurrences, potentially missing important areas where a species also exists. But commission errors are arguably even more problematic, if they lead to species being falsely believed conserved when they are not (Rodrigues et al. 2004a).

**Value of range maps for estimating site responsibility**

Overall responsibility estimated from range maps was generally a good predictor ($R^2 = 0.83$; Table 1, Fig. 3a) and captured the broad spatial patterns of reference values (Fig. 4a, b), even if estimates tended to largely overestimate reference values ($slope < 1$). Agreement in relative order was less impressive ($S = 0.47$), and a manager using solely range maps to prioritize between two sites would be wrong one out of three times ($p = 0.67$).

Within most sites, estimates of specific responsibility obtained from range maps were weak predictors of reference values, in terms of order ($S_{median} = 0.26$) and even more so in terms of absolute values ($R^2_{median} = 0.11$). For most sites, there was also little overlap between the top 25% species identified using estimated vs. reference values of specific responsibility ($J_{median} = 0.26$). In other words, a site manager using only range maps would have limited information on the relative responsibility of its site for the global conservation of species.

Commission errors are likely to contribute substantially to the shortcoming of range maps as bases for estimates of responsibility. Indeed, they lead to species being falsely considered as present in sites, and thus their specific responsibilities being considered the same as the average within the species range (rather than zero, as they should be), which then adds up to overestimated overall responsibility values. In contrast, omission errors are less likely to affect the performance of range maps. Indeed, true omission errors (caused by a lack of knowledge of species presence or imprecise mapping) are likely to correspond to regions where species are relatively rare, and hence sites of low specific responsibility, not very different from the zero value obtained from range maps. In this exercise, as discussed previously, apparent omission errors may also arise from a ‘spillage’ artifact of the smoothing used to generate the BBS abundance layers. Cells affected by such spillage are falsely considered as presences in the BBS data, but have low abundances (Sauer et al. 2007) and consequently low values of specific responsibility (not very different from the actual zero value).

**Performance of approaches for improving the information value of range maps**

Combining range maps with local species lists (Approach 1) improved estimates of responsibility for all metrics of effectiveness tested (Table 1), for both specific ($R^2_{median}$ increased from 0.11 to 0.19; $J_{median}$ from 0.26 to 0.31; $S_{median}$ from 0.26 to 0.39) and overall responsibility ($R^2$ increased from 0.83 to 0.87; $S$ from 0.47 to 0.69; and $P$ from 0.67 to 0.75). It also corrected the tendency of estimates from range maps to largely overestimate reference values ($slope < 1$) (Fig. 3b), and resulted in a more realistically detailed spatial pattern of estimated overall responsibility (Fig. 4c, b).
than in the baseline Approach 1 (corresponding to \( m = 1 \); Supplementary material Appendix 2, Table A2, Fig. A2). For \( m = 8 \), there were marked improvements for all metrics comparing specific responsibility (\( R^2_{\text{median}} \), increased from 0.19 to 0.42; \( f_{\text{median}} \), from 0.31 to 0.35; and \( S_{\text{median}} \), from 0.39 to 0.45), in the spatial pattern of overall responsibility (Fig. 4d) and in all but one metrics comparing overall responsibility (e.g., \( R^2 \), increased from 0.87 to 0.94; \( S \), from 0.69 to 0.84). The only exception was a lowering of slope (from 1.01 to 0.80) indicating an increase in overestimation errors (Supplementary material Appendix 2, Fig. A2), and meaning that these estimates should be seen as informative for comparisons between sites, rather than as predictors of the real values.

Unfortunately, these results do not hold in the more plausible Approach 2b, whereby experts’ field experience is restricted to the regional scale: disappointingly, results were no better than those in the baseline Approach 1 (Table 1; Fig. 4e). This is because species’ relative abundances vary substantially across regions within their ranges

However, this approach fails to highlight sites of particularly high overall responsibility, underestimating by a factor of 2 or 3 those with the highest reference values (Fig. 3b). A site has an exceptionally high overall responsibility when it combines a high species richness (Supplementary material Appendix 1, Fig. A1a) and a prevalence of species with small global ranges (i.e., high mean site endemism; Supplementary material Appendix 1, Fig. A1b) and with high relative abundances in the site in relation to their abundances elsewhere (i.e., high mean species’ normalised local abundance; Supplementary material Appendix 1, Fig. A1c). Approach 1 combines information from the first two facets of responsibility, but not from the third. We explored ways of combining the three in Approaches 2 and 3.

In Approach 2a, species of exceptionally high local abundance in the context of North American ranges (as identified by simulated expert knowledge) were then given a higher weight (\( m > 1 \)) when estimating specific and overall responsibility. For \( m \) values from 2 to 10, estimates of both specific and overall responsibility performed better than in the baseline Approach 1 (corresponding to \( m = 1 \); Supplementary material Appendix 2, Table A2, Fig. A2).
In Approach 3 we simulated the effect of disregarding (assuming zero specific responsibility) the locally rare species in each site, i.e., those with a local abundance $v$. We found that for $v = 1$, estimates of responsibility were much better than in Approach 1 (corresponding to $v = 0$) and as good, or even better, than for Approach 2a (e.g, $S_{\text{median}} = 0.67$, compared with 0.39 in Approach 1 and 0.4 in Approach 2a; $R^2 = 0.95$, compared to 0.87 in Approach 1 and 0.94 in Approach 2a; Table 1; Fig. 3e), and reproduced very high abundance of a species within the context of a region and yet be quite unexceptional across the species' entire range. Unfortunately, these results suggest that relying on expert knowledge to provide a broader context to local species' abundances is of little value if such knowledge is geographically restricted. This is thus an example of how adding more information does not necessarily result in better estimates.

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well the spatial pattern of reference overall responsibility (Fig. 4f). Nonetheless, absolute values of overall responsibility were systematically (and substantially) underestimated (slope = 2.29; Fig. 3e), and so again these estimates should be seen as informative for comparisons between sites, rather than as predictors of the real values. The higher the abundance threshold \( v \), the steeper the slope (the higher the underestimation errors), and for \( v > 10 \), the estimates ceased from being better than in Approach 2a in terms of \( R^2 \) and became progressively worse (Supplementary material Appendix 3, Fig. A4). These results illustrate the advantages as well as the risks with this approach. Disregarding low abundance records improves responsibility estimates because in any given site it excludes those species that are locally rare but have large populations elsewhere (for which the site has little responsibility). However, it also excludes species that are rare everywhere, which may be in need of conservation attention in all sites where they occur. Accordingly, we found that increased \( v \) increased the number of species excluded from all sites where they occurred (Supplementary material Appendix 3, Fig. A4). This suggests that disregarding low abundance records may be an effective strategy for improving estimates of responsibility, but only for low thresholds \( v \). Managers may also want to avoid excluding any species known to occur at low densities throughout their ranges.

**Generalizability of the results**

We investigated how species range maps such as those made freely available by the IUCN can inform local-scale conservations decisions. We have focused on the particular case of breeding land birds in North America because the exceptional BBS dataset allowed us to obtain high quality reference values of responsibility against which we could test different approaches for making the best use of data plausibly available to local managers. It is pertinent to ask whether the results we obtained can be generalized to other regions and to other taxa, i.e., whether managers working at local scales (of a similar order of magnitude as in our analyses: cell size = 460 km\(^2\)) but in other parts of the world and/or focused on other taxa would also find that range maps combined with local data provide a good basis for estimating the responsibility of sites for global species conservation.

Species range maps are likely to be of lower quality (higher levels of commission and of omission errors) in regions with lower levels of expertise (e.g., in Africa; Rodrigues et al. 2010), particularly for the most diverse taxonomic groups (e.g., invertebrates, plants; Gaston and May 1992). Furthermore, in regions with a stronger human footprint (e.g., high deforestation rates), species’ areas of occupancy may have become artificially patchy (e.g., restricted to remaining forest fragments), further increasing commission errors in range maps.

The good estimates we found for birds in North America, even when using range maps alone (Table 1) are thus not necessarily generalizable to lesser-known taxa and/or in lesser-studied regions. Furthermore, even obtaining a complete local species list can be quite an ordeal in high-diversity areas (Lawton et al. 1998). But although these limitations are certain to reduce the quality of estimates of responsibility for hyper-diverse sites or taxa and in poorly known regions, the approaches we propose remain applicable, and making a good use of the available data is even more important in those circumstances.

Furthermore, we believe that our result that the range maps of more narrowly distributed species are generally less prone to commission errors (Fig. 2a) is likely to generalize more widely. Indeed, range boundaries can be drawn with higher accuracy if species are associated with particular geographical features (e.g., islands, valleys, mountaintops) that can be mapped precisely. The maps of widespread species are also more likely to encompass wide areas of unsuitable habitat: for example, a widespread wetland species will be mapped over large territory encompassing much non-wetland habitat, whereas a restricted-range wetland species will be mapped into the specific wetland(s) where it is known to occur. Maps of restricted-range species are thus mapped with a higher precision, and accordingly they have previously been used to pinpoint sites of particular importance to conservation, such as those identified by the Alliance for Zero Extinction (Ricketts et al. 2005), Key Biodiversity Areas (Langhammer et al. 2007), and highly irreplaceable protected areas (Le Saout et al. 2013).

**Conclusions and perspectives**

The on-going global erosion of biodiversity (Butchart et al. 2010) is the cumulative result of myriads of impacts affecting species worldwide, mainly through the destruction and degradation of their habitats (Vié et al. 2009). Halting such erosion – as the World’s governments have committed to do (Secretariat of the Convention on Biological Diversity 2002, 2010) – requires a diversity of actions, not only dedicated conservation measures (e.g., protected areas) but also broader actions to mitigate the impacts of the human enterprise (e.g., of agricultural expansion, urbanisation, new infrastructure).

The combined impact of these local actions relies therefore on the extent to which together they affect options for global biodiversity conservation. In the past few decades, the field of Systematic Conservation Planning developed rigorous methods for taking such broader context into account, in particular by focusing on the complementarity between sites (such that the value of a given site for conservation depends not only on what species are found at the site, but also on the species that occur elsewhere) and through the concept of irreplaceability (a measure of the extent to which options for conservation are lost if a site is lost) (Pressey et al. 1993, Ferrier et al. 2000, Wilson et al. 2010, Moilanen 2013).

However, these methods are very data-hungry: a decision-maker wishing to apply them to inform decisions at the local scale would first require information on the distribution and abundance of species not only on its site of interest but across a much wider region. Here, we focused on approaches for allowing managers to make the best use of data plausibly.
available to them at the local scale, while still taking into account the extent to which such decisions might affect global options for species’ conservation.

Freely available and covering an expanding diversity of taxa, the range maps collated under the IUCN Red List (IUCN 2014) are an increasingly valuable resource (Rodrigues et al. 2006), but given their coarse nature they are not necessarily in the radar of local managers as potentially useful sources of information. Our results suggest that despite their limitations, global range maps can indeed provide very valuable information to guide local conservation decisions in a global context, even more so if combined with local data plausibly available to managers. We thus support calls for an urgent investments in the expansion of the geographic and taxonomic coverage of these data (Stuart et al. 2010). Refinement of species’ range maps through habitat suitability modelling, combining local field records with environmental layers obtained from remote sensing (Rondinini et al. 2005, 2011), could further improve the relevancy of these maps for local decision-making.

Our results also reinforce the value of high quality local data obtained through fieldwork. We found that by combining reliable occurrence data with range maps it was possible to eliminate the commission errors inherent to the latter and thus improve substantially site estimates of responsibility. We were able to go even further by differentiating among populations in each site and thus identify sites of particular importance for each species. Our results indicate that using expert knowledge to distinguish records of exceptional abundance for each species can be a highly effective approach for doing so, but only in the unlikely scenario of experts’ field experience encompassing the species’ entire range. In contrast, discarding species of very low local abundance seems to be a more practical and effective method for improving the estimates of site responsibility based on plausible local data, but particular attention should be given to species of concern that occur in low densities throughout their ranges.

The approaches we propose here could be easily integrated into tools for supporting local-scale decision making, such as the Integrated Biodiversity Assessment Tool (IBAT 2015), complemented by information on other considerations that affect real-life conservation priorities (e.g. conservation costs, Carwardine et al. 2008; conservation opportunities, Knight et al. 2011; other biodiversity features besides species, such as ecosystem services, Naidoo et al. 2008). Approaches for improving the effectiveness of decision making are all the more urgent in hyper-diverse poorly-studied regions under high human pressure, where conservation-relevant decisions with truly global biodiversity impacts must proceed despite a paucity of biodiversity data.

Acknowledgements – We thank the North American Breeding Bird Survey, BirdLife International and NatureServe for making species’ distribution data available, and to the thousands of people who contributed to their collection. We are grateful to Cyril Berard for his technical support, and to Thomas Brooks and John Pilgrim for comments that have greatly improved the manuscript.

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