




Projecting the continental accumulation of alien species through to 2050

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Abstract

Biological invasions have steadily increased over recent centuries. However, we still lack a clear expectation about future trends in alien species numbers. In particular, we do not know whether alien species will continue to accumulate in regional floras

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and faunas, or whether the pace of accumulation will decrease due to the depletion of native source pools. Here, we apply a new model to simulate future numbers of alien species based on estimated sizes of source pools and dynamics of historical invasions, assuming a continuation of processes in the future as observed in the past (a business-as-usual scenario). We first validated performance of different model versions by conducting a back-casting approach, therefore fitting the model to alien species numbers until 1950 and validating predictions on trends from 1950 to 2005. In a second step, we selected the best performing model that provided the most robust predictions to project trajectories of alien species numbers until 2050. Altogether, this resulted in 3,790 stochastic simulation runs for 38 taxon–continent combinations. We provide the first quantitative projections of future trajectories of alien species numbers for seven major taxonomic groups in eight continents, accounting for variation in sampling intensity and uncertainty in projections. Overall, established alien species numbers per continent were predicted to increase from 2005 to 2050 by 36%. Particularly, strong increases were projected for Europe in absolute ($+2,543 \pm 237$ alien species) and relative terms, followed by Temperate Asia ($+1,597 \pm 197$), Northern America ($1,484 \pm 74$) and Southern America ($1,391 \pm 258$). Among individual taxonomic groups, especially strong increases were projected for invertebrates globally. Declining (but still positive) rates were projected only for Australasia. Our projections provide a first baseline for the assessment of future developments of biological invasions, which will help to inform policies to contain the spread of alien species.

KEYWORDS

biodiversity, biological invasions, business-as-usual scenario, future predictions, global, modelling, species richness, trends

1 | INTRODUCTION

Humans are increasingly reshaping global biogeography by transporting species to regions beyond their natural ranges, where they may become permanent additions to local biota (Meyerson & Mooney, 2007; Sax & Gaines, 2008; Turbelin, Malamud, & Francis, 2017). For the last few centuries, numbers of established alien species have increased in many taxonomic groups, and in most cases, so have the rates of species accumulation (Seebens et al., 2017). These trends in biological invasions have been driven particularly by the intensification of trade and transport (Hulme, 2009), land-use change (Pauchard & Alaback, 2004) and access to new source pools (Seebens et al., 2018). Besides being an outcome of globalization, biological invasions are also considered a major driver of global biodiversity change (IPBES, 2019; Pyšek et al., 2020) and have become a defining feature of the Anthropocene (Lewis & Maslin, 2015). Thus, knowledge about future developments in alien species dynamics is of great significance for science and society.

Many studies have considered future trends in biological invasions, though mostly qualitatively (Sax & Gaines, 2008; Walther et al., 2009) or for a limited number of species, using for instance niche modelling approaches (Bellard et al., 2013). A few studies

provide quantitative projections of future invasion dynamics for vascular plants (Bradley et al., 2012; Knapp, Winter, & Klotz, 2017; Rouget et al., 2016; Seebens et al., 2015), insects (Liebhold, Brockerhoff, & Kimberley, 2017) or marine algae (Seebens, Schwartz, Schupp, & Blasius, 2016) based on changes in environmental conditions, dynamics of international trade or assuming a time lag between the actual introduction of species and its subsequent establishment. These studies are, however, restricted to single taxonomic groups or regions. A recent global study addressed future trends in threats from alien species in general based on changes in the drivers of biological invasions rather than the dynamics of alien species themselves (Early et al., 2016). Surprisingly, most previous studies predicting future biological invasions have not considered how alien species numbers have developed over past times (see Knapp et al., 2017; Liebhold et al., 2017; Seebens et al., 2015 for exceptions). Such temporal dynamics are, however, of crucial importance for predictions, as they allow for a general understanding of temporal dynamics as a basis to quantify future dynamics and enable the validation of model results.

The temporal accumulation of alien species varies among taxonomic groups and continents (Seebens et al., 2017). Continents differ in key features relevant to alien species introduction and

establishment: these include evolutionary history and isolation (Fridley & Sax, 2014), prehistoric and modern human colonization (Ellis et al., 2013) and the level of past (di Castri, 1989) and current socio-economic activities that facilitate alien species introduction (Dyer et al., 2017) and establishment (Pyšek et al., 2020). Furthermore, the dynamics of important drivers of alien species introduction and establishment (e.g. trade, human population density and land-use intensity) exhibit considerable variation among continents in recent history (Ellis et al., 2013). The temporal patterns of alien species accumulation may differ among different taxonomic groups on the same continent, as the importance of the underlying mechanisms (e.g. the relevance of different introduction pathways, environmental and habitat characteristics) may vary substantially (Essl et al., 2015). While we are beginning to understand the determinants of current numbers and rates of alien species invasions across space and taxa, comprehensive attempts to analyse and project future dynamics of alien species accumulations across multiple continents and taxonomic groups are largely missing.

Here, we present the first global and cross-taxonomic quantitative projections of alien species numbers for major biogeographical regions (hereafter described as 'continents') through to 2050. The projections and their evaluation are based on the recently established and freely available Alien Species First Records Database (Seebens et al., 2017, 2018). We present a new modelling approach to simulate alien species numbers on continents for different taxonomic groups based on observed long-term historic trends. We developed a modelling approach without the consideration of driver dynamics to be able to apply the same method to all taxonomic groups on all continents considered here. This implies that we assume a continuation of processes in the future as observed in the past. In line with well-known scenario exercises (IPBES, 2016; IPCC, 2014), this can be considered as a business-as-usual scenario. We parameterized and evaluated different model versions by reconstructing the dynamics of alien species numbers between 1950 and 2005. Subsequently, we applied the model to project trends in alien species accumulation for seven taxonomic groups on eight continents up to 2050.

2 | MATERIALS AND METHODS

2.1 | First-records database

The analysis was based on the Alien Species First Records database (Seebens et al., 2017, 2018), which contains first records of alien species that have become established—that is, forming permanent self-sustaining populations (Blackburn et al., 2011)—in one or more mainland or island regions. The regions largely correspond to countries; however, large islands administered politically by a mainland country, but located in biogeographically distinct locations or with a particularly high number of samples are considered as different regions. Each region was assigned to one of eight continents (Antarctica was excluded due to a very low number of first records)

following the classification of the Taxonomic Database Working Group (TDWG) World Geographical Scheme for Recording Plant Distributions Version 2.0 (<https://www.tdwg.org/standards/wgsrpd/>) for region delineation (Figure S1). The first-records database is described in detail in Seebens et al. (2017), was subsequently extended and revised (Seebens et al., 2018) and is available online (<https://doi.org/10.5281/zenodo.3690748>).

We used first records up to the year 2005, as more recent data have been shown to be affected by delays in reporting alien species records into databases (Muñoz-Mas & García-Berthou, 2020; Seebens et al., 2017). We focused on seven major taxonomic groups with the largest number of first records (>1,000), resulting in a dataset of 45,531 first records of 15,628 established alien species across 271 regions worldwide. In this final dataset, most first records are for vascular plants (54% of all records), followed by arthropods other than crustaceans (for simplicity called 'arthropods' hereafter, 28%), birds (6%), fishes (4%), mammals (3%), molluscs (2%) and crustaceans (2%). The geographical distribution of first records is biased towards Europe (38% of all first records), followed by Northern America (16%), Australasia (15%), Southern America (9%), Temperate Asia (9%), Africa (6%), Pacific Islands (5%) and Tropical Asia (2%; Figure S2).

Besides the total number of first records, we additionally extracted the earliest first record of each alien species on a continent. Those species that appeared for the very first time on a continent as alien are called 'emerging alien species' here. This is in contrast to Seebens et al. (2018) where emerging alien species were defined at the global scale. Thus, in the present study, each alien species was considered an emerging alien species at its very first occurrence on a continent. We use the term 'all alien species' to refer to the total set of first records, which includes repeated first records of the same species on the same continent (and of which emerging alien species are a subset).

Long-term dynamics and invasion patterns of alien species may be different for aquatic and terrestrial habitats (Cox & Lima, 2006; Essl et al., 2020). To assess these potential differences, we distinguished between aquatic and terrestrial species. Sensible comparisons between aquatic and terrestrial habitats could only be done for large species groups with members characterized as being either aquatic or terrestrial, which applied only to vascular plants and insects in our dataset. The habitats of aquatic insects have been identified at the family level, whereas those for aquatic plant species have been identified at the species level. Aquatic insects were defined as those species that have at least one developmental stage in water, whereas aquatic plants were defined as species that normally grow to maturity with at least some photosynthetic organs permanently on (floating) or under (submerged) standing or flowing water. Thus, riparian species found only at the margins of waterbodies were not considered as aquatic.

2.2 | Overview of the modelling approach

In what follows, we use the terms 'prediction' and 'simulation' to describe model output in general and 'projection' for trends in future

dynamics, which in our case refers to the time after 2005. Our model estimates changes in the number of emerging alien species over time. More precisely, the model simulates the years of first record of alien populations that have been recorded as established in the region. This implies that the model does not cover dynamics of introduction, establishment or detection before the actual first record. For instance, local population dynamics, Allee effects or time lags cannot be considered in our approach. Simulations are based on two input variables: the size of source pools (i.e. the 'candidate species pools', see below) and the first-record rates of all alien species (i.e. the number of first records per year). Thus, projections can be considered as extrapolations of alien species first records constrained by the finite size of source pools.

The 'candidate species pool' represents a pool of species, which have the potential to become an alien species on the respective continent. It does not necessarily include all species native outside the respective continent, because not all species are likely to become alien somewhere. Characteristics of the candidate species pool such as size or abundance distribution are unknown. However, knowledge about the candidate species pools is essential for the quantification of future dynamics, as the finite size of these pools constrains the total number of emerging alien species and thus provides an upper limit to alien species numbers (Liebhold et al., 2017). Here, we estimated the size of candidate species pools as follows: The candidate species pools are specific for each combination of a continent and a taxonomic group. The species pool has size X , which may be either constant or variable in time t , denoted then as X_t . Initially, X is unknown and thus estimated from observed long-term trends of emerging and all alien species numbers following Seebens et al. (2018). The basic idea behind this approach is that the size of the candidate species pool is a strong determinant of the rate by which emerging alien species are sampled from this pool. For instance, when repeatedly sampling individuals from a large candidate species pool, chances are high to select a new species not already present in the focal region. This would result in high rates of emerging alien species even after a long time (more precisely, after many draws from the pool) and a slow depletion of the candidate species pool. In contrast, only a few species could establish when the candidate species pool is small, which would then result in a rapid decline in the rate of emerging alien species (fast depletion). Similar approaches have been applied to estimate the total number of species on Earth from time series of species descriptions (Mora, Tittensor, Adl, Simpson, & Worm, 2011). Abundances of the species in the source pool are assumed to be log-normally distributed with a fixed mean ($\log \mu = 0$) and variance ($\log \sigma^2 = 1$), which is a common way to describe the distribution of species abundances in natural communities (Bell, 2000; Liebhold et al., 2017). Modifying these parameter values has little influence on projections (Figure S3) as long as the full range of the distribution is sampled, as we do here.

The model simulates changes in the number of emerging alien species of a certain taxonomic group on a continent by randomly selecting individuals from the candidate species pool with size X at a rate in time t , which was obtained from the observed first-record

rates of all alien species of that group, continent and time. The selected individuals are then added to the alien species pool of the particular continent and taxonomic group. Once the simulation finishes in the year 2005, the rate of emerging alien species can be computed for the full time series. The model can be considered as a process of resampling with replacement, where individuals are randomly selected from a log-normal distribution of X species. The only parameter of the basic model is X . In a refined model version, X_t was allowed to change in time. For simplicity, X_t was considered to be a stepwise linear function of moderate complexity, but high flexibility. We selected a function with three linear segments of independent slopes, which has proven to capture observed dynamics very well (Seebens et al., 2018). The flexibility of such a function is much higher compared to, for example, exponential functions as it covers shapes ranging from constant to linear, near-exponential and various other nonlinear forms. Another advantage is that the final shape of the function is determined by the data and the fitting process (see below) rather than pre-defined by the user. Both slopes and position of inflection points along t were estimated by model fitting. The use of three linear segments with a fixed start and end date requires knowledge about the timing of two inflection points and four size values, which altogether resulted in six parameters describing X_t for the refined model version.

2.3 | Modelling steps

In brief, our approach consists of the following data manipulation and modelling steps (Figure 1): (1) correction of first records to account for temporal variation in sampling intensity; (2) identification of the rate of emerging alien species from the first-records dataset; (3) fitting the model to the observed rates of emerging alien species, thereby estimating the size of candidate species pools; (4) future extrapolation of model input variables, namely the estimated sizes of candidate species pools and the first-record rates and (5) projection of the accumulation of emerging alien species constrained by the size of the candidate species pools. As steps 1 and 3 include stochastic elements (see below), all modelling steps 1–5 were repeated 100 times for each taxon–continent combination; this allowed us to explore the full range of predicted future trajectories and to assess the uncertainty inherent in these projections. In more detail, the procedure applied was as follows:

2.3.1 | Step 1 (bias-correction)

Time series of first records are likely biased by temporal variation in sampling intensity, which may affect model predictions. As documentation of sampling intensity is largely missing, we define a time series of likely changes in sampling intensity, which can be used to explore variations of results due to a temporal sampling bias following Seebens et al. (2018). In this approach, the year of each first record was shifted to earlier years according to a presumed recording

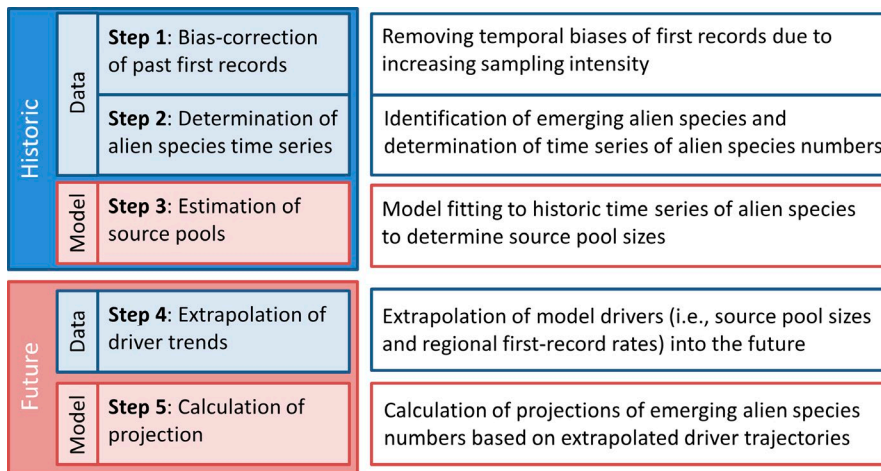


FIGURE 1 Overview of data manipulation and modelling steps to generate future projections (left column) and brief explanations (right column). The applied approach consists of two parts, involving historic and future dynamics, respectively. In each part, time series of model input variables were first generated (indicated by 'Data') and then used for modelling ('Model') to either fit the model to historic time series and determine parameter values ('Historic') or to calculate future projections ('Future')

lag between actual introduction and recording of that species in a given region. Due to missing data, we had to make assumptions about the temporal development of sampling intensity: we assumed that the size of the time lag is a direct result of sampling intensity, and we further assumed an exponentially increasing sampling intensity since 1500 ranging from a time lag of a maximum of 100 years at 1500 to 5 years in 2005 due to increasing effort and efficiency in sampling species. This assumption is related to findings of studies on native species description rates (Appeltans et al., 2012; Costello, Wilson, & Houlding, 2012). This correction resulted in a shift in the year of first record to earlier dates and in a less steep increase in rates of first records in recent decades (Figure S4), and thus provides more conservative predictions of future trajectories. Even without actual data on changes in sampling intensities, we believe such a bias-correction provides more realistic results compared to an approach ignoring temporal biases.

2.3.2 | Step 2 (extraction of first-record rates)

From the bias-corrected time series, first records of emerging alien species were extracted. The numbers of records were aggregated over time intervals of 5 years to reduce the number of gaps and the amount of noise, which is particularly problematic for taxonomic groups with few records such as mammals. The number of first records per time interval is referred to as the first-record rate.

2.3.3 | Step 3 (estimation of source pools)

The model was fitted to the observed time series of emerging alien species. This was done by modifying X in such a way that predictions best match the observed changes in rates of emerging alien species. Deviations between observed and simulated rates were measured using the root mean squared error (RSME) between observed and predicted values. Finding the optimal value of X , which minimized RSME, was done using the Nelder–Mead optimization algorithm implemented in the R software package 'optim' (R Core Team, 2019).

For the model version considering temporal changes in X_t , the six parameters defining X_t were estimated. As it is a stochastic model, only the mean of 100 single runs was used for fitting to avoid extreme trajectories. Further details of the modelling approach can be found in Seebens et al. (2018).

2.3.4 | Step 4 (extrapolation of model input variables)

For projections, input variables of the model (i.e. the size of the candidate species pool and the first-record rate of all alien species) have to be extrapolated into the range of projections. As X or X_t are either constant or linear functions, the size of the candidate species pool was linearly extrapolated (Figure S5), while a more flexible approach was used for the first-record rates to capture their more dynamic nature. We tested six different approaches for extrapolation, which were fitted to each taxon–continent combination separately: a linear [$y = a + bt$], exponential [$y = ae^{bt}$], power law [$y = at^b$], saturating [$y = a(1 - e^{-bt})$], sigmoidal [$y = a(x^b / (t^b + c^b))$] or hump-shaped (Weibull) [$a(c/b * (t/b)^c e^{-(t/b)^c})$] function of first-record rates y with time t , and a , b and c denoting parameters defining the shape and scale of the functions (see Figure S5 for examples). The functions were fitted individually to the time series of first-record rates from 1800 until 1950 or 2005, respectively, using the Nelder–Mead optimization algorithm. The best function was determined as the one with the lowest AIC. In some cases, different functions performed equally well with a $\Delta AIC < 5$ (Burnham & Anderson, 2004). To avoid unnecessary complexity and less robust projections, simpler functions were preferred over more complex ones in these cases. That is, from the set of best-fitting functions ($\Delta AIC < 5$), linear functions were preferred over nonlinear ones and monotonic functions over hump-shaped ones. In addition, we restricted the choice of the hump-shaped function to cases where the time series of first-record rates declined in recent times to avoid false predictions of declining trends due to noisy data. This was tested by additionally fitting a linear function to the last five time intervals (1985–2005). If the slope of this

fit was negative, the selection of the hump-shaped function was accepted. In cases where the slope was positive, the second-best fit was chosen. The simplest of the best-fitting functions thus obtained was used for extrapolation.

2.3.5 | Step 5 (calculation of quantitative projections)

Projections of emerging alien species accumulation were obtained by performing simulations using the candidate species pool sizes obtained from step 3 and the extrapolated time series of first records and candidate species pool sizes from step 4. Since the first-records database does not include first records for all established alien species, we applied a correction for spatial variation in sampling intensity among continents using available databases of established alien species as references (vascular plants, Pyšek et al., 2017; birds, Dyer et al., 2017; fishes, Froese & Pauly, 2015; no databases were available for mammals or invertebrates). This was done by applying a correction factor to simulation results to achieve the same number of emerging alien species on the respective continent in the year 2005 as reported by these databases.

These five steps were repeated 100 times for each taxon-continent combination to assess the uncertainty inherent in this approach. Projections were sensitive to a low number of first records and a low variation in the time series (e.g. constant time series). We therefore defined a set of minimum requirements of a time series to run the model: Projections were only calculated for first-record time series of taxon-continent combinations with at least 15 values, a mean of at least two emerging alien species per time interval and a standard deviation of at least one. Consequently, projections could only be accomplished for 38 out of 56 taxon-continent combinations (68%). Due to data limitations, analyses of dynamics on subsets of the data, such as islands-mainland and aquatic-terrestrial comparisons, could only be conducted at the global scale.

2.4 | Predicting invasion dynamics

To validate model predictions, we performed a back-casting exercise to evaluate model performances by predicting historical dynamics of emerging alien species numbers. For this purpose, the model was fitted to emerging alien species numbers until 1950 to predict species numbers for the period 1950–2005. Observed and predicted numbers during the latter time period were compared to assess the reliability of predictions. The comparison was done using the relative absolute error (RAE). The RAE is a measure of improvement of the model compared to using a single constant rather than the more complex model, with a value of 1 or larger denoting no improvement. In addition, the reliability of predictions for each taxon-continent combination was assessed. We excluded taxon-continent combinations with an RAE > 1 for the calculation of projections. We made an exception to this rule for mammals, as the observed first-record

rates of mammals changed at around 1950 from an increasing to a decreasing trend, which is difficult to predict using our approach. Note that an RAE of 1 does not necessarily imply poor model performance as the trends in emerging alien species numbers can be close to constant.

In a sensitivity analysis, we tested different versions of the model and data-manipulation steps of varying complexity to assess their influences on model results. As mentioned above, the size of the candidate species pool was either set constant or allowed to vary through time. In an alternative approach, first-record rates were assumed to remain constant in the future. For this, we extrapolated the mean value of first-record rates observed during the last 50 years into the future. Furthermore, we tested the influence of different bin sizes (1, 3, 5, 10 and 15 years), used to aggregate first records (step 2), on model results. In the selection of the optimal bin size, we had to find a compromise between the number of observations and the variability of the time series. Using a large bin size of, for example, >10 years would have resulted in a low number of observations reducing the chance of convergence of the fit algorithm. In contrast, using a low bin size of, for example, <5 years may result in noisy time series or time series with very low variability (e.g. constant), which are difficult to fit as well. For the selection of the optimal bin size, model performance was therefore also assessed by the number of successful model runs, constrained by our minimum requirement for a time series (step 5). Finally, we tested whether bias correction using a linear versus an exponential increase in sampling intensity (see step 1) influenced model results.

After the identification of the model version that best described past observed invasion dynamics, the modelling steps were applied to first-record rates over the whole time series until 2005 to project trajectories of emerging alien species until 2050.

3 | RESULTS

In a sensitivity analysis, we validated the performance of different model versions in our back-casting exercise by predicting emerging alien species numbers during 1950–2005, which revealed that model performance varied considerably among model versions, regions and taxa. For most model versions, only a few taxon-continent combinations provided a reasonable performance with an RAE lower than 0.75 (Figure S6). For instance, only 16% of all simulations using the model with a constant candidate species pool and a non-constant extrapolation of first-record rates (Figure S6c) reached an RAE < 0.75, while this was the case for 40% of the simulations using the best performing model (Figure S6a). Across all combinations, the poorest performance was obtained for the two model versions assuming a constant candidate species pool size, with a mean RAE of 0.9 ± 0.18 (standard deviation calculated over all taxon-continent combination) and 0.88 ± 0.2 , respectively. Allowing for changes in the size of the candidate species pool and a constant extrapolation of first-record rates clearly improved performance (RAE = 0.78 ± 0.22). Best

model performance was obtained when allowing for dynamic (rather than constant) future developments of both candidate species pools and first-record rates (Figure S6a): that is, when the sizes of candidate species pools were allowed to change in time (increase or decrease) and first-record rates were predicted using one of the set of different functional forms (linear or nonlinear) described in Section 2 (step 4). In addition, we tested the influence of varying parameter values of the log-normal distribution, which determines the shape of the candidate species pool (step 3), on model performance. Overall, selecting a mean of the distribution of around zero, as done here, provided the best model performance, albeit the differences to other parameter values were low (Figure S3). Furthermore, testing the influence of different bin sizes used to aggregate first records (other than 5 years; step 2) revealed that model performance as well as the number of successful model runs clearly decreased using both smaller bin sizes (1 or 3 years) and larger bin sizes (10 or 15 years; Figure S7). In particular, using bin sizes of 1 year (no aggregation) or 15 years resulted in a very low number of successful model runs, which is due to the fact that variability of time series was too low (i.e. constant time series) and/or the number of observations was too low, respectively. Differences in using 5 or 10 year bin sizes were minor and in some cases, model performance even improved using a bin size of 10 years. However, for reasons of consistency, we selected a bin size of 5 years for all model runs as this provided the best compromise across all taxon–continent combinations.

The best model performance was obtained using the model version with a nonlinear development of candidate species pools, flexible extrapolations of first-record rates and a temporal resolution of 5 years ($RAE = 0.78 \pm 0.22$; Figure S6). We removed two taxon–continent combinations (fishes and molluscs in Australasia) with low numbers of successful simulation runs ($n < 15$). Using this configuration, we obtained projections of emerging alien species numbers (i.e. number of earliest first records of species on a continent, see Section 2) through 2050 for a total of 3,790 simulation runs for 38 taxon–continent combinations (99.7 simulations each on average). Model validation revealed that in general consistently good model fits ($RAE < 0.5$) were obtained for projections for the European continent, vascular plants and to some degree for arthropods, while poorer fits with RAE often between 1 and 0.75 were gained for mammals and birds (Figure 2).

Mean projections of future developments of emerging alien species numbers roughly fell into three groups: (a) a steep increase, as shown, for instance, for many European trajectories; (b) a low increase, as shown for arthropods in Australasia; or (c) saturation, as shown for fishes and vascular plants on Pacific Islands (Figure 3). In general, variation in the mean projections was comparatively low for taxonomic groups with high species numbers, such as vascular plants and arthropods, and high for those with low numbers, such as vertebrates. For mammals in particular, the range of obtained trajectories was large, which indicates a high uncertainty in these projections. To account for large-scale spatial variation in sampling intensity, projections were adjusted to reach the same level of species numbers as reported in the literature in recent years. This adjustment, which could only be performed for vascular plants, birds

and fishes due to the lack of comprehensive alien species data for other taxonomic groups, resulted in a more uniform picture for most continents (compare Figure 3 and Figure S8). That is, after accounting for spatial variation in sampling intensities, trajectories for less well-sampled regions such as Tropical Asia increased distinctly and revealed similar shapes of alien species increases as neighbouring regions such as Temperate Asia and the Pacific Islands.

According to our model, numbers of emerging alien species on a continent were predicted to increase from the year 2005 to 2050 by 36% on average but with a large variation across all taxonomic groups and continents (Tables 1 and 2). The total number of emerging alien species on a continent was projected to increase by 1,195 species averaged over projections of those taxonomic groups, which are available for most continents to enable a fair comparison (vascular plants, fishes, birds and arthropods; Table 1). Highest increases can be expected for Europe in both relative and absolute terms (64% , $2,543 \pm 237$ alien species) followed by Temperate Asia (50% , $1,597 \pm 197$), Northern America (23% , $1,484 \pm 74$) and Southern America (49% , $1,391 \pm 258$). Lowest increases in relative values were projected for Australasia (16%), while in absolute terms this was the case for the Pacific Islands ($+132$ species). In a more detailed analysis, projected relative increases in numbers of emerging alien species

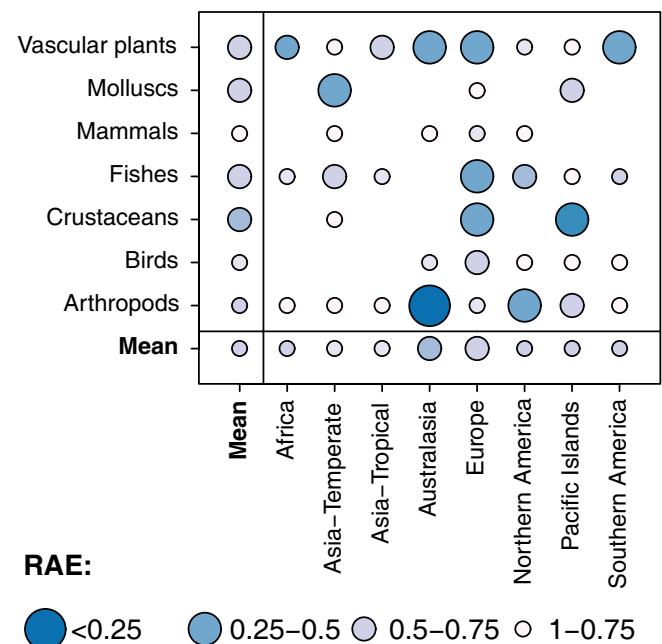


FIGURE 2 Performance of the best-fitting model for all taxonomic groups and continents. Performance was tested by parameterizing the model with first records until 1950 to predict dynamics from 1950 to 2005. Model performance was measured using the relative absolute error (RAE) between observed and predicted rates of emerging alien species. Size of bubbles and colours indicate the median RAE averaged over 100 model runs for each combination of taxa and continent. Lower values of RAE denote better model performance. Gaps indicate taxon–continent combinations for which the resulting time series of first records did not fulfil our minimum requirements for simulations (step 5 in Section 2), and thus projections could not be calculated

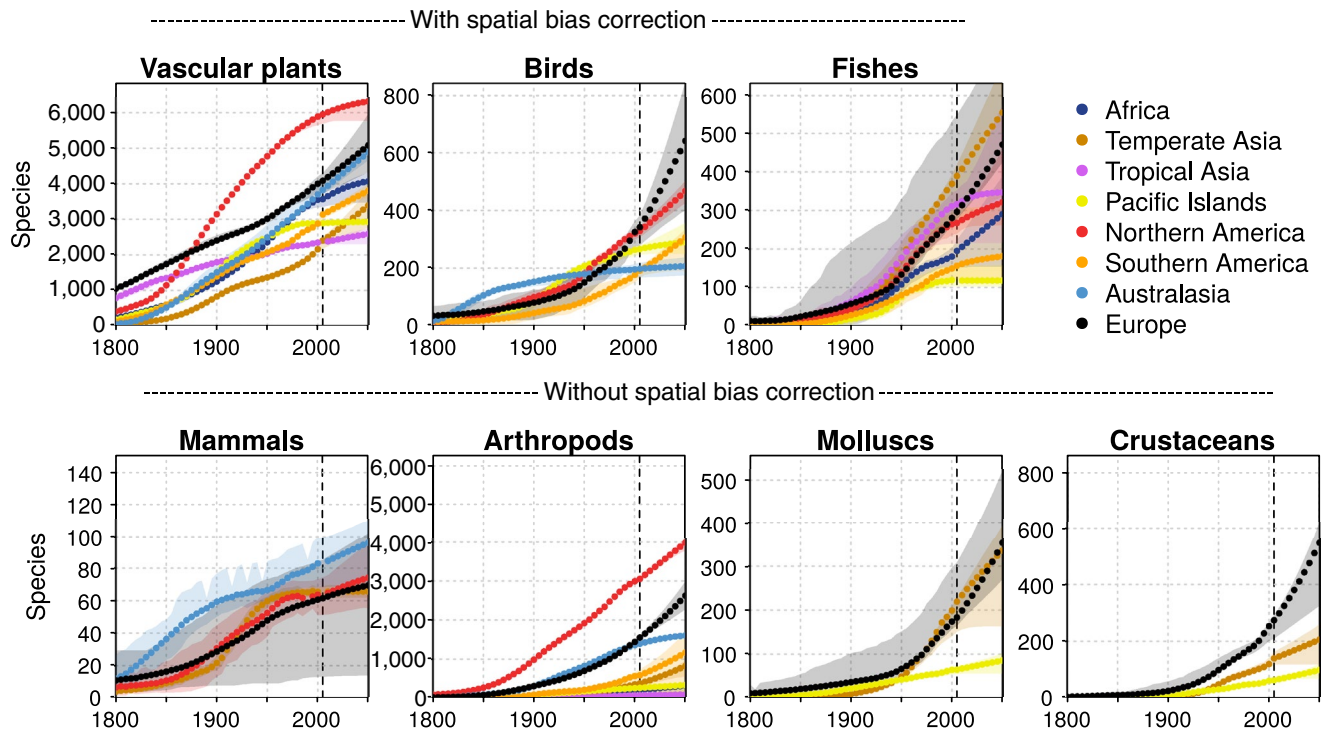


FIGURE 3 Predicted developments of alien species numbers on different continents for seven taxonomic groups until 2050. The dots represent means of up to 100 model runs, while the full range of all predicted trajectories is indicated by shaded areas. To account for spatial variation in sampling intensity among continents, trajectories are corrected to have the same value of alien species numbers in 2000 as reported in recent publications for vascular plants, birds and fishes. This was not possible for mammals and invertebrates. Uncorrected trajectories are shown in Figure S8. See Section 2 for more information

ranged from 0% (fishes on Pacific Islands and mammals in Temperate Asia) to 117% (arthropods in Temperate Asia; Table 2). Invertebrates showed the highest relative increases. In absolute terms, highest numbers of alien species among all continents were expected for Europe (for birds, fishes, arthropods, molluscs and crustaceans), Australasia (mammals) and Temperate Asia (vascular plants) by 2050 (Figure 3; Table 2). For arthropods, high increases in absolute terms were also predicted for Northern America. This was not only due to the already high numbers of alien species on these continents but also due to the steep rises in alien species numbers projected up to 2050 (Figure S9).

Predicted trajectories of alien species numbers were surprisingly similar for both mainland and island regions across taxonomic groups (Figure S10). While numbers of predicted alien species were generally higher in mainland regions, those for islands were at a similar magnitude. Trends in alien species numbers are predicted to show further increases across all taxonomic groups, albeit less clearly for mammals in mainland regions, with a tendency towards steeper increases in mainland regions. The comparison of trajectories in aquatic and terrestrial habitats for vascular plants and insects revealed that total alien species numbers are much lower in aquatic habitats (Figure S11). Trends of alien species numbers are predicted to distinctly increase further, with higher relative increases predicted for aquatic vascular plants and terrestrial insects. However, due to the lower number of aquatic alien species in these two taxonomic groups, the variation around the mean is very high and predictions for aquatic alien species are less robust compared to those of terrestrial alien species.

TABLE 1 Projected relative and absolute increases in emerging alien species numbers from 2005 to 2050 across taxonomic groups for each continent. Relative increases represent mean relative increases and range of increases across taxonomic groups, while absolute increases denote sums of emerging alien species numbers and standard deviations across taxonomic groups

Continent	Relative increase 2005–2050 (%)	Absolute increase ^a 2005–2050
Africa	39 (14, 51)	767 ± 133
Temperate Asia	50 (0, 117)	1,597 ± 197
Tropical Asia	30 (10, 67)	360 ± 78
Australasia	16 (5, 28)	1,286 ± 44
Europe	64 (13, 100)	2,543 ± 237
Northern America	23 (6, 42)	1,484 ± 74
Pacific Islands	21 (0, 56)	132 ± 29
Southern America	49 (16, 99)	1,391 ± 258
Average	36 (0, 117)	1,195 ± 131

^aAbsolute increases were calculated only including projections of those taxonomic groups that were available for most continents (i.e. vascular plants, fishes, birds and arthropods) to avoid biases towards continents with a high number of projections.

To evaluate changes in the dynamics of alien species numbers, we calculated increases over the same period of time (45 years) before and after the end of reported first records (2005). A comparison

TABLE 2 Projected relative (%) increases in numbers of emerging alien species in the years 2005–2050. Values represent means over up to 100 individual runs and 0.025% and 0.975% quantiles of 100 simulation runs in square brackets. Mean absolute increases in alien species numbers together with standard deviations across individual simulation runs are provided in parentheses. Grey-shaded numbers represents predictions corrected for spatial variation in sampling intensity among continents

Taxon	Africa	Temperate Asia	Tropical Asia	Australasia	Europe	Northern America	Pacific Islands	Southern America
Vascular plants	14 [4, 19] (503 ± 113)	41 [28, 54] (987 ± 170)	10 [0, 17] (227 ± 67)	28 [22, 29] (1,065 ± 41)	24 [16, 39] (997 ± 209)	6 [1, 7] (365 ± 33)	1 [0, 2] (38 ± 9)	21 [18, 25] (669 ± 52)
Mammals								
Birds	42 [12, 67] (59 ± 26)		67 [40, 86] (78 ± 15)	5 [1, 8] (9 ± 4)	88 [61, 120] (299 ± 53)	42 [35, 46] (138 ± 11)	9 [2, 26] (24 ± 22)	60 [22, 69] (115 ± 20)
Fishes	49 [7, 73] (96 ± 39)	42 [12, 56] (165 ± 48)	10 [1, 42] (31 ± 34)		59 [52, 87] (175 ± 32)	20 [3, 61] (54 ± 57)	0 [0, 0] (0 ± 0)	16 [2, 86] (25 ± 39)
Arthropods	51 [0, 69] (109 ± 51)	117 [80, 141] (445 ± 87)	35 [0, 53] (24 ± 13)	15 [13, 17] (212 ± 14)	69 [65, 84] (1,072 ± 92)	30 [29, 32] (927 ± 31)	26 [13, 32] (70 ± 17)	99 [15, 124] (582 ± 249)
Molluscs		53 [5, 69] (116 ± 40)			93 [69, 131] (170 ± 31)		32 [2, 43] (21 ± 7)	
Crustaceans		47 [27, 71] (66 ± 18)			100 [77, 112] (273 ± 34)		56 [37, 82] (36 ± 8)	

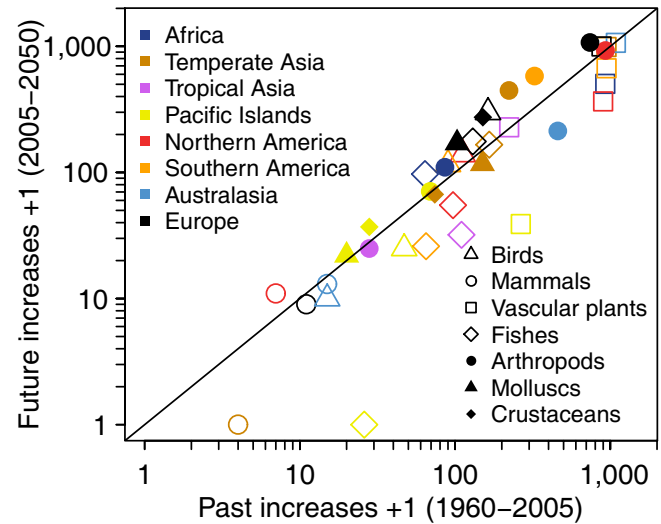


FIGURE 4 Comparison of observed past (1960–2005) and projected (2005–2050) increases in alien species numbers for all combinations of continents and taxonomic groups. Values above the 1:1 line indicate a larger increase in alien species numbers projected for the period 2005–2050 than observed between 1960 and 2000, while values below the line denote a lower increase in alien species numbers in the future than observed in the past. Alien species numbers plus one is shown to include zeros in the log–log plot as well. Hence, values at one actually indicate zero change in alien species numbers

of past (1960–2005) and future (2005–2050) increases in alien species numbers revealed that the rates of emerging alien species were projected to accelerate particularly for arthropods worldwide but also for birds (Figure 4). In contrast, rates were projected to decline in the future for mammals and fishes. Among continents, a uniquely accelerating rise in alien species numbers was projected for Europe, where rates of emerging alien species were expected to increase in all taxonomic groups except mammals (Figure 4). In Australasia, projected increases in alien species numbers were consistently lower than in the past, indicating a consistent decline in the rate of emerging alien species.

The sizes of the candidate species pools were predicted to increase for all taxa and continents with two exceptions: mammals in Temperate Asia and birds in Australasia (Figure S12). Particularly strong increases since the early 20th century were simulated in nearly all cases. However, the trajectories of the sizes of candidate species pools of individual simulation runs may differ widely (see large bands around the mean in Figure S12). The predicted sizes of candidate species pools ranged from around 800 species of mammals to over 100,000 species for arthropods (Figure S12).

4 | DISCUSSION

This study provides the first quantitative estimates of future alien species accumulation trajectories at a global scale, resolved to continents and major taxonomic groups and based on historic information on alien species trends. Our projections indicate that

by the mid-21st century there will be distinct increases in alien species numbers particularly for Europe but also for Temperate Asia and Northern America, and for invertebrates in all regions (Table 2). Our results suggest that the observed past trends of biological invasions will continue to accelerate also in the next decades for many taxonomic groups and continents. Likewise, the increases in global candidate species pools in the past are likely to continue into the future, resulting in the emergence of more new alien species.

The highest increases, both in absolute and relative terms, were predicted for Europe, the continent with arguably the longest history of alien species recording. Indeed, comparing the number of first records of alien vascular plants with published alien species numbers (van Kleunen et al., 2015) revealed that first records seem to be most complete for Europe (a total of 3,895 alien vascular plant species for Europe in the first-record database compared to known 4,140 alien vascular plants in this continent, 94%). The proportion is often much lower for other continents, such as Northern America (43% coverage), Temperate Asia (45%) or Africa (26%), which may affect model predictions. Accounting for spatial variation in sampling intensities increased alien species numbers particularly for less sampled continents such as Tropical Asia, Southern America and Africa, resulting in a more balanced picture with similar trajectories among continents (compare Figure 3 and Figure S8). Due to the lack of publicly available global databases, this correction could only be applied to alien vascular plants, birds and fishes, but it seems reasonable to expect similar results for other mammals and invertebrates as well. Thus, the low alien species numbers observed, for example, for mammals in Southern America or arthropods in Asia are likely an effect of low sample sizes for those continents and taxonomic groups in the first-records database. True numbers of alien species can be expected to be higher. In contrast to absolute values, relative increases and the shape of the accumulation curves can be regarded as more robust to spatial variation in sampling intensities since even distinct modifications of the first records resulted in only slight modifications of the time series (Figure S4).

The predicted increases in alien species numbers were consistent across different geographical units such as islands and mainland regions or across habitats such as aquatic and terrestrial (Figures S10 and S11). For islands, it has been shown that both the total number of alien species and the first-record rate are comparable to those in mainland regions, although usually a bit lower (Seebens et al., 2017), which is confirmed by our projections. This is remarkable as the area of mainland regions in our database is 12 times larger than the area covered by islands. Although alien species are certainly not equally distributed across regions, this imbalance further emphasizes the high numbers of alien species observed on islands compared to mainland regions (Dawson et al., 2017). However, our simulations indicate that the future rise in the number of alien species will be greater for mainland regions than for islands, as a consequence of the observed recent accelerations in mainland alien species numbers (Figure S10). This acceleration will lead to larger differences in numbers of alien species between islands and mainland regions in the

future. Nevertheless, islands should still be viewed as being particularly vulnerable to the introduction of alien species (Russell, Meyer, Holmes, & Pagad, 2017), with major drivers in the future being trade, transport, tourism, land-use changes and climate change (Lenzner et al., 2020).

Absolute and relative accumulation rates provide complementary insights into future trajectories of alien species accumulation (Figure 3; Figure S9). While the former are sensitive to characteristics of continents such as geographical size, human population size or economic growth and to the level of recording of alien species (the better historical recording has been and the more records are in our database, the higher the projected absolute number of new established aliens, given a particular trajectory), relative accumulations are less sensitive to these confounding factors and provide insights into the relative increases of alien species compared to the status quo. On the other hand, the relative increase in alien species numbers does not indicate if these increases are based on small or large absolute numbers of alien species currently present in a continent.

The predicted rises in alien species numbers may not come as a surprise, given the continuous increases observed during recent centuries, and the general lack of indications of slowdowns worldwide (Aukema et al., 2010; Blackburn, Dyer, Su, & Cassey, 2015; Liebhold et al., 2017; Muñoz-Mas & García-Berthou, 2020; Seebens et al., 2017). Furthermore, major drivers of alien species introductions and establishment are predicted to rise as well, with clear implications for biological invasions. In a recent expert-based assessment on future dynamics of drivers of biological invasions, trade and transport are assumed to highly likely play a dominant role in driving future increases in alien species numbers, even in the best case scenario considering a human society addressing threats to biodiversity appropriately (Essl et al., 2020). Other drivers such as climate change, biodiversity loss, land-use change and human migration are assumed to become more dominant in the less optimistic scenario (Essl et al., 2020). All of these drivers are predicted to intensify in the future as well, increasing the probability of introduction and establishment. Countermeasures would need to be put in place to alter these trajectories, and indeed legislations relating to alien species have increased in number during recent decades (Turbelin et al., 2017). However, the capacity of most countries to proactively counter the rising tide of alien species is still poor for most regions worldwide (Early et al., 2016), and it seems likely that this will not change substantially in the near future. Overall, numbers of alien species will very likely increase as predicted by our model.

Projecting alien species accumulation rates into the future is subject to a number of caveats. We have explicitly accounted for several of the most important factors (e.g. the size of the candidate pool of potential alien species, recording bias and some sources of uncertainty), but two main limitations remain:

First, the statistical extrapolations assume that the patterns observed for past alien species accumulation will continue in the future. Thus, potential sudden changes in the underlying dynamics

driving alien species introductions were ignored. Examples of such potential changes are the implementation of new mitigation strategies (Albert, Lishman, & Saxena, 2013), strengthening of biosecurity regulations such as those now implemented in Australia and New Zealand (Sikes et al., 2018), changes in the global transportation network (e.g. use of the standardised intermodal container since the 1950s, Cudahy, 2006), accelerated climate change and changes in land use (such as increased cultivation of biofuels). The consequences of these types of changes for biological invasions are difficult to project, even if the respective drivers could explicitly be incorporated in the model. Assuming a continuation of past dynamics into the future seemed to be a strong assumption, but it is a standard practice in well-known scenario-based assessments of future dynamics such as for climate change (IPCC, 2014) or biodiversity loss (IPBES, 2016). Accordingly, our results can be regarded as a business-as-usual scenario, where distinct shifts in underlying drivers are not considered. For such a scenario, we are confident that the calculated trajectories provide a reasonable baseline for the exploration of future alien species accumulations under diverging future scenario assumptions.

Second, projections of alien species numbers were calculated without explicit information on underlying drivers except the size and dynamics of the candidate species pools. Data on many drivers of biological invasions are currently lacking for the period before the last 50–100 years and not consistently available for all taxonomic groups. An integration of driver dynamics into the model would have restricted the analysis to a limited time span and to just a few drivers with sufficient historical data (Early et al., 2016; Seebens et al., 2018), resulting in a higher risk of errors in the overall trends used for extrapolation of first-record rates into the future. In addition, observed trends of emerging alien species numbers during the 20th century were surprisingly stable despite distinct political and socio-economic changes during that time (Figure S4). The reason may be the coarse spatial and temporal resolution of our analysis, which reduces much of the variation observed on finer scales.

Our approach takes advantage of the long time series currently available for first records. We see our model approach as complementary to more mechanistic approaches and as an important baseline for comparison to get confidence in projections of alien species dynamics. However, mechanistic model approaches on a comparable scale are currently lacking and still need to be developed. Thus, this study provides an important first step towards thorough quantifications of future dynamics of biological invasions (Lenzner et al., 2019).

5 | CONCLUSIONS

Our analyses indicate that for multiple taxonomic groups on most continents, alien species numbers will continue to increase—often at accelerating rates—in the next three decades (Figures 3 and 4). We show that although the absolute values of predicted alien species numbers may be uncertain, the continental-scale trends are

robust. The overall results may not be entirely surprising given the continuous increase in alien species numbers observed during the last decades (Seebens et al., 2017), but we here provide a first quantitative baseline for the assessment of how biological invasions may unfold in the future. The availability of such a baseline enables comparisons of scenario projections of potential future dynamics (Lenzner et al., 2019) and can help to direct management strategies efficiently to mitigate the spread of alien species. The predicted increases in alien species numbers would be expected to slow down with the application of additional regulations aimed at preventing alien species incursions. Indeed, it has been repeatedly shown that the implementation of targeted biosecurity efforts can reduce the numbers of new alien species becoming established (Bacon, Bacher, & Aebi, 2012; Leung, Springborn, Turner, & Brockerhoff, 2014; Sikes et al., 2018). However, a significant decrease in rates of alien species numbers on a large scale can only be achieved by a coordinated attempt across political borders.

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AUTHOR CONTRIBUTION

H.S. designed and implemented the model, conducted the analysis and wrote the initial draft of the manuscript together with F.E. Information about aquatic species was provided by Z.P. and A.M.L. All authors contributed to the overall study design and paper writing.

DATA AVAILABILITY STATEMENT

Data of first records are provided at <https://doi.org/10.5281/zenodo.3690748>.

Code availability statement: Computer code to reproduce results of this study is provided at <https://doi.org/10.5281/zenodo.3690797>.

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REFERENCES

- Albert, R. J., Lishman, J. M., & Saxena, J. R. (2013). Ballast water regulations and the move toward concentration-based numeric discharge limits. *Ecological Applications*, 23(2), 289–300. <https://doi.org/10.1890/12-0669.1>
- Appeltans, W., Ah Yong, S. T., Anderson, G., Angel, M. V., Artois, T., Bailly, N., ... Costello, M. J. (2012). The magnitude of global marine species diversity. *Current Biology*, 22(23), 2189–2202. <https://doi.org/10.1016/j.cub.2012.09.036>
- Aukema, J. E., Cullough, D. G. M., Holle, B. V., Liebhold, A. M., Britton, K., & Frankel, S. J. (2010). Historical accumulation of nonindigenous forest pests in the continental United States. *BioScience*, 60(11), 886–897. <https://doi.org/10.1525/bio.2010.60.11.5>
- Bacon, S. J., Bacher, S., & Aebi, A. (2012). Gaps in border controls are related to quarantine alien insect invasions in Europe. *PLoS One*, 7(10), e47689. <https://doi.org/10.1371/journal.pone.0047689>
- Bell, C. (2000). The distribution of abundance in neutral communities. *The American Naturalist*, 155(5), 606. <https://doi.org/10.2307/3078983>
- Bellard, C., Thuiller, W., Leroy, B., Genovesi, P., Bakkenes, M., & Courchamp, F. (2013). Will climate change promote future invasions? *Global Change Biology*, 19(12), 3740–3748. <https://doi.org/10.1111/gcb.12344>
- Blackburn, T. M., Dyer, E. E., Su, S., & Cassey, P. (2015). Long after the event, or four things we (should) know about bird invasions. *Journal of Ornithology*, 156(1), 15–25. <https://doi.org/10.1007/s10336-015-1155-z>
- Blackburn, T. M., Pyšek, P., Bacher, S., Carlton, J. T., Duncan, R. P., Jarošík, V., ... Richardson, D. M. (2011). A proposed unified framework for biological invasions. *Trends in Ecology & Evolution*, 26(7), 333–339. <https://doi.org/10.1016/j.tree.2011.03.023>
- Bradley, B. A., Blumenthal, D. M., Early, R., Grosholz, E. D., Lawler, J. J., Miller, L. P., ... Olden, J. D. (2012). Global change, global trade, and the next wave of plant invasions. *Frontiers in Ecology and the Environment*, 10(1), 20–28. <https://doi.org/10.1890/110145>
- Burnham, K. P., & Anderson, D. R. (2004). Model selection and inference – A practical information-theoretic approach. *Sociological Methods & Research*, 33(2), 261–304.
- Costello, M. J., Wilson, S., & Houlding, B. (2012). Predicting total global species richness using rates of species description and estimates of taxonomic effort. *Systematic Biology*, 61(5), 871–883. <https://doi.org/10.1093/sysbio/syr080>
- Cox, J. G., & Lima, S. L. (2006). Naiveté and an aquatic-terrestrial dichotomy in the effects of introduced predators. *Trends in Ecology & Evolution*, 21(12), 674–680. <https://doi.org/10.1016/j.tree.2006.07.011>
- Cudahy, B. J. (2006). The containership revolution: Malcom McLean's 1956 innovation goes global. *Transportation Research News*, 246, 5–9.
- Dawson, W., Moser, D., van Kleunen, M., Kreft, H., Pergl, J., Pyšek, P., ... Essl, F. (2017). Global hotspots and correlates of alien species richness across taxonomic groups. *Nature Ecology & Evolution*, 1, 0186. <https://doi.org/10.1038/s41559-017-0186>
- di Castri, F. (1989). History of biological invasions with special emphasis on the Old World. In J. A. Drake, H. A. Mooney, F. di Castri, R. H. Groves, F. J. Kruger, M. Rejmánek, & M. Williamson (Eds.), *Biological invasions: A global perspective* (pp. 1–30). Chichester: John Wiley and Sons.
- Dyer, E. E., Cassey, P., Redding, D. W., Collen, B., Franks, V., Gaston, K. J., ... Blackburn, T. M. (2017). The global distribution and drivers of alien bird species richness. *PLoS Biology*, 15(1), e2000942. <https://doi.org/10.1371/journal.pbio.2000942>
- Early, R., Bradley, B. A., Dukes, J. S., Lawler, J. J., Olden, J. D., Blumenthal, D. M., ... Tatem, A. J. (2016). Global threats from invasive alien species in the twenty-first century and national response capacities. *Nature Communications*, 7, 12485. <https://doi.org/10.1038/ncomms12485>
- Ellis, E. C., Kaplan, J. O., Fuller, D. Q., Vavrus, S., Goldewijk, K. K., Verburg, P. H., ... Verburg, P. H. (2013). Used planet: A global history. *Proceedings of the National Academy of Sciences of the United States of America*, 110(20), 7978–7985. <https://doi.org/10.1073/pnas.1217241110>
- Essl, F., Bacher, S., Blackburn, T. M., Booy, O., Brundu, G., Brunel, S., ... Jeschke, J. M. (2015). Crossing frontiers in tackling pathways of biological invasions. *BioScience*, 65(8), 769–782. <https://doi.org/10.1093/biosci/biv082>
- Essl, F., Lenzner, B., Bacher, S., Bailey, S., Capinha, C., Daehler, C., ... Roura-Pascual, N. (2020). Drivers of future alien species impacts: An expert-based assessment. *Global Change Biology*, 26(9), 4880–4893. <https://doi.org/10.1111/gcb.15199>
- Fridley, J. D., & Sax, D. F. (2014). The imbalance of nature: Revisiting a Darwinian framework for invasion biology. *Global Ecology and Biogeography*, 23(11), 1157–1166. <https://doi.org/10.1111/geb.12221>
- Froese, R., & Pauly, D. (Eds.). (2015). FishBase. Retrieved from <http://www.Fishbase.de/>
- Hulme, P. E. (2009). Trade, transport and trouble: Managing invasive species pathways in an era of globalization. *Journal of Applied Ecology*, 46(1), 10–18. <https://doi.org/10.1111/j.1365-2664.2008.01600.x>
- IPBES. (2016). *Summary for policymakers of the methodological assessment of scenarios and models of biodiversity and ecosystem services of the Intergovernmental Science-Policy Platform on Biodiversity and Ecosystem Services*. In V. C. S. Ferrier, K. N. Ninan, P. Leadley, R. Alkemade, L. A. Acosta, H. R. Akçakaya, L. Brotons, W. Cheung, N. H. R. K. A. Harhash, J. Kabubo-Mariara, C. Lundquist, M. Obersteiner, H. Pereira, G. Peterson, R. Pichs-Madruga, & B. W. C. Rondinini (Eds.). Bonn, Germany: Secretariat of the Intergovernmental Science-Policy Platform on Biodiversity and Ecosystem Services.
- IPBES. (2019). *Summary for policymakers of the global assessment report on biodiversity and ecosystem services of the Intergovernmental Science-Policy Platform on Biodiversity and Ecosystem Services*. In S. Diaz, J. Settele, E. S. Brondizio, H. T. Ngo, M. Guèze, J. Agard, & C. N. Zayas (Eds.). Bonn, Germany: IPBES Secretariat.
- IPCC. (2014). *Climate change 2014: Synthesis report. Contribution of working groups I, II and III to the fifth assessment report of the Intergovernmental Panel on Climate Change*. Geneva, Switzerland: IPCC.
- Knapp, S., Winter, M., & Klotz, S. (2017). Increasing species richness but decreasing phylogenetic richness and divergence over a 320-year period of urbanization. *Journal of Applied Ecology*, 54(4), 1152–1160. <https://doi.org/10.1111/1365-2664.12826>
- Lenzner, B., Latombe, G., Capinha, C., Bellard, C., Courchamp, F., Diagne, C., ... Essl, F. (2020). What will the future bring for biological invasions

- on islands? An expert-based assessment. *Frontiers in Ecology and Evolution*, 8. <https://doi.org/10.3389/fevo.2020.00280>
- Lenzner, B., Leclère, D., Franklin, O., Seebens, H., Roura-Pascual, N., Obersteiner, M., ... Essl, F. (2019). A framework for global twenty-first century scenarios and models of biological invasions. *BioScience*, 69(9), 697–710. <https://doi.org/10.1093/biosci/biz070>
- Leung, B., Springborn, M. R., Turner, J. A., & Brockerhoff, E. G. (2014). Pathway-level risk analysis: The net present value of an invasive species policy in the US. *Frontiers in Ecology and the Environment*, 12(5), 273–279. <https://doi.org/10.1890/130311>
- Lewis, S. L., & Maslin, M. A. (2015). Defining the Anthropocene. *Nature*, 519(7542), 171–180. <https://doi.org/10.1038/nature14258>
- Liebold, A. M., Brockerhoff, E. G., & Kimberley, M. (2017). Depletion of heterogeneous source species pools predicts future invasion rates. *Journal of Applied Ecology*, 54(6), 1968–1977. <https://doi.org/10.1111/1365-2664.12895>
- Meyerson, L. A., & Mooney, H. A. (2007). Invasive alien species in an era of globalization. *Frontiers in Ecology and the Environment*, 5(4), 199–208. [https://doi.org/10.1890/1540-9295\(2007\)5\[199:IASIAE\]2.0.CO;2](https://doi.org/10.1890/1540-9295(2007)5[199:IASIAE]2.0.CO;2)
- Mora, C., Tittensor, D. P., Adl, S., Simpson, A. G. B., & Worm, B. (2011). How many species are there on earth and in the ocean? *PLoS Biology*, 9(8), 1–8. <https://doi.org/10.1371/journal.pbio.1001127>
- Muñoz-Mas, R., & García-Berthou, E. (2020). Alien animal introductions in Iberian inland waters: An update and analysis. *Science of the Total Environment*, 703, 134505. <https://doi.org/10.1016/j.scitotenv.2019.134505>
- Pauchard, A., & Alaback, P. B. (2004). Influence of elevation, land use, and landscape context on patterns of alien plant invasions along roadsides in protected areas of south-central Chile. *Conservation Biology*, 18(1), 238–248. <https://doi.org/10.1111/j.1523-1739.2004.00300.x>
- Pyšek, P., Hulme, P. E., Simberloff, D., Bacher, S., Blackburn, T. M., Carlton, J. T., ... Richardson, D. M. (2020). Scientists' warning on invasive alien species. *Biological Reviews*, brv.12627. <https://doi.org/10.1111/brv.12627>
- Pyšek, P., Pergl, J., Essl, F., Lenzner, B., Dawson, W., Kreft, H., ... Kleunen, M. V. (2017). Naturalized alien flora of the world: Species diversity, taxonomic and phylogenetic patterns, geographic distribution and global hotspots of plant invasion. *Preslia*, 89(3), 203–274. <https://doi.org/10.23855/preslia.2017.203>
- R Core Team. (2019). *R: A language and environment for statistical computing*. Vienna, Austria: R Foundation for Statistical Computing. Retrieved from <https://www.r-project.org/>
- Rouget, M., Robertson, M. P., Wilson, J. R. U., Hui, C., Essl, F., Renteria, J. L., & Richardson, D. M. (2016). Invasion debt – Quantifying future biological invasions. *Diversity and Distributions*, 22(4), 445–456. <https://doi.org/10.1111/ddi.12408>
- Russell, J. C., Meyer, J. Y., Holmes, N. D., & Pagad, S. (2017). Invasive alien species on islands: Impacts, distribution, interactions and management. *Environmental Conservation*, 44(4), 359–370. <https://doi.org/10.1017/S0376892917000297>
- Sax, D. F., & Gaines, S. D. (2008). Species invasions and extinction: The future of native biodiversity on islands. *Proceedings of the National Academy of Sciences of the United States of America*, 105, 11490–11497. <https://doi.org/10.1073/pnas.0802290105>
- Seebens, H., Blackburn, T. M., Dyer, E. E., Genovesi, P., Hulme, P. E., Jeschke, J. M., ... Essl, F. (2017). No saturation in the accumulation of alien species worldwide. *Nature Communications*, 8(1), 14435. <https://doi.org/10.1038/ncomms14435>
- Seebens, H., Blackburn, T. M., Dyer, E. E., Genovesi, P., Hulme, P. E., Jeschke, J. M., ... Essl, F. (2018). Global rise in emerging alien species results from increased accessibility of new source pools. *Proceedings of the National Academy of Sciences of the United States of America*, 115(10), E2264–E2273. <https://doi.org/10.1073/pnas.1719429115>
- Seebens, H., Essl, F., Dawson, W., Fuentes, N., Moser, D., Pergl, J., ... Blasius, B. (2015). Global trade will accelerate plant invasions in emerging economies under climate change. *Global Change Biology*, 21(11), 4128–4140. <https://doi.org/10.1111/gcb.13021>
- Seebens, H., Schwartz, N., Schupp, P. J., & Blasius, B. (2016). Predicting the spread of marine species introduced by global shipping. *Proceedings of the National Academy of Sciences of the United States of America*, 113(20), 5646–5651. <https://doi.org/10.1073/pnas.1524427113>
- Sikes, B. A., Bufford, J. L., Hulme, P. E., Cooper, J. A., Johnston, P. R., & Duncan, R. P. (2018). Import volumes and biosecurity interventions shape the arrival rate of fungal pathogens. *PLOS Biology*, 16(5), e2006025. <https://doi.org/10.1371/journal.pbio.2006025>
- Turbelin, A. J., Malamud, B. D., & Francis, R. A. (2017). Mapping the global state of invasive alien species: Patterns of invasion and policy responses. *Global Ecology and Biogeography*, 26(1), 78–92. <https://doi.org/10.1111/gcb.12517>
- van Kleunen, M., Dawson, W., Essl, F., Pergl, J., Winter, M., Weber, E., ... Pyšek, P. (2015). Global exchange and accumulation of non-native plants. *Nature*, 525(7567), 100–103. <https://doi.org/10.1038/nature14910>
- Walther, G.-R., Roques, A., Hulme, P. E., Sykes, M. T., Pyšek, P., Kühn, I., ... Bugmann, H. (2009). Alien species in a warmer world: Risks and opportunities. *Trends in Ecology & Evolution*, 24(12), 686–693. <https://doi.org/10.1016/j.tree.2009.06.008>

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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