Mind the gap: Can downscaling Area of Occupancy overcome sampling gaps when assessing IUCN Red List status?

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Abstract

Aim: The Area of Occupancy (AOO) of a species is often utilized to assess extinction risk for determining IUCN Red List status. However, the recommended raw-counts method of summing occupied grid cells likely reflects only sampling effort, as the majority of species have not been sampled across their entire range at the fine grains required by IUCN. More accurate measurements can be generated at coarser grains (so-called atlas data) as false absences are reduced. If we fit the occupancy-area relationship to these data, we can extrapolate the relationship down to estimate occupancy at finer grains. Numerous models have been proposed to carry out such occupancy downscaling, but have only been tested on a limited range of species.

Methods: We test the ability of downscaling models to recover fine grain AOO against the raw-counts method for 28,900 virtual species with a wide range of prevalence and aggregation characteristics, subsampled to reflect common spatial biases in sampling effort. We address several questions for ensuring accurate downscaling: How to generate accurate atlas data? How far can we accurately extrapolate the occupancy-area relationship given perfect data? Can occupancy downscaling overcome false absences at fine grain sizes? And how does sampling bias and coverage affect accuracy?

Results: Downscaling was more accurate than the raw-counts method in all scenarios except where sampling coverage was very high and/or the sampling bias was positively related to the species distribution. However, if atlas data contained many false absences, then even downscaling under-estimated actual occupancy.

Main conclusions: Occupancy downscaling has the potential to be a useful tool for estimating AOO for IUCN Red List assessments, especially when sampling coverage is low and the currently recommended method is ineffective. However, its application should be tailored to the species' characteristics, as well as the sampling coverage and bias of the species' records.

KEYWORDS

AOO, area of occupancy, conservation, IUCN Red List, OAR, occupancy downscaling, occupancy-area relationship, red listing
1 | INTRODUCTION

The geographic range size of a species is an important characteristic describing a species' rarity, as it is correlated with species' local and total abundances (Gaston, 1991; Gaston & Lawton, 1990) which require more detailed information to estimate. Range size can be quantified at two extremes (Gaston, 1994); the Extent of Occurrence (EOO) is the geographic range that encompasses all occurrences of a species, and the Area of Occupancy (AOO) is the total area within that range that is actually occupied. The measurement of either requires only the spatial coordinates of readily available species record data and forms the basis of one of the criteria used to assess extinction risk for the IUCN Red List (Criterion B, IUCN, 2001, 2017). For example, the recommended method of calculating AOO is simply to overlay a grid over all known records, and sum the area of occupied cells (hereafter, the ‘raw-counts’ method). As a result, the proportion of species assessed as threatened through the estimate of their AOO varies between 37% (amphibians) and 97% (gymnosperms), depending upon the taxonomic group (Gaston & Fuller, 2009).

The issue, however, is that AOO is intrinsically scale-dependent (Kunin, 1998): a species will be seen to occupy different amounts of area if grids of different spatial “grain” are used. Therefore, a species does not have a single AOO value, but rather AOO is a function of grain size (Figure 1), the scale-area or occupancy-area relationship (OAR, He & Condit, 2007), the shape of which is dependent upon the characteristics of the species’ distribution, such as the degree of clumpiness and prevalence (Kunin, Hartley, & Lennon, 2000). The coarser the grain, the larger the measurement of AOO and thus the less threatened the status of that species will appear to be. The finer the grain size used, the closer the correlation with total abundance, so that if a grain size is set to cover a single individual then AOO will eventually equal population size (Kunin, 1998).

The IUCN guidelines require a grain size of 2 × 2 km (IUCN, 2017) and certainly no larger than 3.16 × 3.16 km, as a single occupied grid cell larger than this would give an AOO beyond the threshold for

**FIGURE 1** Example of a hypothetical species distribution where only 50% of true occupancies are sampled at the finest grain, leading to false absences in unsampled areas (red cells) and observed presences (black cells). As grain size increases, the proportion of false absences decreases (bottom right, red) and our estimate of occupancy (bottom left, black) approaches the true area of occupancy (bottom left, blue); however, the lower the proportion of cell area that each occupied cell is actually occupied by the species at the fine scale (second row; bottom right, black). The grain size where the number of false absences approaches zero is a reliable atlas scale (grey line). Models can be fit to the relationship at the atlas scale and above and then be extrapolated back down to fine grains (dashed line).
potential classification as Critically Endangered (10 km$^2$). Although the IUCN Red List Guidelines used to permit use of a different scale ‘dependent on the taxon’, others have suggested that grain size should be based upon the spread of points (Willis, Moat, & Paton, 2003), and in fact, a great many assessments of AOO used grain sizes much larger than the IUCN suggestion (Gaston & Fuller, 2009) as biodiversity atlases are typically compiled at 10 × 10 km or larger.

Regardless of the grain size selected, there are several other challenges potentially preventing the accurate assessment of AOO. The first is insufficient sampling coverage. The finer the grain size used, the greater the sampling effort required to identify all occupied cells for accurate measurement, but the vast majority of species do not have sample data across their full ranges at a grain size of 2 × 2 km. Not only are omission errors (false absences) important in assigning a species’ conservation status (Visconti et al., 2013), with the magnitude of error varying between species of different range sizes (Gaston, 1996), but the sheer volume of records required to attain lower threat categories is prohibitive for the majority of species—put simply, the value of AOO assigned for poorly recorded species may not attain lower threat categories is prohibitive for the majority of species.

The second, and perhaps more pressing, challenge is that the certainty of absences is increased (Figure 1; bottom right, red). Whereas only a single record is required to confirm a presence within a cell (although there are still possibilities of false presences through misidentification, incorrect spatial coordinates or local extinctions since sampling), it is much more difficult to confirm a species’ absence (Kéry, 2002). Therefore at small grain sizes many false absences are likely, but these are reduced as grain size increases (Graham & Hijmans, 2006). This principle lies behind biodiversity atlas data: by collating information over long time spans at large-grain sizes, accurate representations of a species’ distribution can be generated (Gibbons, Donald, Bauer, Fornsar, & Dawson, 2007). Therefore, as grain size increases and the proportion of false absences decreases, the measurement of AOO moves from being largely reflective of sampling effort to being a more accurate estimate of that species’ AOO at the scale of measurement, where accuracy is measured as the false omission rate (the number of false absences divided by the sum of false and true absences) in this case.

Although accuracy, using this measure, increases with grain size, there is, however, a reduction in within-cell precision for coarse-grain atlases; for an occupied coarse-grain cell, the cell area will be composed of a larger proportion of area that it is not occupied at a finer grain (Figure 1; bottom right, black) and are therefore less information-rich than an accurate fine grain atlas. As accuracy and within-cell precision have opposing scaling relationships, they must therefore be compromised between each other.

Published atlases of biodiversity have a distinguished tradition and have proliferated over the last decade (Powney & Isaac, 2015) as larger volumes of biodiversity data have become publicly available. However, even for extremely well-recorded taxa in highly sampled regions, atlases are typically collated at 10–20 km

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**1.2 | A solution to sampling gaps: atlas data**

A potential solution where sampling gaps are large is to increase the spatial grain at which data are aggregated. As grain size is increased, the quantity of sampling within each sampled cell grows higher and the number of cells with little-to-no sampling is reduced. In particular, the certainty of absences is increased (Figure 1; bottom right, red). Whereas only a single record is required to confirm a presence within a cell (although there are still possibilities of false presences through misidentification, incorrect spatial coordinates or local extinctions since sampling), it is much more difficult to confirm a species’ absence (Kéry, 2002). Therefore at small grain sizes many false absences are likely, but these are reduced as grain size increases (Graham & Hijmans, 2006). This principle lies behind biodiversity atlas data: by collating information over long time spans at large-grain sizes, accurate representations of a species’ distribution can be generated (Gibbons, Donald, Bauer, Fornsar, & Dawson, 2007). Therefore, as grain size increases and the proportion of false absences decreases, the measurement of AOO moves from being largely reflective of sampling effort to being a more accurate estimate of that species’ AOO at the scale of measurement, where accuracy is measured as the false omission rate (the number of false absences divided by the sum of false and true absences) in this case.

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**1.1 | Spatial sampling biases in data**

If sampling intensity is equally spread across a species’ distribution, then an accurate AOO estimate may be achieved even at relatively fine grain sizes at low efforts (Gaston & Fuller, 2009). Unfortunately, however, even for well-sampled species there are likely to be distinct spatial biases in where data have been collected and therefore in the location of false absences, and furthermore, the patchiness in collection effort is rarely random (Beck, Boller, Erhardt, & Schwanghart, 2014; Isaac & Pocock, 2015).

Spatial biases occur at distinct spatial scales. At global scales, sampling intensity is concentrated in developed countries with high political stability (Hortal, Jiménez-Valverde, Gómez, Lobo, & Baselga, 2008), particular colonization histories and a cultural proclivity to natural history (Stropp et al., 2016). At regional scales, occurrences are often clustered around areas of high accessibility, such as close proximity to roads (Reddy & Dávalos, 2003) or around research centres and universities (the ‘botanist effect’; Moerman & Estabrook, 2006). There is typically a distance-decay in sampling effort away from these well-recorded regions (Ladle & Hortal, 2013). At more local scales, effort is often directed towards good habitat areas the recorder believes a priori to be suitable for the species or sites of high biodiversity, such as reserves or known areas of occurrence of particular rare species (Freitag, Hobson, Biggs, & Jaarsveld, 1998).

Unfortunately, only occurrence records are available for the majority of species, so it is extremely difficult to estimate sampling effort across space for a given species lacking associated absence data (Isaac, Strien, August, Zeeuw, & Roy, 2014). It is therefore difficult to distinguish between a species that is genuinely rare and does only occur in a few locations and one that is simply under-recorded and for which there are large sampling gaps across its distributions.
One solution is so-called occupancy downscaling. First, we generate the OAR at coarse grains using atlas data and fit likely mathematical functions to approximate the relationship, which can then be extrapolated down to estimate occupancy at fine grains. The initial step is therefore to generate accurate atlas data at large-grain sizes; however, currently there is no method for selecting the most suitable grain size. The larger the grain size, the greater the accuracy of cell occupancies and the better that sampling gaps may be overcome by minimizing false absences. This comes with several trade-offs, however. First, the further up the OAR towards large grains that modelling begins, the further it must be extrapolated back down to predict occupancy at fine grain sizes, with potential increases in subsequent prediction error. Second, as grain size increases atlases may reach the scale of saturation—the grain size where all cells are occupied—or the scale of endemism—the grain size where all occurrences occur within a single cell (Hartley & Kunin, 2003). If these are reached, the data for all grain sizes larger than this point must be discarded before modelling, so reducing the number of data points for model fitting, which may reduce prediction accuracy, or worse, leave insufficient data points to fit the downscaling models.

A number of models for downscaling the OAR are available (see Azaele, Cornell, & Kunin, 2012; Barwell, Azaele, Kunin, & Isaac, 2014), ranging from simple Poisson models to more complicated models that incorporate species aggregations through pattern-point processes. The ability of these models to accurately extrapolate AOO to finer grain sizes has been found to be relatively consistent, dependent upon species’ prevalences (Groom et al., 2018). Furthermore, the direction of error is predictable. For example, the power-law model, the IUCN-suggested method for translating the number of samples and their distribution pattern are likely to affect the number of false absences detected in the atlas data and thereby the shape of the sampled OAR. In general, we predict that the closer the sampled OAR curve is to the true curve at the atlas scales, the greater the accuracy of the downscaling estimates at fine scales.

2 | METHODS

Our analyses followed four main steps (Figure 2): (a) generating the virtual species, (b) sampling the virtual species, (c) selecting the appropriate scale to generate the atlas data and finally (d) predicting fine-scale occupancy using downscaling. All simulations were carried out using R 3.4.3 (R Core Team, 2017); occupancy downscaling was carried out using the ‘downscale’ package (Marsh et al., 2018).

2.1 | Generating virtual species

Previous research has shown that the prevalence of a species (the proportion of occupied cells in the landscape) has a large effect on our ability to downscale the OAR (Groom et al., 2018), and it is likely that models will also differ in their ability to recover the OARs generated from species with different degrees of aggregation, as this also affects the shape of the OAR. We distributed species across 512 x 512 cell grids (262,144 cells) using a spherical variogram (sill = 1.5, beta = 1) and explored seventeen prevalence levels between 0.00005 (13 occupied cells) and 0.5 (131,072 occupied cells) and seventeen clumping values (the range parameter) between 2 (highly disaggregated) and 512 (highly aggregated), both distributed evenly in log space.

We created 100 replicates for each prevalence-clumping combination, generating a total of 28,900 virtual species spanning the full parameter space of realistic species distributions in the given extent. A description and R script for creating the virtual species is available in the Supporting information (examples are presented in Figure 3).

2.2 | Sampling virtual species

The number of samples and their distribution pattern are likely to affect the number of false absences detected in the atlas data and thereby the shape of the sampled OAR. In general, we predict that the closer the sampled OAR curve is to the true curve at the atlas scales, the greater the accuracy of the downscaling estimates at fine scales.

For each of the 28,900 species, we explored various combinations of sampling coverage and sampling distribution. For the sampling coverage, we drew between 0.005 (1,311 cells) and 0.232 (60,818 cells) proportions of cells in six equally spaced increments in log-space. We used three different protocols to distribute the samples in the landscape (Figure 3) but ensured that at least one presence was recovered (see details in the Supporting information; Figure S1.1).
1. Random sampling—every cell has an equal probability of being sampled until the designated sampling coverage has been reached.

2. Aggregated neutral sampling bias—a spatially biased distribution of samples, where the bias is neutral in relation to the species distribution. This represents the bias that occurs where sampling effort is independent of whether the data recorder is expecting to encounter the species. This bias occurs frequently where certain regions or countries are over- or under-sampled due to cultural and economic reasons, or there is increased sampling intensity around easily accessible locations. A probability of sampling surface was created by splitting the grid into 64 ‘regions’ of 64 × 64 cells. Each region was assigned a probability of sampling with a large degree of spatial autocorrelation that approximates patterns found at continental scales (Figure S1.2). When samples are drawn from the sampling surface, sampling coverage is therefore concentrated in a few closely associated regions.

3. Aggregated positive sampling bias—a bias positively correlated with the species distribution such that presences are more likely to be sampled than absences, but discovering new populations may take some time as effort is focussed around known locations. This represents scenarios of increased sampling effort in suitable habitat where the sampler expects to encounter the species due to previous knowledge, but is unlikely to sample previously unsampled areas. Samples were drawn using the probability map created during the virtual species creation process. In order to further distinguish high-suitability areas, probability values were first raised to the power of ten. Samples are then drawn from this probability surface (\( \text{prob} \)). Once a presence has been detected, we calculate a new probability surface as a function of an exponential decay curve with a mean of 25 cells (halving distance = ~17 cells), so that the probability of sampling of cell \( i \) is calculated as \( \text{prob}_i \times e^{-\left(\frac{d_i}{25}\right)} \), where \( d_i \) is the centre-to-centre distance between cell \( i \) and the occupied cell. Further samples are drawn from this probability surface until the next presence is detected and the process is repeated until sufficient samples have been accumulated (Figure S1.3).

2.3 | Selecting the atlas scale

To examine the impact of grain size on atlas accuracy, we first generated the atlases and OARs for each of the 100 replicates for each species (prevalence-clumping combination) using the complete data and each sampling coverage and protocol combination. Each OAR was created by aggregating the sampled cells in arrays of 2 × 2, 4 × 4, 8 × 8, 16 × 16, 32 × 32, 64 × 64, 128 × 128 and 256 × 256 cells. This resulted in 41,616,000 atlases for testing.

We explored the trade-off between atlas accuracy and model performance. Atlas accuracy was evaluated as the mean proportion of false absences across the 100 replicates at each scale for each sampling protocol and coverage combination. We expect that as grain size increases the proportion of absences will decrease and...
that larger grain sizes will be required when sampling coverage is low or when sampling is independent of the species’ distributions.

A larger grain size increases atlas accuracy (but reduces within-cell precision), but may prevent model fitting if the scales of saturation (the grain size where all cells are occupied) or endemism (the grain size where all occurrences occur within a single cell) is reached. We calculated the median scale of saturation and endemism at each grain size for each OAR over the 100 species replicates. We also examined three aspects that may then impact the accuracy of occupancy downscaling. First, extrapolating from a finer grain size means that a larger number of coarser grain data points are available for model fitting, which should therefore result in more accurate estimates of the OAR. Second, the finer the grain size, the fewer steps down between model fit and prediction. However, and finally, the finer the grain size, the greater the likelihood that atlas data will be inaccurate as we increase the likelihood of false absences. We explored this by predicting AOO at the 1 × 1 grain size using atlas data generated at grains sizes of 4 × 4, 8 × 8, 16 × 16 and 32 × 32. Models were also fitted with three to six fitting scales, depending on atlas scale (see following section on modelling procedure and evaluation). As species were distributed over a square area, we did not explore the effects of standardizing the extent or the position of the grid origin but these methods can also be important (Groom et al., 2018; Marsh et al., 2018).

Upon inspection of the results (see Figures S2.1-S2.7), the most appropriate atlas data were dependent upon the species prevalence and clumping, as well as on the sampling bias and effort. For example, rare species require atlas data created at very large-grain sizes, but atlas data at these scales have already reached saturation in common species. It was therefore decided that for further analyses, the atlas scale would be run-specific; it would be created at the largest scale that still allowed for modelling before the scale of saturation or endemism was reached.
2.4 | Predicting fine-scale occupancy

We fitted the downscaling models to the OAR generated from the atlas scale and larger grain sizes and extrapolated them to predict occupancy back at the 1 × 1 grain size. As successfully fitting some of the models can be difficult or require long processing times, in this study we used an ensemble approach, averaging the estimates in log-space from the Poisson, power law, Nachman, exponential and negative binomial downscaling models. These models were selected as they provide robust estimates that are rapidly computed while still maintaining good accuracy (Groom et al., 2018).

We first examined individual models’ ability to accurately predict the OAR given complete atlas data across all species (‘downscaled prevalence, full data’ in Figure 2) and then repeated the analyses for the three subsampling methods and six sampling coverages (‘downscaled prevalence, sampled data’). We also calculated occupancy generated through the raw-counts method recommended by IUCN, by simply summing the number of occupied cells after subsampling at 1 × 1 grain size (‘raw-counts, sampled data’). Accuracy was calculated as the difference between the predicted occupancy and true occupancy (‘true prevalence, full data’) in log-space. Negative values indicate the method under-predicted occupancy and positive values indicate the method over-predicted occupancy. For each prevalence-clumping combination, we calculated the mean accuracy across all 100 species replicates. For the downscaled predictions, if fewer than five repeats could be evaluated then accuracy was given as non-assessed. We then compared downscaling models and raw-counts through the difference in absolute values of accuracy.

3 | RESULTS

3.1 | Selecting the atlas scale

For a given level of sampling effort, the proportion of false absences increased as the virtual species became either rarer or more clumped (Figures S2.1–S2.3). For the rarest species, some false absences remained even at the highest sampling effort (0.232 of cells sampled) at the largest grains examined under random or neutral sampling biases (Figures S2.1–S2.2). If the sampling bias was positively correlated with the species distribution, then false absences could be removed at fine grains with much lower sampling effort (Figure S2.3).

If the scale of saturation or endemism occurs at fine grains, then it can prevent the fitting of downscaling models. All but very scarce (low prevalence) species reached the scale of saturation at some scale (Figures S2.4–S2.6) but the scale was only small enough to cause modelling issues for common, dispersed species with higher sampling effort (red in Figure 4). A greater problem was reaching the scale of endemism (blue in Figure 4): scarce species reached scales of endemism at even the smallest scales (Figures S2.4–S2.6). As we used a variable cell size, we were still able to model all but the lowest prevalence species at low sampling efforts, but atlases may be as fine as 2 × 2 cells in these cases (Figure S2.7). Overall, reducing prevalence
and sampling effort increased the chances of reaching the scale of endemicism. Under a positive sampling bias, the potential for reaching the scale of saturation was increased, but reduced for the scale of endemicism, whereas the opposite occurred under a neutral sampling bias.

When examining the OARs generated using complete data, most species’ OARs were linear in log-log space before quickly levelling off if full occupancy was reached (Figure S2.8), which reflects the variogram used to generate them. Some species of medium prevalence but high aggregation may actually show a slightly concave OAR, as well as higher variation between species. Examining the OARs after subsampling reveals that there is much more variance between the subsampled OARs as species aggregation increases (Figures S2.9–S2.11). Where species prevalence and aggregation are low, the subsampled OAR does not approach the true OAR even at the highest sampling coverages.

### 3.2 | Predicting fine-scale occupancy

When predicting fine grain AOO through downscaling, there was considerable variation between the predictions of different downscaling models, even when using complete atlas data (Figures S2.12–S2.16). The Poisson, power law and negative binomial downscaling models tended to over-predict occupancy, whereas the Nachman and logistic models generally under-predict occupancy. Where species were rare and aggregated, there was much greater within-model variation (Figure S2.16).

For fitting the downscaling models, the effect of selecting a particular atlas scale was greater than the effect of having a larger number of scales (Figure 5). The coarser the atlas scale, the more the models were able to overcome sampling gaps, but with increased variation in predicted occupancies. Increasing the number of scales for fitting the models also increased the accuracy of predictions even though the atlas data were identical in these cases.

Occupancy downscaling provided more accurate predictions than did the raw-counts method in the majority of cases (Figure 6), but the variance was also much higher, suggesting that there is a large amount of error associated with the downscaling models themselves beyond the variation present in the set of species examined. However, downscaling was still mostly unable to fully overcome the sampling gaps at low sampling coverages. The raw-counts method was instead highly correlated with sampling coverage, although it could provide high accuracy where there are high sampling coverage and a positive sampling bias associated with the species distribution (Figure S2.17).

Performance was not consistent across species (Figure 7). Downscaling produced a rather unusual ‘yin-and-yang’ pattern of accuracy where downscaling tended to under-predict for clumped, medium-high prevalence species but over-predict for dispersed low-medium prevalence species, when sampling coverage was high (upper row in Figure 7). At lower sampling coverages (lower rows), downscaling tended to under-predict at all but very low prevalence
values. We were often unable to provide predictions for extremely low prevalence species under random or regional sampling with low effort due to the scale of endemism preventing model fitting.

Occupancies predicted through downscaling were substantially closer to true occupancy than the raw‐counts method unless sampling coverage was very high, or at lower coverages if there was a positive sampling bias and species were highly clumped (Figure 8).

**4 | DISCUSSION**

The area occupied by a species is one of the most widely applied estimates of a species’ conservation status (Gaston & Fuller, 2009). However, the recommended method of measurement by summing the area of occupied cells of a 2 × 2 km grid is inaccurate where there are large sampling gaps. A more accurate estimate can be made if occurrences are aggregated across larger spatial (and temporal) scales to create atlas scale data, reducing the number of false absences (Figures S2.1–S2.3). We can then fit likely functions to the OAR at large‐grain sizes, where accuracy is high, and downscale them to predict occupancy at the fine grain size required by IUCN.

There are numerous challenges to this approach, listed in the introduction, that have been explored here. These questions have been impossible to address using real species data as so few species have been mapped at sufficiently high resolution and accuracy across large extents, but can be approached using virtual species as demonstrated here.
4.1 | Selecting the appropriate atlas scale

Published biodiversity atlases use grain sizes that are understandably standardized across species and highly correlated with atlas extent (Gibbons et al., 2007). In parallel, the open data revolution allows the creation of biodiversity ‘atlases’ for specific objectives directly from distribution data stored in online repositories, allowing greater freedom in the choice of grain and extent. We found that for an atlas to be relatively accurate (i.e. have a limited number of false absences), not only is the sampling coverage and bias critical to selecting grain size, but this scale should be specific to the species’ prevalence and clumping (Figures S2.1–S2.3). For example, accurate atlas data can be generated at fine grains for common, clumped species but if species are rare and dispersed atlas data may be inaccurate even at large-grain sizes and high sampling coverage.

Issues are further complicated by a species reaching the scale of either saturation or endemism before enough scales are present to fit downscaling models. In this study, we used the largest grain size that still allowed for modelling, but in our example, this led to very small grain sizes for species with very low prevalence. During Red List assessments, if a species reaches the scale of endemism at a fine grain size where sampling coverage is believed to be too low to generate accurate atlas data, this may be an appropriate reason to assign the species as Data Deficient with regard to AOO.

4.2 | Predicting fine-scale occupancy

In the majority of cases for our virtual species, the estimates from ensemble downscaling were more accurate than were the raw-counts method but downscaling is still likely to underestimate AOO to some extent (Figures 6 and 8), unless models that systematically over-predict AOO given perfect data are utilized (Figure S2.16). Prevalence is an important indicator of downscaling accuracy (Groom et al., 2018), but we also found that degree of aggregation will impact the ability to accurately recover the OAR. For example, even given perfect data, downscaling over-predicts the AOO of scarce, dispersed species but under-predicts the AOO of abundant, clumped species (Figures S2.9–S2.11).

Underestimation of AOO was greater the lower the sampling coverage (Figure 7) but to a large extent this is due to atlas data at even large-grain sizes containing a large proportion of false absences (Figures S2.4–S2.6), as underestimation was reduced when a positive sampling bias was applied. Every effort should therefore be made to ensure atlas data are accurate before downscaling is attempted. The increased accuracy through downscaling does come, however, with increased variance (Figure 6).
4.3 | Guidelines on downscaling the OAR

The most appropriate approach to estimating AOO should, as far as possible, vary depending upon the characteristics of the species' distribution, the expected sampling coverage and any suspected spatial sampling bias (Table 1). More generally, we propose the following recommendations:

- Where sampling coverage is low, downscaling provides a better estimate of AOO than the raw-counts method does, but it is still likely to be an underestimate of true AOO (Figures 6 and 8).
- Where sampling coverage is very high, and particularly where any sampling bias is likely to be positive to the species distribution, it may be better to use the raw-counts method (Figure 8).
- An accurate estimate of AOO is more likely using atlas data at larger grain sizes, but the estimates have greater uncertainty from the downscaling predictions (we must extrapolate further back; Figure 5).
- Where possible downscaling accuracy will be increased by using more scales for fitting, but this should not be done at the expense of using a finer-grained atlas (Figure 5).
- Where the scale of endemism is reached at fine grain sizes before accurate atlas data can be generated, it may be appropriate to assign the species as Data Deficient with regard to Criterion B2.
- Where no other information on the appropriate atlas grain size is available, atlas data should be generated at the largest scale that still allows for modelling before the scale of saturation or endemism is reached, providing this will not result in very fine grain sizes.
- Care should be taken when using downscaling to assess trends over time, as occupancy changes will be manifested over longer time periods at coarse grains (regional extinctions/colonizations) than at fine grains (local extinctions/colonizations; Hartley & Kunin, 2003).

4.4 | Potential improvements and future directions

Where the downscaling approach could provide additional information over the raw-counts method is to give an estimate of uncertainty or error around the measurement of AOO, which is not provided by current methods but could be critical when determining if there are trends in changing AOO over time (Akçakaya et al., 2000). There are two errors associated with downscaling.

The first is the error within the downscaling models, which are relatively predictable at least in the direction of error (Figure S2.16; Groom et al., 2018). Here, we also show that variation between models is dependent upon the species’ prevalence and clumping, as some models appear to struggle to recover the shape of certain
TABLE 1  Guidelines for how to approach downscaling the OAR depending upon the prevalence and clumping of the species and the spatial coverage of sampling

| Species type | Scarcity | Rare | Medium | Common |
|--------------|----------|------|--------|--------|
| Complete distribution data | Typical OAR (black line) | Clumped (bottom right) | Dispersed (bottom left) | Clumped (top right) | Dispersed (top left) |
| Downscaled accuracy (red line) | Over-predicts | Accurate | Highly accurate | Unfeasible | Under-predicts |

| Sampling | Sampling bias | Random/neutral | Positive | Random/neutral | Positive | Random/neutral | Positive | Random/neutral | Positive |
|----------|--------------|----------------|----------|----------------|----------|----------------|----------|----------------|----------|
| Generating good atlas data after subsampling | High | Low | Extremely high/complete | High | Medium | Medium | Low | Medium | Low |
| Scales of endemism reached? | Even at medium sampling coverage | At low sampling coverage | Even at high sampling coverage | Even at medium sampling coverage | Rarely | Rarely | Never | Never | Never | Never |
| Scales of saturation reached? | Never | Never | Rarely | Rarely | Rarely | Rarely | Rarely | Rarely | At medium sampling coverage | At medium sampling coverage |

| Downscaled accuracy | Low coverage | Infeasible | Accurate | Infeasible | Accurate | Under-predicts | Under-predicts | Under-predicts | Under-predicts | Under-predicts | Under-predicts |
|---------------------|--------------|------------|----------|------------|----------|----------------|----------------|----------------|----------------|----------------|----------------|
|                     | Medium coverage | Accurate | Accurate | Slightly over-predicts | Under-predicts | Slightly under-predicts | Under-predicts | Under-predicts | Accurate | Accurate |
|                     | High coverage | Accurate | Slightly over-predicts | Over-predicts | Accurate | Accurate | Under-predicts | Under-predicts | Accurate | Accurate |

| Downscaled predictions versus raw counts | Low coverage | Downscaling infeasible | Downscaling infeasible | Downscaling | Downscaling | Similar | Downscaling | Downscaling | Downscaling |
|-----------------------------------------|--------------|------------------------|------------------------|------------|------------|---------|------------|------------|------------|
| Medium coverage                         | Downscaling Raw counts | Downscaling Raw counts | Downscaling Raw counts | Downscaling | Downscaling | Downscaling |
| High coverage                           | Downscaling Raw counts | Similar | Raw counts | Downscaling | Similar | Raw counts | Downscaling | Similar |
OARs. Error will also increase with the distance of extrapolation (the larger the atlas scale, Figure 5). To some extent, by exploring a range of possible OARs in simulations such as these, we can therefore predict the uncertainty of our estimates given the species’ characteristics in the atlas data and could weight models during the averaging of the ensemble process accordingly.

The second uncertainty is associated with generating coarse-grain atlas data without sampling gaps and is more difficult to estimate as generally we cannot distinguish true absences from areas that have not been sampled. Published atlases ensure accuracy through the accumulation of data over long time periods, but they are generally confined only to well-known taxa in well-recorded regions. Unfortunately, for less well-recorded taxa, particularly in the tropics, the majority of species are represented by fewer than 30 records (Brummitt, Bachman, Aletrari, et al. 2015a; Brummitt, Bachman, Griffiths-Lee, et al., 2015b).

The issue is exacerbated in that sampling coverage will also be uneven and likely to be spatially biased. It is extremely difficult to estimate sampling coverage or bias from the presence-only data generally available from biological records, although some models are available that utilize information from the records of similar taxa to ascertain this (Isaac et al., 2014). We urge those that collect biological records data to also collect and publish absences where they are certain, as absence data are just as valuable as presence data in nearly all applications for predicting species distributions.

Finally, downscaling methods themselves could also be developed further. For example, models could account for potential false absences in the data if we know sampling to be low in particular regions, randomly assigning a presence or absence to uncertain atlas cells. Repeated many times this could produce a distribution of predicted occupancies. Additionally, although most downscaling models account for saturation (occupancy cannot exceed one), none account for the ‘slope of endemism’ where the maximum slope possible is when only one of four cells at grain size n is occupied for each cell occupied at grain size n + 1. Therefore, some models produce OARs that we know to be impossible. In such cases, the log-log slopes could be set at 0.25.

5 | CONCLUSION

Downscaling occupancy from coarse-grain atlas data is potentially a valuable method for estimating AOO in IUCN Red List assessments. In previous studies and here we have:

1. Created a new R package which makes ten published downscaling models accessible (Marsh et al., 2018).
2. Shown that an ensemble approach can accurately predict the occupancy of a large number of real species (Groom et al., 2018).
3. Shown that for many virtual species, differing in their prevalence and clumping, ensemble downscaling can fill in information gaps resulting from low sampling coverage and spatial biases better than the currently advocated method of using raw-counts (Figure 6).
4. Provided information on the limitation of downscaling and guidelines on when it should and should not be used (Table 1).

Given the increased availability of open-access biodiversity data that allows considerable freedom in creating bespoke atlas data, the potential to automate the fitting of downscaling models, and their ability to provide a more accurate AOO estimate at the recommended scale of 2 × 2 km, we hope that downscaling can usefully contribute to the IUCN Red Listing toolbox. We note that downscaling is only one of the tools suggested in the literature to assess AOO (Marsh et al., in review). Which of these methods is the most appropriate for various species under various sampling scenarios remains under-explored. Repeating a similar analysis with virtual species for a wide range of AOO methods may reveal further methods that are complimentary to one another, each more suitable for different species and data characteristics, and lead to more holistic guidelines.

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Biosketch

Charlie Marsh is an ecologist interested in how biodiversity and ecosystem function are affected by anthropogenic disturbance, particularly fragmentation and habitat modification. Recently, the focus has been on the spatial scaling of ecological processes, and decoupling the effects of scales of observation from interpretations of spatial patterns of diversity. This used to involve lots of exciting running around rain forests, but sadly mainly involves computer simulations these days.

Author contributions: All authors conceived the idea and designed the simulations. C.J.M. and Y.G. wrote the scripts and carried out the analyses. C.J.M. led the writing with input from all authors.

Supporting Information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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