Extinction debt and colonization credit delay range shifts of eastern North American trees

Matthew V. Talluto¹*^(D), Isabelle Boulangeat², Steve Vissault³, Wilfried Thuiller¹ and Dominique Gravel³

Global climate change is already having an impact on species ranges. For species with slow demography and limited dispersal, such as trees, lags between climate change and distribution shifts are likely to increase in the future. Such lags can be of critical importance to management and biodiversity of forests, because they can result in 'extinction debts', where populations temporarily persist under unsuitable conditions, and 'colonization credits', where suitable locations are not occupied owing to slow demography and limited dispersal. Here we use a range dynamics model based on metapopulation theory to show that the distributions of 21 dominant trees in eastern North America are out of equilibrium with climate and demonstrate both extinction debt and colonization credit. Moreover, lags are more severe at northern range limits, suggesting that range contraction in response to warming temperatures will outpace expansion.

ecause climate is as a major driver of species distributions, global warming is expected to induce species range shifts¹. The close relationship between climate and species distribution is the basis of correlative species distribution models^{2,3}, which have been extensively used to derive biodiversity scenarios4-7. However, species do not respond to change immediately, and consequently distributions can lag behind climate change⁸. This is particularly the case for long-lived and sessile organisms. Demographically 'slow' species such as trees are vulnerable to such lags, leading to so-called extinction debts, where populations persist for some time after unsuitable conditions render eventual extinction likely9-11, and corresponding colonization credits, where suitable but unoccupied locations are only slowly filled owing to slow demography. Such mismatches between optimal and realized climates have already been observed¹² and are likely to be exacerbated in the future, producing lags between climate change and shifts in species distributions that may extend well beyond the timescales typical of ecological studies. It is critical to predict these situations for forest management and biodiversity conservation, which rely on understanding which species can persist at a given location, and which are often interested in short-time predictions (that is, when lags are most likely to occur).

It is not straightforward to identify disequilibrium between climate and species distributions using correlative species distribution models (SDMs). Instead, dynamic models provide the advantage of linking local processes with species ranges. In addition, these models need to be able to explain species distributions at the scale of their entire range, and capture the variation in key processes according to climate variables. Therefore, although process-based models (for example, PPA13, JABOWA14. LANDIS II15, FATE-HD16) can easily capture local dynamics, they are inherently data demanding and computationally intensive^{17,18}, making the estimation of entire distributions at continental scales challenging. Here we propose a novel approach inspired by metapopulation theory¹⁹. Our approach stochastically models the dynamics of presence and absence at a locality as a function of two fundamental ecological processes: colonization of empty localities and extinction from occupied localities. Although theoretical studies have shown how

colonization–extinction dynamics can lead to range limits^{20,21}, this is, to our knowledge, the first time the theory is applied to explain species distributions.

We fit this model to a database of repeated observations of more than 90,000 forest inventory plots spanning the eastern United States and Canada (Supplementary Fig. 1). This approach is ideal to represent forest dynamics owing to the strong imprint that succession and natural disturbance leave on forest landscapes (Fig. 1). We analysed range dynamics to look for lags in the response to climate change in 21 dominant trees across temperate, transitional and boreal forests (Supplementary Table 1) and to evaluate future rates of response to climate change at the northern and southern range limits. This transition is particularly relevant because of the major shifts in ecosystem processes that are expected to follow colonization of boreal forests by deciduous and temperate species. Our principal hypotheses were that range limits are set by a balance between colonization and extinction dynamics and that the slow nature of these demographic processes results in lags in the response to climate change. Consequently, even though local conditions could allow populations to persist temporarily in a given location²², we expect that range limits of forest tree species are ultimately determined by regional processes such as dispersal limitation, succession and natural disturbance.

Results

The metapopulation theory of range dynamics predicts that range limits will occur where colonization is balanced by extinction. Although originally developed to represent a set of populations connected by dispersal¹⁹, it was later applied to continuous landscapes, even representing the dynamics of individuals in a spatially structured population²³. We computed the probability of colonization (c(E)) and extinction (m(E)) as functions of climate E for all species, then solved these for the environmental conditions E^* such that $c(E^*) = m(E^*)$ (Fig. 1 and Supplementary Figs 2–4). This yields the range limits at equilibrium, allowing for comparisons between contemporary distributions and long-term equilibria. We found realistic ranges for all of the species (Fig. 2 and Supplementary Fig. 5) and surprisingly low uncertainty in the predicted range limits

¹Université Grenoble Alpes, CNRS, UMR 5553, Laboratoire d'Écologie Alpine (LECA), F-38000 Grenoble, France. ²Department of Biosciences, Ecoinformatics and Biodiversity, Aarhus University, 8000 Aarhus, Denmark. ³Département de biologie, Université de Sherbrooke, Sherbrooke, Quebec, Canada. *e-mail: matthew.talluto@univ-grenoble-alpes.fr

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Figure 1 | Schematic demonstrating how metapopulation dynamics can lead to range limits. A metapopulation approach to range dynamics hypothesizes that range limits are determined by colonization (c, blue lines) and extinction (m, red lines) probabilities as they vary along an environmental gradient⁹. \mathbf{a} , c and m for Acer saccharum vary with temperature, resulting in a latitudinal gradient. More generally, we describe c and m as functions of the environment: c(E) and m(E). The theory predicts that the species will be present at equilibrium (shaded region) where c(E) > m(E); therefore the temperature at which c(E) = m(E) determines the latitude of the range limits. **b**, A simulated 2.5 °C of warming produces a northward shift in the suitable range (grey and blue shading). However, over short time scales, slow extinction probabilities will cause the species to persist where m(E) > c(E) (red shading) until all occupied populations have gone extinct. A similar lag in colonization ability will result in the species being absent in many areas where c(E) > m(E) (blue shading). Thin lines show 90% credible intervals around mean c and m probabilities.

(Supplementary Fig. 6), despite considerable variability in the colonization and the extinction functions (Supplementary Figs 2,3). However, compared with prior expectations about where the species occur (for example, from published range maps), we observed some systematic discrepancies at the borders of most species' ranges. We therefore compared the difference between the predicted equilibrium ranges from our fitted metapopulation model with current distributions interpolated from the forest inventory data using the random forest algorithm. Because this algorithm is extremely flexible and was parameterized with a large dataset, these maps presented a highly accurate picture of the current distributions (Supplementary Fig. 7), in contrast to the long-term climatic equilibrium predicted by the metapopulation model. For all species, we found extensive areas of extinction debt, where the current ranges indicate that the species is present while the longterm expectation is absence. These areas were generally found at the southern edge of current ranges. All species also had a corresponding colonization credit, where the current maps indicate absence but the long-term expectation is presence (Fig. 2). Colonization credit was usually found at the north and west of the current range. This strong latitudinal pattern in extinction debt and colonization credit suggests that the disequilibrium in species ranges reflects the signal of contemporary climate change, where climatic optima have already shifted northward of where the species are presently located.

In order to better understand the transient conditions leading from the present ranges of species to their equilibrium distributions, we further analysed our model with respect to the rate of change in response to climate change at the range boundaries. The simplicity of the theory facilitates this analysis, since the intrinsic growth rate of a metapopulation, defined as $\lambda(E) = c(E) - m(E)$ within the range and $\lambda(E) = m(E) - c(E)$ outside the range, approximates the rate of response to a slight change in the environmental conditions. Thus, the responsiveness of the system, given by the partial derivative $\partial \lambda/\partial E$ at the range boundary, indicates how quickly the system will track new equilibrium conditions in response to climate warming.

NATURE ECOLOGY & EVOLUTION

For species that had both range boundaries within the study area, responsiveness was much greater at southern range boundaries than in the north (grand mean difference across species = 0.14; 95% confidence interval = 0.046-0.24; n = 18), although there was substantial interspecies variation (Fig. 3). Therefore, future increases in temperature will probably result in faster responses at southern range limits, where northward contraction is expected, and slower responses in the north, where range expansion is expected (Fig. 2). We further expect that temperate species will respond more quickly than boreal species due to the overall trend of increased responsiveness at southern latitudes.

Ecological theory predicts that the climate dependence of colonization-extinction dynamics should be related to individual-scale demographic variation^{24,25}. It follows that regional-scale extinction debts are a result of increased mortality or a failure of recruitment at the individual scale. To test this hypothesis, we extracted information on individual trees from our database and analysed mortality and recruitment rates. Grouping across all species, average mortality was lower within equilibrium ranges than in areas facing extinction debt, with a mean rate of 0.044 individual⁻¹ year⁻¹ (0.038-0.050) within the range and 0.063 individual⁻¹ year⁻¹ (0.051-0.078) in areas facing extinction debt (values are mean predictions from a hierarchical model with 90% credible intervals). Recruitment rates did not differ, with rates of 0.038 individual⁻¹ year⁻¹ (0.034–0.043) in the equilibrium range and 0.045 (0.037-0.055) in areas facing extinction debt. Mortality estimates varied considerably among species, but were substantially higher in the regions facing extinction debt in 16 of the 21 species (Fig. 4), with boreal, transitional and temperate species all showing the same trend. Therefore, elevated mortality rates at the range margins, rather than recruitment failure, are likely a principal cause of range limits and may contribute to the loss of populations in regions facing extinction debt.

Discussion

Our model predicts that the distributions of many of the most abundant trees occurring in the boreal-temperate ecotone in eastern North America are out of equilibrium with climate at the range margins (Fig. 2). Similar results have been observed in other systems, suggesting that the distributions of many species^{8,12,26} (and therefore the ecosystems in which they are present) may be in disequilibrium with climate. Models failing to account for this bias could therefore produce overly optimistic estimates of the tolerance of many species to climate change. In particular, these results call for caution when forecasting and planning using SDMs. When species distributions are out of equilibrium with climate, SDM estimates of species' optimal climates and thus potential future ranges are likely to be biased, because the species may be inhabiting unsuitable regions and may not yet have colonized all suitable regions. The strength of our modelling approach rests on the strong theoretical foundation, which predicts that range limits occur where colonization and extinction rates are balanced²⁰. We have shown for the first time and with high precision that this theory is well-suited to explain the distribution of forest tree species, and that we can predict the static distribution of a species using only dynamic data. This offers us the unique opportunity to identify current extinction debts and colonization credits.

A further benefit to using a simple and tractable theory to understand range dynamics is having a comprehensive understanding of transient states, such as the responsiveness of range limits to climate change. We have shown that many dominant trees in eastern North American forests have low capacities to track climate change (Fig. 3). In the future, most species are expected to experience range losses at the southern end of the distribution and gains at the northern end even with no warming beyond the present climate. Given the expectation that northern regions of our study area will warm by as much as 5 °C by 2100 (ref. ²⁷), changes to range limits (and corresponding

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Figure 2 | Maps of projected species ranges with extinction debt and colonization credit for selected species. The maps indicate, for the present climatic conditions, where the species is (1) currently present and predicted to be present at equilibrium (continued presence; blue), (2) currently absent but predicted to be present at equilibrium (colonization credit; green), and (3) currently present but predicted to be absent at equilibrium (extinction debt; pink). Equilibrium presence was assessed using the equilibrium solution of the state and transition model (that is, c(E) > m(E)), and current presence was assessed using the random forest model with a threshold (computed using a semi-independent evaluation dataset) applied to the probabilities.

changes to community composition) will almost certainly be larger than those we have predicted here. However, like in other studies^{11,28-31}, we find that such changes will happen slowly, on the one hand because the high longevity of trees creates extinction debt in warmer regions, on the other hand because slow colonization constrains range expansion in the north. Moreover, our results should be interpreted with caution with respect to boreal ecosystems, as our database does not include the boreal-tundra transition. Thus, the large losses we predict may be offset by boreal encroachment into tundra. Such shifts may be complex and difficult to predict with respect to warming temperature³². Overall, temperature increases are likely to continue to outpace distributional changes, resulting in increasing ecological tension between extant populations and the climate required for sustainable populations. Effective management strategies and climatic adaptation would have to account for such increasing disequilibrium in the future.

As with SDMs, the need to choose from a limited set of key variables presents a challenge when modelling some species. It is likely that additional variables can influence the colonization-extinction probabilities. If these variables are spatially structured, failing to account for them can result in a spatial bias in the colonizationextinction functions and incorrect projections of species ranges. An analysis of the residuals in our models (Supplemental Methods and Supplemental Fig. 9) suggested that spatially structured variables may indeed be missing for certain species. However, a limited spatially explicit version of the model showed no evidence of bias (Supplemental Figs 10-12). It is clear, however, that for some species a complete picture of what variables influence range dynamics is lacking, and thus dynamic models must be interpreted with caution. Even when the models accurately capture species dynamics in the calibration range, important variables and interactions that are missing can make prediction and analysis of transient states unreliable.

One of the main strengths of our approach is to shift from modelling species ranges and their dynamics from a static, niche-based perspective to a dynamic perspective. However, the model is unable to distinguish fine-scale mechanisms underlying the relationships between climate and colonization and extinction probabilities. For example, interactions between climate and local age-specific demographic rates could result in nonlinearities in the relationships between climate and colonization and extinction that would not be detectable using our method (for example, through storage effects³³ or mass effects²⁸). To properly account for such mechanisms, our dynamic approach to modelling species ranges could be extended using more detailed models³⁴⁻³⁶ when their parameterization is feasible. More generally, the perspective of modelling species and community changes instead of their distributions offers numerous opportunities to improve the predictive power of biodiversity models.

Methods

Model description. Our model design is a simple mathematical model following classical metapopulation theory¹⁹, extended to incorporate environmental gradients²⁰. A metapopulation consists of a set of local populations connected together by dispersal. Patches are either occupied (that is, the species is present) or unoccupied. We used a discrete-time variant of the Holt model²⁰. In discrete time, the patch occupancy dynamics of the metapopulation can be conceptualized as a spatially inhomogeneous Markov chain with state vector $\mathbf{P}_{i.1}$ (with $\mathbf{P}_{i.0}$ unoccupied patches and $\mathbf{P}_{i.1}$ occupied patches at time *t*) and transition matrix *S*, such that:

 $\mathbf{P}_{t+1} = \mathbf{P}_t S$

The model is made inhomogeneous by parameterizing the transition matrix with colonization and extinction probabilities that are estimated as functions of the environment:

$$S = \begin{bmatrix} 1 - p(E) c(E) & p(E) c(E) \\ m(E) & 1 - m(E) \end{bmatrix}$$
(1)

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where *E* represents environmental conditions, *t* represents time, p(E) is the fraction of the landscape where the species is present, c(E) is the colonization probability and m(E) is the extinction probability. For example, for a colonization to occur in a patch with environmental conditions *E*, the patch must be initially empty (with probability 1-p(E)), it must have a source of available seeds (with probability p(E) representing the prevalence of the species in the immediate neighbourhood of the focal patch) and seeds must arrive and establish in the patch (with probability (c(E)). At equilibrium, the species persists where c(E) > m(E); thus the range limit is found at environmental conditions *E**, where $c(E^*) = m(E^*)$ (Fig. 1).

Data preparation. We estimated the climate-dependence of colonizationextinction probabilities of dominant tree species found in a region encompassing 27 degrees of latitude in eastern North America and spanning the transition from temperate to boreal forest. To perform these analyses, we assembled a database from regional forest inventory databases from the Forest Inventory and Analysis National Program in The United States³⁷, Domtar (a private forestry company in Quebec), the Ministry of Forests, Wildlife and Parks in Quebec³⁸, the Ministry of Natural Resources and Forestry in Ontario³⁹, and the Ministry of Natural Resources in New Brunswick⁴⁰. Although protocols and data storage differed among databases, we were able to build a combined database that included common measurements from each database, including plot locations and sizes, years of measurement, and the diameter at breast height (DBH), species and status (living or dead) of every tree in each plot with a DBH > 127 mm. Smaller trees were not recorded consistently across databases and sampling years, and were therefore excluded from our analysis to minimize geographic biases due to the differing protocols. Initial investigations using models containing smaller trees showed no detectable differences in colonization-extinction curves, therefore we chose to filter these trees to maintain consistency among data sources. To account for potential bias (because the DBH threshold may represent different ages for different species, influencing the probability of colonization), we used ages derived from tree cores to (available for 16 out of 21 species) to estimate age-DBH curves (Supplementary Fig. 8). We then used linear models to estimate the mean age at which each species reached 127 mm (Supplemental Table 2).

From this database, we extracted repeated surveys ($2 \le n \le 10$ surveys per plot, where a survey is the record of all trees in a given year) from 99,724 total plots (Supplementary Fig. 1). Owing to the variety in the survey methods represented in our database, plot sizes ranged from 400–1,513 m² (median = 672 m²).

NATURE ECOLOGY & EVOLUTION



Figure 4 | Predicted recruitment and mortality rates in equilibrium ranges and in areas facing extinction debt. Mortality rates were systematically greater in extinction debt zones (red) than in the equilibrium range (blue), whereas differences in recruitment rates varied less systematically. Bars show mean predicted rates (with 90% credible intervals) from a hierarchical model with species as a grouping factor. Data are for all individuals with DBH > 127 mm.

To select our study species, we computed the basal area (in $m^2 ha^{-1}$) of all species occurring in plots in the 42–50° latitude range (representing the approximate limits of the temperate–boreal transition zone). We then ranked the species by basal area within these plots and selected the most abundant species that summed to at least 90% of the basal area. This procedure resulted in 21 species representing 90% of the basal area, including nine primarily boreal species, eight temperate species and four transitional species.

To identify observed colonizations and extinctions, we searched the database for plots that were surveyed multiple times. We defined a sampling unit as an initial observation within a plot and its subsequent repeat (therefore, for example, a plot surveyed in 1990, 1995 and 2000 had 2 samples: 1990–1995 and 1995–2000). For each species we classified every sampling unit as an absence (that is, absent for both observations; n = 37,265-79,137 observations per species), a presence (present during both observations; n = 1,763-30,403), a colonization (absence followed by presence, n = 112-2,684), or an extinction (presence followed by absence, n = 108-2,182) (Supplementary Table 1). We removed intervals greater than 15 years from the dataset. This minimizes the risk of multiple transitions within an interval (for example, a colonization followed by an extinction, which would be observed as an absence).

To fit climatic relationships, we used annual 10-km² climate rasters for the entire study area covering the period from 1945–2010 that included six bioclimatic variables: annual mean temperature (°C), mean diurnal temperature range (°C), the mean temperature in the wettest quarter (°C), total annual precipitation (mm), precipitation seasonality and mean precipitation in the warmest quarter (mm). All bioclimatic rasters were obtained from ANUSPLIN, a non-parametric, multi-dimensional curve fitting technique that interprets data from meteorological stations into continuous surfaces⁴¹. To minimize the effects of extreme years on model calibration, climate for a plot in a given year was then computed as the mean of the climate over the 15 years preceding the observation. Because the full dataset includes forest plots for many species, for any given species many plots were well outside the species' climatic range and provided little additional information for fitting colonization–extinction curves (which are most sensitive

NATURE ECOLOGY & EVOLUTION

ARTICLES

at the range boundaries). Fitting a model with the full dataset thus potentially puts a large amount of likelihood weight in the tails of the colonization-extinction curves where both colonization and extinction probabilities are near zero (because the species is never present). Therefore, to avoid overfitting to the tails and to put the emphasis on climatic conditions where the species is potentially present, we restricted the calibration data range by eliminating all plots that were outside a 10° latitude and longitude buffer surrounding all observed presences. For similar reasons, we truncated climatic data that extends beyond the northern limit of the forest surveys and do not project our models beyond these limits.

Calibration. Our primary modelling goal was to provide a simple interpretation of how the colonization-extinction probabilities varied as functions of temperature and precipitation. Therefore, we selected a priori a single representative of each for model fitting. On the basis of exploratory analyses, mean annual temperature and total annual precipitation demonstrated good relationships to colonizationextinction probabilities and simple interpretations, and these were therefore selected for all species. The colonization-extinction functions (equation (1)) were fit as second-order polynomials of temperature and precipitation using a Bayesian model with binomially distributed errors and a logistic link function. All model parameters were fit with weakly informative Cauchy priors⁴². Because the sampling interval was not equal across samples (range 5-15 years), it was necessary to standardize the parameters to a single sampling interval and then compute the likelihood of an observation given the (constant interval) transition probability and the observation interval. We selected five years as the standard interval because the majority of plots were sampled at five-year intervals. Therefore, the likelihood of an observed extinction (for instance) over an *n*-year interval (m_n) given the standardized probability over a five-year interval (m_5) can be computed as:

$$m_n(T,P) = 1 - [1 - m_5(T,P)]^{\binom{n}{5}}$$
(2)

where *T* and *P* are mean annual temperature and total annual precipitation, respectively. The likelihood of an observed colonization, $c_n(T,P)p(T,P)$, was similarly computed, with the added complication that, under metapopulation theory, an observed colonization depends on both the probability of colonization and establishment ($c_n(T,P)$) and the prevalence (p(T,P)), representing the availability of a nearby seed source.

Computing the exact prevalence in the neighbourhood of the plots was impossible with the forest inventory data, as landscape-level surveys of surrounding forests were unavailable and most forest community maps and remote sensing data sources do not resolve to abundances of individual species. Instead, we estimated prevalence using the random forest algorithm. The response variable with this model was the binomial variable ($\phi(E)$, N(E)), where $\phi(E)$ is the number of plots in which the species was present under environmental conditions *E* and N(E) is the total number of plots under the same environmental conditions. The predictor variable *E* consisted of the six climatic variables described above. This model was then projected to the individual plots, where the prevalence p_c of a species at a particular plot *x* was interpreted as the conditional expectation of the random forest model θ : $p_c = \mathbf{E}(\theta|E_x)$.

To estimate the posterior distributions of the parameters of the colonizationextinction functions, we used Markov chain Monte Carlo (MCMC) with a Metropolis-Hastings sampler. To assure that the samplers had forgotten their initial conditions and were sampling from the stable distribution, we ran four chains with overdispersed starting values and computed the potential scale reduction factor⁴³ (\hat{r}). Chains were considered to be converged when the upper bound of \hat{r} was less than 1.1 for all parameters. We then selected a single chain at random, ran it for 50% longer than required for convergence, and discarded the first 50% of samples. Finally, to reduce autocorrelation in the samples, reduce disk storage requirements and simplify derived computations, we thinned the samples, retaining a single sample at a regular interval such that the final sample size was 10,000. To prevent skewed posterior distributions from influencing parameter estimates, we report posterior medians, rather than means, for all results. We also report posterior uncertainty intervals for all parameters and derived statistics. Unless otherwise specified, these intervals are reported as 90% credible intervals and are computed as the 90% quantiles across all (thinned) MCMC iterations.

Analysis. To determine predicted extinction debt and colonization credit, we first projected the metapopulation model spatially by computing the proportion of simulations in which each species was predicted to be present within each cell of the climate rasters used to calibrate the model. This prediction is a stochastic equilibrium fit from dynamic data and represents the long-term dynamics of the system. We then estimated the present species distributions using the random forest models. This prediction is an instantaneous prediction fit from static data and represents the best estimate of the present distribution of the species. Presence and absence thresholds for each model were determined by maximizing the evaluation statistic (see 'Model evaluation'). The status in each raster cell was then determined by the presence or absence prediction of the two models; continued presence, when both models predicted presence, extinction debt, when the metapopulation model predicted absence at equilibrium, but the random forest predicted presence, and colonization credit for the inverse.

To explore the transient states of the model, we first computed the rate at which patches go extinct when the climate is unsuitable and the rate and which empty patches are colonized when the climate is suitable, defined as:

$$\lambda(E) = \begin{cases} c(E) - m(E), & c(E) > m(E) \\ 0, & c(E) = m(E) \\ m(E) - c(E), & c(E) < m(E) \end{cases}$$

We then simulated the effect of climate change on the range boundary by calculating the partial derivative of this curve with respect to temperature $(\partial \lambda / \partial T)$, evaluated at the range boundary (that is, $\lambda(T) = 0$). This is equivalent to a standard local stability analysis, where the perturbation to the equilibrium is caused by a permanent change in parameters⁴⁴ and is therefore given by linearization of the system around the equilibrium. This quantity, termed the 'responsiveness', yields the change in colonization and extinction of new patches following a small change in temperature at the range limit and indicates the relative rate of movement of the range boundary.

We fit hierarchical linear models to investigate whether there were differences in individual-level mortality and recruitment rates. To perform this analysis, we extracted information on individual trees from the forest inventory database (as before, considering only trees with DBH > 127 mm). Because trees were individually marked during sampling, we were able to track the status of individuals across successive sampling periods. Thus, we were able to determine, for any given sampling interval, the number of trees that survived (that is, were present and alive in both visits), died (present initially and absent or marked as dead in the second visit) and recruited (not present initially and present in the second visit). We then fit two models, one with recruitment probability and one with mortality probability as the response. In both cases, the explanatory variable was an indicator taking a value of 1 if the plot was considered to be subject to extinction debt (see Fig. 2) or 0 if the plot was within the long-term persistence range. Although some plots from the absence and colonization credit portions of the range had observations (owing to the stochastic nature of the system), these plots were excluded for analysis. We included species as a (random) hierarchical grouping factor.

Model evaluation. A large number of plots in the database were sampled only once during the study period. Lacking repeated observations, these data are unsuitable for calibrating the colonization-extinction functions, and were therefore not used in fitting the metapopulation model. However, they represent a large source of information about species distribution that is independent of the data used to calibrate the model. Therefore, we used this dataset (n = 64,848) to evaluate the model predictions for species distributions. For each posterior sample, we computed the conditional prediction of presence or absence of each species using the climatic data for the validation plots. The proportion of presences thus represents the conditional posterior probability that a species *s* is present at equilibrium under climatic regime $\hat{E}: p(X_s = 1 | \hat{E})$. We then used the observed presences and absences in each plot along with the posterior probability of presence to compute the area under the receiver operating curve (ROC), an estimate of classification ability that ranges from 0.5 to 1, where 0.5 is random classification and 1 is perfect classification. This is the most stringent validation test we could think of, as the model was fitted from dynamic data and validated on an integrated prediction about static distribution. Models for 19 out of 21 species performed well ($0.70 < \text{ROC} \le 0.91$), suggesting that the core range for these species is predicted well by the model (Supplemental Table 3).

Data availability. Some data reported in this paper are publicly available via figshare: https://doi.org/10.6084/m9.figshare.4906535. Data not included in this repository are available from the authors upon request.

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Author contributions

M.V.T., D.G., S.V. and I.B. conceived the study, S.V. built the database and M.V.T. wrote the code for the model, MCMC samplers, and ran the analyses. All authors contributed to writing the manuscript.

Additional information

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Correspondence and requests for materials should be addressed to M.V.T.

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Competing interests

The authors declare no competing financial interests.