Historical drivers of extinction risk: using past evidence to direct future monitoring

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Global commitments to halt biodiversity decline mean that it is essential to monitor species’ extinction risk. However, the work required to assess extinction risk is intensive. We demonstrate an alternative approach to monitoring extinction risk, based on the response of species to external conditions. Using retrospective International Union for Conservation of Nature Red List assessments, we classify transitions in the extinction risk of 497 mammalian carnivores and ungulates between 1975 and 2013. Species that moved to lower Red List categories, or remained Least Concern, were classified as ‘lower risk’; species that stayed in a threatened category, or moved to a higher category of risk, were classified as ‘higher risk’. Twenty-four predictor variables were used to predict transitions, including intrinsic traits (species biology) and external conditions (human pressure, distribution state and conservation interventions). The model correctly classified up to 90% of all transitions and revealed complex interactions between variables, such as protected areas (PAs) versus human impact. The most important predictors were: past extinction risk, PA extent, geographical range size, body size, taxonomic family and human impact. Our results suggest that monitoring a targeted set of metrics would efficiently identify species facing a higher risk, and could guide the allocation of resources between monitoring species’ extinction risk and monitoring external conditions.

1. Introduction

Despite a growing international commitment to conservation, the current biodiversity crisis is characterized by increasing human pressures and continuing decline in the status of many species and habitats [1]. Reversing this trend has become the aim of one of the ambitious Aichi biodiversity targets proposed for 2020 [2]: reducing the extinction risk of known threatened species. If this target is achieved, it will in turn have a positive synergistic effect on other targets (such as the protection of forests and the maintenance of carbon stocks [3]). Progress towards meeting this global biodiversity target relies on monitoring the extinction risk of species. Over recent decades, the International Union for Conservation of Nature (IUCN) has assessed the extinction risk of more than 70,000 species of plants, vertebrates and invertebrates on the Red List of threatened species [4]. The classification of threatened species is clearly an effective conservation tool [5], with the IUCN Red List underpinning both international policy processes [2] and research aimed at improving conservation responses [6].

However, classifying and monitoring species’ extinction risk requires intensive expert effort and considerable financial resources, which is unsustainable without change in either the strategy for assessment or funding [7]. Approaches such as sampling of taxa can be used to provide short cuts, but it remains a substantial task [8]. Overall statistics from the IUCN Red List are used for measuring the status and trends of biodiversity [1,6] and for designing global-scale...
strategies for conservation interventions [9]. In addition, species-specific assessments inform direct actions to address particular threats at specific times and sites, requiring a comprehensive species-level approach [10].

The extinction risk of species, assessed using the IUCN Red List criteria [11], is a consequence of their biological traits, past and current environmental conditions, direct human pressures and the interactions between these factors [12,13]. Environmental changes and pressures on species are increasing in intensity and are the main cause of current increases in extinction rates. Extinction risk modelling has been used to better represent and quantify these external drivers, which can change and intensify over a short time frame [14,15]. Biological traits, by contrast, change very slowly, and determine the way in which species respond to external pressures [13]. Historical information on species’ extinction risk, and the way in which risk has changed in response to known pressures, could therefore be a good way to predict future biodiversity trends, particularly when the pressures can be effectively monitored or forecast.

Di Marco et al. undertook a retrospective assessment of the extinction risk of the world’s carnivores and ungulates between 1975 and 2008 [16] by applying the current IUCN Red List criteria [11] to historical information. Studying past trends in extinction risk can indicate the circumstances under which conservation policies and strategies are or are not successful. Retrospective assessments can also guide the interpretation of future scenarios of emerging threats, for example, inferring the likely consequences of land use change or climate change [17]. Therefore, one approach to reducing the logistical and financial constraints of constant extinction risk monitoring could be to use well-validated models, based on past trends, to predict the effect of changing external pressures on future extinction risk [18,19].

In many cases, Red List categories remain stable over long periods of time, especially for the large number of species listed as Least Concern (LC) [11]. The most useful information therefore concerns those species whose extinction risk is likely to escalate. We use historical records to develop and refine models of change in extinction risk, to identify those species for which high-risk combinations of biological vulnerability and extrinsic threats occur. We use current [4] and historical [16] information on Red List categories for 497 species of mammalian carnivores and ungulates in the period 1975–2013, to represent ‘transitions’ in species’ extinction risk (figure 1). We classified species in two groups: ‘lower risk’ transitions, for those species not facing a significant increase in their extinction risk over time, and ‘higher risk’ transitions, for those species facing a significant increase in their extinction risk over time (see Material and methods; electronic supplementary material, table S1). This approach is not analogous to measuring ordinal transitions between Red List categories (e.g. [20]), as we deliberately highlight species that will be of greatest concern to conservation, namely those that remain at a relatively high risk of extinction over time, and those that move from lower to higher risk categories.

We acknowledge that our study species are not a representative subset of all mammals, let alone life on the Earth. For example, carnivores and ungulates are generally characterized by longer generation times [21] and higher risk of extinction [4] relative to other mammals. Nonetheless, the high conservation attention devoted to these groups makes a perfect case for testing our analytical approach.

We predicted higher and lower extinction risk transitions for species, using a comprehensive set of variables, which represent the conditions faced by the species during the study period. Our analyses therefore mimic a hypothetical situation in which relevant biological datasets and reliable forecast environmental and conservation metrics were available in the 1970s. This would have enabled conservation planners to predict which species would be in a higher or lower risk condition over the next 40 years.

Figure 1. Transition of species’ extinction risk categories in the period 1975–2013. The plot reports the number of species (carnivores and ungulates) in each Red List category for each time period. Circle size is proportional to the number of species, while arrows represent the proportion of species moving from an initial category to a final category (arrows’ width scales with the proportion of species in the original category). Data were obtained from [4,16]. CR, Critically Endangered; EN, Endangered; EW, Extinct in the Wild; LC, Least Concern; NT, Near Threatened; VU, Vulnerable. (Online version in colour.)
2. Material and methods

(a) Obtaining extinction risk transitions

We included all species of carnivores (Carnivora), ungulates (Perissodactyla and terrestrial Cetartiodactyla) and Proboscidea (discussed below together with ungulates) currently assessed in the IUCN Red List [4]. We excluded those species identified as being historically (less than 1970) Extinct or Data Deficient. We also excluded the Saudi gazelle (Gazella saudia), declared extinct in the 1980s, as we had no detailed information available for its life-history traits (apart from body mass) or spatial distribution.

We considered 497 species in our analyses, representing 93% of all extant species in the study groups.

We compared the most recent species’ extinction risk categories assessed in the IUCN Red List [11] with a retrospective assessment for 1975 [16]. We calculated an extinction risk transition value for each species between the two time periods in terms of the number of Red List categories changed (figure 1). A negative transition (less than 0) characterized species that moved towards a lower category of risk, a stable transition (equal to 0) characterized species that maintained the same Red List status, and a positive transition (more than 0) characterized species that moved towards a higher category of risk.

We considered changes in species’ extinction risk over a roughly 40-year period (1975–2013). This is a reasonable reference period for species in our study groups, as it corresponds to more than 10 generations for small carnivores and approximately two generations for large-bodied species such as elephants and rhinos [21].

(b) Classifying extinction risk transitions

Because we were most interested in species that had fared unusually badly compared with those following an average trend over the study period, we identified species with a transition value significantly higher than random, when compared with other species within the same original extinction risk category. To do this, we (i) randomly re-assigned the observed transitions across all species within each original Red List category, (ii) compared the observed transitions with the randomly assigned transitions, and (iii) repeated the previous steps 10 000 times.

As an example, the transition of a species moving from LC (in 1975) to Near Threatened (in 2013) was higher than a transition randomly selected from other originally LC species in approximately 85% of the comparisons. Species with a transition value higher than random in less than or equal to 5% of the comparisons were included in the ‘lower risk’ group. Species with a transition value higher than random in more than 5% of the comparisons were included in the ‘higher risk’ group. Importantly, a species retaining the same category over the time period (net change equal to 0) may have a transition value higher than random if several other species in the same original category had moved to lower categories of risk (net change less than 0).

The randomization resulted in two groups containing species characterized by different extinction risk trajectories (electronic supplementary material, table S1). The ‘lower risk’ group included species that were LC throughout the study period, together with species that underwent a change from any category to a lower category of risk. The ‘higher risk’ group included all species that underwent a change from any category to a higher category of risk, together with species that were originally threatened or near threatened and retained their category. This classification reflects the intrinsic properties of the Red List criteria, in particular the fact that remaining within the same Red List category has different implications depending upon the category. For example, a species classified as LC throughout the time period does not face any significant decline over time. By contrast, a species classified as Vulnerable (VU) throughout the time period faces a strong continuing decline in abundance (more than or equal to 30%) and/or remains at a very low population size. The species in the latter case therefore has a much higher probability of extinction (more than or equal to 10% in 100 years) [11].

(c) Modelling the drivers of extinction risk transition

We modelled the probability that a species is included in the higher risk or in the lower risk group, based on its original extinction risk category and the conditions in place over the study period. Extinction risk has been shown previously to be attributable to a combination of intrinsic and extrinsic factors [13]. Following recent work [22], our model included three classes of external predictor variables and one class of intrinsic (biological) predictors (see table 1 for a complete list and description). The external variables are intended to reflect conditions faced by the species during the study period. We measured: (i) distribution state variables, such as species’ range size (measured in orders of magnitude); (ii) human pressure variables, such as the human influence index [37]; and (iii) conservation response variables, measured as the proportional coverage and absolute extent of protected area (PA) within species ranges (again the extent was measured as an order of magnitude). The fourth group of predictor variables reflects species life-history traits (i.e. species biology) including physical characteristics (e.g. body size), reproductive timing (e.g. weaning age) and reproductive output (e.g. weight at birth) [32]. We used an existing dataset [33], in which multiple imputation techniques had been used to fill gaps in life-history data [38].

Obtaining measures of external predictor variables corresponding to exactly the same years as the assessment period was not always possible. Nonetheless, most of these data refer to the second half of the study period (i.e. more than or equal to 1990s), where the highest decline in species status was observed [16]. We assumed that changes that occurred within a relevant part of the 40-year study period (especially the second half of the period) would serve as a valid approximation for the entire period. In addition, this reduces the risk of collinearity between predictor variables (including levels of habitat loss and other proxies of human pressure) and original threat status (derived from retrospective assessments of extinction risk in the 1960s–1970s). We decided not to include variables that could not reasonably be used as predictors of future extinction risk change. For example, measures related to species distribution such as biogeographic realm—while probably acting as a proxy for regional pressure levels—could not reasonably be used by conservation planners to predict future changes in extinction risk of species.

We used random forest (RF) modelling to estimate the probability that a species was included in the higher risk or in the lower risk group. RF modelling is a powerful tool for ecological analysis [39], and it has been successfully used to model extinction risk in mammals [23,35] and amphibians [40]. RF is a machine learning technique with a number of characteristics that make it suitable for extinction risk prediction [15], including: limited assumptions about data distributions, high classification stability and performance, and ability to cope with collinear predictors. In a recent test, RF showed the highest performance in predicting global mammal extinction risk among several machine learning methods [23]. Our model included several variables which are external to species biology (human pressures, habitat state and conservation responses); hence, in common with other studies [15], we did not include phylogenetic constraints in our analyses. However, we tested whether this could influence our results by independently examining the effect of including taxonomy for predicting extinction risk [23].

We ran a full RF model, including all predictor variables, and ranked the variables according to their relative importance (i.e.
their contribution to model’s classification accuracy). Variable importance, as well as the classification accuracy of the model, was calculated using an automated bootstrapped cross-validation procedure (implemented within the RF routine). During each iteration of the RF model, one-third of the data were left out and used to cross-validate the classification ability of the model (see [41] for additional details).

Based on the final variable importance scores, we ran a series of partial RF models, each time including one additional variable following the variables’ ranked importance. First, we ran the

Table 1. Description of the variables used in the model. Variables are organized in different classes: human pressure (P), species biology (B), distribution state (D) and conservation response (R). Examples of previous use of the variables for predicting extinction risk in terrestrial mammals, and the original data sources for each variables are also provided.

<table>
<thead>
<tr>
<th>class</th>
<th>variable</th>
<th>description and justification</th>
<th>examples</th>
<th>source</th>
</tr>
</thead>
<tbody>
<tr>
<td>—</td>
<td>dependent variable</td>
<td>extinction risk transition as described in electronic supplementary material, table S1</td>
<td></td>
<td>[4,16]</td>
</tr>
<tr>
<td>P</td>
<td>Acc_50</td>
<td>travel distance from major cities (accessibility), measured as the median value of the variable within species ranges (percentiles tested: 5, 10, 20, 50); a proxy of human encroachment</td>
<td>[22,23]</td>
<td>[24]</td>
</tr>
<tr>
<td>P</td>
<td>AOIloss</td>
<td>proportional loss of suitable habitat within species ranges (1970 – 2010); a proxy of the main driver of mammal species decline calculated from back casts of global land cover changes, from the IMAGE integrated assessment model [25]</td>
<td>[22]</td>
<td>[26,27]</td>
</tr>
<tr>
<td>P</td>
<td>HII_5</td>
<td>human influence index, measured as the proportion of species ranges where the variable had values larger than 5 (values tested: 5, 10, 20); a proxy of the human impact on the environment</td>
<td>[22,23]</td>
<td>[28]</td>
</tr>
<tr>
<td>P</td>
<td>HPD90_50</td>
<td>human population density in 1990, measured as the median value the variable within species ranges (percentiles tested: 5, 10, 20, 50); a proxy of human encroachment</td>
<td>[13,22,23]</td>
<td>[29]</td>
</tr>
<tr>
<td>P</td>
<td>PopChange</td>
<td>proportional change in human population count in 1990 – 2010, measured as the mean value observed within species range</td>
<td></td>
<td>[30]</td>
</tr>
<tr>
<td>P</td>
<td>ForestCG</td>
<td>proportional change in forested habitat within species ranges between 2000 and 2012. A proxy of natural habitat loss</td>
<td></td>
<td>[31]</td>
</tr>
<tr>
<td>B</td>
<td>AFB_d</td>
<td>age at first birth</td>
<td>[32,33]</td>
<td>[34]</td>
</tr>
<tr>
<td>B</td>
<td>BirthW</td>
<td>birth weight</td>
<td>[22]</td>
<td>[34]</td>
</tr>
<tr>
<td>B</td>
<td>BodySize</td>
<td>body mass</td>
<td>[13,23,35]</td>
<td>[34]</td>
</tr>
<tr>
<td>B</td>
<td>DietBrdth</td>
<td>number of dietary categories eaten by the species</td>
<td>[22]</td>
<td>[34]</td>
</tr>
<tr>
<td>B</td>
<td>InterbInt</td>
<td>interbirth interval</td>
<td>[32]</td>
<td>[34]</td>
</tr>
<tr>
<td>B</td>
<td>LitPY</td>
<td>litters per year</td>
<td>[34]</td>
<td></td>
</tr>
<tr>
<td>B</td>
<td>LitSiz</td>
<td>litter size</td>
<td>[22,23,32]</td>
<td>[34]</td>
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<tr>
<td>B</td>
<td>WeanAge</td>
<td>weaning age</td>
<td>[13,32]</td>
<td>[34]</td>
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<tr>
<td>B</td>
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<td>[4]</td>
<td></td>
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<tr>
<td>B</td>
<td>Ord</td>
<td>taxonomic order</td>
<td>[13,22]</td>
<td>[4]</td>
</tr>
<tr>
<td>B</td>
<td>Genlen</td>
<td>generation length</td>
<td>[32]</td>
<td>[21]</td>
</tr>
<tr>
<td>B</td>
<td>HabBrdth</td>
<td>number of habitat layers used by each species</td>
<td>[34]</td>
<td></td>
</tr>
<tr>
<td>D</td>
<td>TreeCover_50</td>
<td>median tree cover within species range in 2000 (percentiles measured: 5, 10, 20, 50). A proxy of forests state</td>
<td>[31]</td>
<td></td>
</tr>
<tr>
<td>D</td>
<td>Hab</td>
<td>species habitat preferences, classified as: forest, grassland, shrubland, bareland, coastal or generalist (when more than 1 of the previous applied)</td>
<td>[27]</td>
<td></td>
</tr>
<tr>
<td>D</td>
<td>RangeSize</td>
<td>species range size, measured as an order of magnitude (e.g. 1 for ranges of 10 – 100 km², 2 for ranges of 100 – 1000 km², etc.)</td>
<td>[13,22,35]</td>
<td>[4]</td>
</tr>
<tr>
<td>R</td>
<td>RangeProt_prop</td>
<td>proportion of species range covered by PAs with an IUCN category I to IV</td>
<td>[22]</td>
<td>[36]</td>
</tr>
<tr>
<td>R</td>
<td>RangeProtkm</td>
<td>extent of PAs within species ranges, measured as an order of magnitude (as described for ‘RangeSize’)</td>
<td>[36]</td>
<td></td>
</tr>
</tbody>
</table>
model including only the most important variable, then added the second most important variable and re-ran the model, and so on until the last variable was included. We measured the performance of each partial RF model in terms of proportion of correctly classified species, proportion of correctly classified higher risk species (sensitivity), proportion of correctly classified lower risk species (specificity) and true skill statistic (TSS = sensitivity + specificity - 1) [42].

In order to account for the effect of including the original (1975) species Red List status in the model, we re-ran the full model after removing this variable. Because of its potential role in Red List assessments and its representation of past threat conditions [43], we also re-ran the model after removing species’ range size (RangeSize). In this latter case, we also removed the variable representing extent of PA within the species range (RangeProtkm), as it has a weak positive correlation with range size ($R^2 = 0.56$). We used degraded values of both range size and PA extent (i.e. order of magnitude rather than actual values; as for previous work [43]), to better represent the availability of coarse and approximate information during the study period. Finally, we built a single conditional inference classification tree to visually represent the interaction between predictor variables.

We adopted alternative classifications of extinction risk transitions and tested the performance of our model under different formats of the response variable. First, we repeated our RF modelling using ordinal changes in Red List categories as a numeric response variable (e.g. +2 for a species moving from LC to VU; see also [20]). Second, we repeated our RF modelling after removing all species that did not change their Red List category between 1975 and 2013; in this case, we classified the remaining species in two categories: ‘uplisted’ for species moving to higher extinction risk categories and ‘downlisted’ for species moving to categories of lower risk. Third, we divided species in three categories of lower risk. Third, we divided species in three categories of extinction risk and ‘downlisted’ for species moving to higher categories of lower risk (RangeSize). In this latter case, we also removed the variable representing extent of PA within the species range (RangeProtkm), as it has a weak positive correlation with range size ($R^2 = 0.56$). We used degraded values of both range size and PA extent (i.e. order of magnitude rather than actual values; as for previous work [43]), to better represent the availability of coarse and approximate information during the study period. Finally, we built a single conditional inference classification tree to visually represent the interaction between predictor variables.

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The quantification of spatial variables was performed in GRASS GIS [44]. Statistical analyses were performed in R [45] using the packages ‘randomForest’ [41] and ‘party’ [46].

### 3. Results

Our classification of extinction risk resulted in 277 species being included in the lower risk group (55% of all species) and 220 species in the higher risk group (45% of species). The full RF model for classification of higher risk versus lower risk species performed well in cross-validation (table 2): 89% of all species were correctly classified, with a sensitivity of 0.84, and a specificity of 0.93 (TSS = 0.77). After removing the Red List category in 1975 from the model (i.e. the most important predictor), 82% of the species were still correctly classified, but the ability to correctly classify higher risk transitions was reduced (sensitivity = 0.78; TSS = 0.64). Subsequent removal of range size caused further deterioration in the model performance; although 79% of species were still correctly classified, there was a substantial reduction in sensitivity and TSS (sensitivity = 0.73; TSS = 0.57).

The six most important variables in the full RF model were: Red List category in 1975, PA extent (representing conservation response), range size (representing distribution state), body size (representing biology), family (representing taxonomy) and human impact index (representing human pressure; see figure $2b$). A sequence of partial RF models, adding one variable at a time from the most important to the least important, showed that some of the variables had a contrasting effect on sensitivity and specificity. For example, adding the taxonomic family to the model substantially increased sensitivity, but reduced specificity. By contrast, adding the human influence index slightly increased both sensitivity and specificity.

The extinction risk transition of 87% of species could be correctly predicted from one variable alone (Red List category in 1975), highlighting the importance of knowing the initial condition when modelling changes in extinction risk. However, this was biased towards lower risk species (specificity = 0.95 versus sensitivity = 0.78). Adding five additional variables did not substantially alter the overall classification ability, but improved the balance between specificity and sensitivity (figure $2n$). Even after removing the Red List categories in 1975 from the model, the performance remained fairly good, but then several variables had to be included in order to correctly classify approximately 78% of the higher risk and approximately 86% of the lower risk species (figure $2b$). Subsequent removal of range size required the use of more than 50% of all variables to achieve a sensitivity of approximately 73% and specificity of approximately 83% (electronic supplementary material, figure S1).

A single conditional inference tree (figure $3$) represents the interplay between correlates of extinction risk transitions. For example, species that were LC in 1975 had a much higher probability of being in the higher risk group if they had a relatively low coverage of PAs during the study period (less than 1000 km$^2$) and faced a substantial increase in human population change, generation length, age at first birth and proportion of PAs (electronic supplementary material, figure S2).

When excluding species that did not undergo a change in their Red List category, our sample reduced to 15 downlisted and 113 uplisted species. The RF model then gave highly

<table>
<thead>
<tr>
<th>metric</th>
<th>full model</th>
<th>RL75 removed</th>
<th>RL75 and RangeSize removed*</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCC</td>
<td>0.89</td>
<td>0.82</td>
<td>0.79</td>
</tr>
<tr>
<td>sensitivity</td>
<td>0.84</td>
<td>0.78</td>
<td>0.73</td>
</tr>
<tr>
<td>specificity</td>
<td>0.93</td>
<td>0.86</td>
<td>0.84</td>
</tr>
<tr>
<td>TSS</td>
<td>0.77</td>
<td>0.64</td>
<td>0.57</td>
</tr>
</tbody>
</table>

*When removing the variable RangeSize, the extent of PAs within the range was also removed to avoid a potential surrogate effect.

### Table 2. Performance of the RF models. The full model is compared with partial models, where the original species status (RL75) and the range size (RangeSize) were removed. PCC, proportion of correctly classified species; TSS, true skill statistic.
biased results in this case, due to the high class imbalance, and classified all species as being uplisted (i.e. a complete imbalance towards sensitivity). The overall classification accuracy in this case was misleadingly high (88%), as the model was unable to predict improvement in species conservation status.

When dividing species into three groups, there were 15 downlisted species, 262 LC to LC species and 220 higher risk species. Here again, the overall classification accuracy of the model was high (89%), but the predictive ability for the downlisted class was very low (only one correct prediction; electronic supplementary material, table S2).

4. Discussion
By focusing on extinction risk transitions, we were able to distinguish between two groups of species. The higher risk group included species that remained at high extinction risk and those whose extinction risk increased between 1970 and 2010. The lower risk group included species that remained at, or improved their status to, low extinction risk during the same period. This classification is different from the Red List status, as it identifies species that are undergoing an unusual increase in extinction risk compared with other species that started the period in the same risk category.

We included candidate predictor variables from a range of classes (see Material and methods) and found that a small number of variables (from different classes) can efficiently predict the extinction risk transition of ungulates and carnivores. These variables have been highlighted previously [13,35] and include initial conservation status, certain biological traits (represented by body mass), levels of human encroachment and the degree of conservation action (represented by PA coverage). The importance of considering...
conservation interventions in extinction risk modelling has already been demonstrated for Australian birds [20] and for African mammals [22], and we confirm it here in a global-scale analysis.

Our results show that the probability of a species being at higher risk was reduced by some adequate level of PAs coverage (1000 km² or more; figure 3), while it was increased by limited PA coverage and high levels of human pressure. To a first approximation, this indicates the conditions under which PAs deliver positive conservation outcomes [47]. Monitoring the progress of PA expansion and the extent of human encroachment within species ranges can therefore be strategic. Future projections of these variables may be translated into global projection of species extinction risk and allow for a proactive planning of conservation interventions [48].

Our models included measures of environmental change (e.g. the amount of suitable habitat for a species during the study period) and static measures of human impact (e.g. human influence index). These classes of variables were both important predictors in our model. Among general proxies of human pressures and habitat state, we also included information on levels of tree cover and tree cover change (see also [22]). While the role of these variables is probably more influential for forest-dependent than for non-forest species, it is known that habitat clearance has a contagious effect [49], and we use tree cover, a well-mapped habitat feature at a global scale [31], to estimate the general condition of natural habitats within species ranges.

The extinction risk transition model performed well in cross-validations; the classification ability was high for both lower risk and higher risk species. The availability of a dataset with retrospective extinction risk assessments [16] made it possible for us to validate our extinction risk model. This type of validation is common in other environmental science areas and has been used to validate models of climate change effects on species distribution [50]. As our knowledge of past extinction risk improves, this approach could become standard practice in extinction risk modelling.

Unlike many previous studies, we did not convert IUCN Red List categories into numerical measures of extinction risk (e.g. LC to Extinct, from 0 to 5 [20,51]) or use extinction risk probabilities described in Red List Criterion E [52]. These involve assumptions about the relationship between categories and probability of extinctions that are not supported in theory or in practice [11]. We simply assumed that species in the higher risk group have higher conservation requirements than those in the lower risk group, and found that predicting ordinal changes in Red List categories (as in [20]) was substantially less efficient than predicting extinction risk transitions. We also found that excluding those species with no change in their Red List category, or assigning stable LC species to a separate group, resulted in a biased allocation of model error, with downlisted species being systematically misclassified. In this case, the model is unable to predict the outcome of conservation success (i.e. those situations in which the extinction risk of a species is reduced over years).

Our results on the relative importance of different predictor variables can be used to identify priorities for future data gathering. We suggest that monitoring a set of such variables over time would allow conservationists to effectively anticipate future extinction risk. The accuracy of these predictions will rest on the assumption that these variables represent the drivers of transitions in species extinction risk. Our results demonstrate that this was the case for past extinction risk transitions, but the emergence (or the exacerbation) of new threats (such as climate change) would need to be accounted for to have a robust forecasting of extinction risk [17,53]. However, this is not a weakness unique to our approach: threats to biodiversity change over time [54] and any model used to forecast extinction risk would require continuing updates and recalibration to account for emerging threats. Monitoring the emergence of new threats and the occurrence of rapid changes in external conditions will be necessary, yet

![Figure 3. Conditional inference classification tree for extinction risk transition. Each terminal node reports (in dark grey) the proportion of higher risk species. See table 1 for a description of the variables.](http://rspb.royalsocietypublishing.org/Downloaded from rspb.royalsocietypublishing.org)
even this would probably be easier than continuously assessing the extinction risk category of all species.

McCarthy et al. [20] investigated optimal investment strategies to prevent the extinction and minimize the number of threatened Australian birds, using conservation investments to model the probability of species moving between Red List categories. A similar approach could be combined with our modelling framework here, to measure the probability of undergoing a high-risk transition. In this case, the probability can be modelled as a function of the intrinsic and extrinsic conditions in place for the species, plus the conservation budget available. However, adequate information on global conservation expenditure for threatened species needs to be available to reliably model the relationship between investments and status change.

Our approach can provide guidance on how to allocate resources among monitoring of species extinction risk and monitoring of external conditions; it can inform the identification of key variables to be monitored. There is great potential for the application of our approach to other taxa, especially considering the increasing availability of retrospective extinction risk assessments for groups such as amphibians [55] and corals [56], and the potential to use historical information to perform retrospective assessments for other groups [16].

Data accessibility. Data used in this paper come from published sources which have been appropriately cited in the ‘Material and methods’ section.

Authors’ contributions. M.D.M. and G.M.M. conceived the study design; M.D.M. performed the analyses; all authors interpreted the results, contributed to the writing and approved the final version of the paper.

Competing interests. We declare we have no competing interests.

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