

# Climate and socio-economic factors explain differences between observed and expected naturalization patterns of European plants around the world

Robin Pouteau<sup>1,2</sup>  | Wilfried Thuiller<sup>3</sup> | Carsten Hobohm<sup>4</sup> | Caroline Brunel<sup>1</sup>  | Barry J. Conn<sup>5</sup>  | Wayne Dawson<sup>6</sup> | Michele de Sá Dechoum<sup>7,8</sup> | Aleksandr L. Ebel<sup>9,10</sup>  | Franz Essl<sup>11</sup> | Ori Fragman-Sapir<sup>12</sup> | Trevor Fristoe<sup>13</sup> | Nejc Jogan<sup>14</sup> | Holger Kreft<sup>15,16</sup>  | Bernd Lenzner<sup>11</sup> | Carsten Meyer<sup>17,18,19</sup>  | Jan Pergl<sup>20</sup>  | Petr Pyšek<sup>20,21</sup>  | Alla Verkhozina<sup>22</sup> | Patrick Weigelt<sup>15</sup>  | Qiang Yang<sup>13</sup> | Elena Zykova<sup>10</sup> | Svetlana Ačić<sup>23</sup>  | Emiliano Agrillo<sup>24</sup>  | Fabio Attorre<sup>25</sup> | Ariel Bergamini<sup>26</sup>  | Christian Berg<sup>27</sup>  | Erwin Bergmeier<sup>16,28</sup>  | Idoia Biurrun<sup>29</sup>  | Steffen Boch<sup>26</sup>  | Gianmaria Bonari<sup>30,31</sup>  | Zoltán Botta-Dukát<sup>32</sup>  | Helge Bruelheide<sup>17,33</sup>  | Juan Antonio Campos<sup>29</sup>  | Andraž Čarni<sup>34</sup>  | Laura Casella<sup>35</sup>  | Maria Laura Carranza<sup>36</sup>  | Milan Chytrý<sup>30</sup>  | Renata Čuštereška<sup>37</sup> | Michele De Sanctis<sup>25</sup>  | Jürgen Dengler<sup>17,38,39</sup>  | Panayotis Dimopoulos<sup>40</sup> | Rasmus Ejrnæs<sup>41</sup> | Jörg Ewald<sup>42</sup>  | Giuliano Fanelli<sup>25</sup> | Federico Fernández-González<sup>43</sup> | Rosario G. Gavilán<sup>44</sup>  | Jean-Claude Gegout<sup>45</sup> | Rense Haveman<sup>46</sup> | Maike Isermann<sup>47,48</sup> | Ute Jandt<sup>17,33</sup>  | Florian Jansen<sup>49</sup>  | Borja Jiménez-Alfaro<sup>50</sup>  | Ali Kavgaci<sup>51</sup> | Larisa Khanina<sup>52</sup> | Ilona Knollová<sup>30</sup> | Anna Kuzemko<sup>53</sup>  | Maria Lebedeva<sup>54</sup> | Jonathan Lenoir<sup>55</sup> | Tatiana Lysenko<sup>56,57</sup>  | Corrado Marcenò<sup>30</sup>  | Vasiliy Martynenko<sup>58</sup>  | Jesper Erenskjold Moeslund<sup>41</sup>  | Ricarda Pätzsch<sup>59</sup>  | Remigiusz Pielech<sup>60</sup>  | Valerijus Rašomavičius<sup>61</sup> | Iris de Ronde<sup>46</sup> | Eszter Ruprecht<sup>62</sup> | Solvita Rūsiņa<sup>63</sup> | Pavel Shirokikh<sup>58</sup>  | Jozef Šibík<sup>64</sup>  | Urban Šilc<sup>34</sup> | Angela Stanisci<sup>36</sup>  | Zvezdana Stančić<sup>65</sup>  | Jens-Christian Svenning<sup>66</sup>  | Grzegorz Swacha<sup>67</sup>  | Pavel Dan Turtureanu<sup>68</sup>  | Milan Valachovič<sup>64</sup>  | Kiril Vassilev<sup>69</sup>  | Sergey Yamalov<sup>54</sup> | Mark van Kleunen<sup>1,13</sup> 

<sup>1</sup>Zhejiang Provincial Key Laboratory of Plant Evolutionary Ecology and Conservation, Taizhou University, Taizhou, China

<sup>2</sup>AMAP, Univ. Montpellier, IRD, CIRAD, CNRS, INRAE, Montpellier, France

<sup>3</sup>Univ. Grenoble Alpes, Univ. Savoie Mont Blanc, CNRS, LECA, Laboratoire d'Ecologie Alpine, 38000, Grenoble, France

<sup>4</sup>Ecology and Environmental Education Working Group, University of Flensburg (EUF), Flensburg, Germany

<sup>5</sup>School of Life and Environmental Sciences, University of Sydney, Sydney, New South Wales, Australia

<sup>6</sup>Department of Biosciences, Durham University, Durham, UK

<sup>7</sup>Department of Ecology and Zoology, Federal University of Santa Catarina, Florianópolis, Brazil

<sup>8</sup>The Horus Institute for Environmental Conservation and Development, Florianópolis, Brazil

<sup>9</sup>Department of Botany, Tomsk State University, Tomsk, Russia

<sup>10</sup>Central Siberian Botanical Garden, Siberian Branch of Russian Academy of Sciences, Novosibirsk, Russia

- <sup>11</sup>Bioinvasions, Global Change, Macroecology-Group, Department of Botany and Biodiversity Research, University of Vienna, Vienna, Austria
- <sup>12</sup>Jerusalem Botanical Gardens, The Hebrew University of Jerusalem, Jerusalem, Israel
- <sup>13</sup>Ecology, Department of Biology, University of Konstanz, Konstanz, Germany
- <sup>14</sup>Department of Biology, Biotechnical Faculty, University of Ljubljana, Ljubljana, Slovenia
- <sup>15</sup>Biodiversity, Macroecology & Biogeography, University of Göttingen, Göttingen, Germany
- <sup>16</sup>Centre of Biodiversity and Sustainable Land Use (CBL), University of Göttingen, Göttingen, Germany
- <sup>17</sup>German Centre for Integrative Biodiversity Research (iDiv) Halle-Jena-Leipzig, Leipzig, Germany
- <sup>18</sup>Institute of Biology, Leipzig University, Leipzig, Germany
- <sup>19</sup>Institute for Geosciences and Geography, Martin Luther University Halle-Wittenberg, Halle (Saale), Germany
- <sup>20</sup>Institute of Botany, Department of Invasion Ecology, Czech Academy of Sciences, Průhonice, Czech Republic
- <sup>21</sup>Department of Ecology, Faculty of Science, Charles University, Prague, Czech Republic
- <sup>22</sup>Siberian Institute of Plant Physiology & Biochemistry, Siberian Branch of Russian Academy of Sciences, Irkutsk, Russia
- <sup>23</sup>Faculty of Agriculture, Department of Botany, University of Belgrade, Belgrade, Serbia
- <sup>24</sup>Operational Center for Environmental Monitoring, Italian Institute for Environmental Protection and Research (ISPRA), Rome, Italy
- <sup>25</sup>Department of Environmental Biology, Sapienza University of Rome, Rome, Italy
- <sup>26</sup>WSL Swiss Federal Research Institute, Birmensdorf, Switzerland
- <sup>27</sup>Institute for Biology, Universität of Graz, Graz, Austria
- <sup>28</sup>Department of Vegetation Science and Phytodiversity, University of Göttingen, Göttingen, Germany
- <sup>29</sup>Department of Plant Biology and Ecology, University of the Basque Country UPV/EHU, Bilbao, Spain
- <sup>30</sup>Department of Botany and Zoology, Faculty of Science, Masaryk University, Brno, Czech Republic
- <sup>31</sup>Faculty of Science and Technology, Free University of Bozen-Bolzano, Bozen, Italy
- <sup>32</sup>Institute of Ecology and Botany, MTA Centre for Ecological Research, Vácrátót, Hungary
- <sup>33</sup>Institute of Biology/Geobotany and Botanical Garden, Martin Luther University Halle-Wittenberg, Halle, Germany
- <sup>34</sup>Research Center of the Slovenian Academy of Sciences and Arts, University of Nova Gorica, Nova Gorica, Slovenia
- <sup>35</sup>Department for Biodiversity Conservation and Monitoring, Italian Institute for Environmental Protection and Research (ISPRA), Rome, Italy
- <sup>36</sup>Envixlab, Department of Biosciences and Territory, University of Molise, Pesche, Italy
- <sup>37</sup>Institute of Biology, Faculty of Natural Sciences and Mathematics, University of Ss. Cyril and Methodius, Skopje, Republic of North Macedonia
- <sup>38</sup>Vegetation Ecology, Institute of Natural Resource Sciences (IUNR), Zurich University of Applied Sciences (ZHAW), Wädenswil, Switzerland
- <sup>39</sup>Plant Ecology, Bayreuth Center of Ecology and Environmental Research (BayCEER), University of Bayreuth, Bayreuth, Germany
- <sup>40</sup>Laboratory of Botany, Department of Biology, University of Patras, Patras, Greece
- <sup>41</sup>Section for Biodiversity, Department of Bioscience, Aarhus University, Rønde, Denmark
- <sup>42</sup>Institute for Ecology and Landscape, Weißenstephan-Triesdorf University of Applied Sciences, Freising, Germany
- <sup>43</sup>Institute of Environmental Sciences, University of Castilla-La Mancha, Toledo, Spain
- <sup>44</sup>Department of Pharmacology, Pharmacognosy and Botany, Complutense University, Madrid, Spain
- <sup>45</sup>AgroParisTech, INRAE, Université de Lorraine, Nancy, France
- <sup>46</sup>Central Government Real Estate Agency, Ministry of the Interior and Kingdom Relations, Wageningen, the Netherlands
- <sup>47</sup>Institute of Ecology, Bremen University, Bremen, Germany
- <sup>48</sup>Lower Saxon Wadden Sea National Park Authority, Nature Conservation, Wilhelmshaven, Germany
- <sup>49</sup>Faculty of Agricultural and Environmental Sciences, University of Rostock, Rostock, Germany
- <sup>50</sup>Department of Organisms and Systems Biology and Research Unit of Biodiversity (UMIB, UO-CSIC-PA), University of Oviedo, Oviedo, Spain
- <sup>51</sup>Department of Forest Botany, Faculty of Forestry, University of Karabük, Karabük, Turkey
- <sup>52</sup>Institute of Mathematical Problems of Biology RAS, Branch of the M.V. Keldysh Institute of Applied Mathematics of the Russian Academy of Sciences, Moscow, Russia
- <sup>53</sup>Department of Geobotany and Ecology, M.G. Kholodny Institute of Botany NAS of Ukraine, Kyiv, Ukraine
- <sup>54</sup>South-Ural Botanical Garden-Institute, Ufa Federal Research Centre, Russian Academy of Science, Ufa, Russia
- <sup>55</sup>UR "Ecologie et Dynamique des Systèmes Anthropisés" (EDYSAN, UMR 7058 CNRS), Université de Picardie Jules Verne, Amiens, France
- <sup>56</sup>Komarov Botanical Institute of the Russian Academy of Sciences, Saint-Petersburg, Russia
- <sup>57</sup>Institute of Ecology of the Volga River Basin of the Russian Academy of Sciences, Branch of the Samara Scientific Center of the Russian Academy of Sciences, Togliatti, Russia
- <sup>58</sup>Laboratory of Geobotany and Plant Resources, Ufa Institute of Biology of Ufa Federal Scientific Centre of Russian Academy of Sciences, Ufa, Republic of Bashkortostan, Russia
- <sup>59</sup>Wagner Ecology Lab, Department of Biological Sciences, University of Alberta, Edmonton, Alberta, Canada
- <sup>60</sup>Department of Forest Biodiversity, Faculty of Forestry, University of Agriculture, Kraków, Poland

<sup>61</sup>Institute of Botany, Nature Research Centre, Vilnius, Lithuania

<sup>62</sup>Hungarian Department of Biology and Ecology, Babeş-Bolyai University, Cluj-Napoca, Romania

<sup>63</sup>Faculty of Geography and Earth Sciences, University of Latvia, Rīga, Latvia

<sup>64</sup>Institute of Botany, Plant Science and Biodiversity Center, Slovak Academy of Sciences, Bratislava, Slovakia

<sup>65</sup>Faculty of Geotechnical Engineering, University of Zagreb, Varaždin, Croatia

<sup>66</sup>Center for Biodiversity Dynamics in a Changing World (BIOCHANGE) and Section for Ecoinformatics and Biodiversity, Department of Biology, Aarhus University, Aarhus C, Denmark

<sup>67</sup>Botanical Garden, University of Wrocław, Wrocław, Poland

<sup>68</sup>A. Borza Botanical Garden, Emil G. Racoviță Institute, Center for Systematic Biology, Babes-Bolyai University, Cluj-Napoca, Romania

<sup>69</sup>Institute of Biodiversity and Ecosystem Research, Bulgarian Academy of Sciences, Sofia, Bulgaria

#### \*Correspondence

Robin Pouteau, Zhejiang Provincial Key Laboratory of Plant Evolutionary Ecology and Conservation, Taizhou University, Taizhou 318000, China.  
Email: robin.pouteau@ird.fr

Editor: Melodie McGeoch

#### Abstract

**Aim:** The number of naturalized (i.e. established) alien species has increased rapidly over recent centuries. Given the differences in environmental tolerances among species, little is known about what factors determine the extent to which the observed size of the naturalized range of a species and hence the extent to which the observed richness of naturalized species of a region approach their full potential. Here, we asked which region- and species-specific characteristics explain differences between observed and expected naturalizations.

**Location:** Global.

**Time period:** Present.

**Major taxa studied:** Vascular plants.

**Methods:** We determined the observed naturalized distribution outside Europe for 1,485 species endemic to Europe using the Global Naturalized Alien Flora (GloNAF) database and their expected distributions outside Europe using species distribution models. First, we investigated which of seven socio-economic factors related to introduction pathways, anthropogenic pressures and inventory effort best explained the differences between observed and expected naturalized European floras. Second, we examined whether distributional features, economic use and functional traits explain the extent to which species have filled their expected ranges outside Europe.

**Results:** In terms of suitable area, more than 95% of expected naturalizations of European plants were not yet observed. Species were naturalized in only 4.2% of their suitable regions outside of Europe (range filling) and in 0.4% of their unsuitable regions (range expansion). Anthropogenic habitat disturbance primarily explained the difference between observed and expected naturalized European floras, as did the number of treaties relevant to invasive species. Species of ornamental and economic value and with large specific leaf area performed better at filling and expanding beyond their expected range.

**Main conclusions:** The naturalization of alien plant species is explained by climate matching but also by the regional level of human development, the introduction pressure associated with the ornamental and economic values of the species and their adaptation to disturbed environments.

#### KEYWORDS

alien species, anthropogenic pressure, environmental driver, functional trait, global change, introduction pathway, naturalization, ornamental plant, sampling bias, species distribution model

## 1 | INTRODUCTION

Species that maintain self-sustaining populations outside their native range independently of direct human intervention (i.e. naturalized species; Blackburn et al., 2011; Richardson et al., 2000) pose a serious concern to native biodiversity and ecosystem functioning and services worldwide (IPBES, 2019). Their number has increased substantially over the last few centuries and does not show any indication of saturation (Seebens et al., 2017). Regions differ considerably in their richness of naturalized species of vascular plants, and species vary considerably in their extent of naturalization (Essl et al., 2019; Pyšek et al., 2017). Part of this variation may result from regional differences in the size of the potential naturalized alien pool and the size of the range potentially available for naturalized species. Little is known about the relationship between the observed and potential richness of naturalized plant species (i.e. the maximum numbers of alien species that these regions could host given species' environmental tolerances; Stohlgren et al., 2008).

The actual distribution of naturalized alien plant species results from the number of introductions by humans (intentional or not) followed by the interplay of factors that determine the environmental space available for a species and factors that prevent or promote the extent to which an equilibrium with an environmental space is reached. Environmental suitability (e.g. climate, habitat) determines the potential (expected) distribution range by filtering the subset of introduced species that can maintain populations over long periods by successfully reproducing, that is, naturalized species (Richardson & Pyšek, 2012). Human activities can further influence the observed distribution range of species by altering environmental conditions and dispersal processes that dictate opportunities for introductions. For instance, human transport, trade and increasing urban development tend to broaden observed ranges, while, conversely, biosecurity responses can contain this expansion (Hulme, 2009).

An increasing body of literature is investigating what proportions of species' potential alien ranges are actually occupied (Hill et al., 2017; Liu et al., 2020; Petitpierre et al., 2012; Strubbe et al., 2013, 2015). Indeed, most alien species are occupying only a subset of the niche corresponding to suitable environments that are occupied in their native range ('niche filling'). Certain alien species are also able to colonize new environments that are not occupied in their native range ('niche expansion'). Patterns of niche filling and niche expansion vary highly depending on the type of organisms. For instance, the average niche filling was found to be 80% for vascular plants considered invasive (i.e. spreading into areas away from sites of introduction; Richardson et al., 2000), 37% for invasive insects and 20% for invasive birds, whereas the average niche expansion was 26% for birds, 20% for insects and 5% for plants (Hill et al., 2017; Petitpierre et al., 2012; Strubbe et al., 2013). Niche changes during invasions appear to be primarily driven by introduction history (e.g. propagule pressure and colonization processes) in birds, and by human disturbance in insects (Hill et al., 2017). However, little is known about what causes possible differences between observed and expected ranges in naturalized plants.

Much of the research effort has been devoted to understanding the socio-economic drivers of the number of naturalized alien plant species (Dawson et al., 2017; Essl et al., 2019; Pyšek et al., 2010; van Kleunen et al., 2015), some of these studies focusing on the subset of invasive species (Bellard et al., 2016; Essl et al., 2015). However, studies dealing with alien species richness always face the challenge of disentangling the specific role of socio-economic drivers from that of large-scale biogeographical factors also known to influence species richness, irrespective of species origin (e.g. latitudinal gradient, resource availability). Moreover, while it is recognized that environmental matching between native and alien ranges largely controls naturalization (Feng et al., 2016; Richardson & Pyšek, 2012), focusing on alien species richness does not allow accounting for species-specific environmental requirements. Hence, determining the expected distributional range of naturalized species should be an important step in understanding the influence of socio-economic factors on observed patterns of naturalization (Xu et al., 2019).

Another advantage of analysing species composition is that it allows for the identification of the species that explain the pattern of naturalized species richness and their characteristics. Many other studies have attempted to determine which species' performance or natural distribution traits are associated with naturalization success by comparing naturalized and non-naturalized plant species (e.g. Pyšek & Richardson, 2007; Pyšek et al., 2015; van Kleunen et al., 2010). Conclusions from these studies delivered little consensus and often appear to be context- and species-dependent. This might partly be because these studies only investigated trait differences of native versus alien or non-invasive versus invasive species (i.e. a binary variable), or tested for relationships between traits and the observed naturalized range without considering the potential range. Therefore, we believe we can gain additional insight into the drivers of naturalization patterns by searching for traits that explain the extent to which species fill or expand beyond their suitable alien range (i.e. a continuous variable that accounts for the potential range).

Europe is the second-most important donor continent of naturalized alien plant species, after Asia. However, the European flora is by far the most successful in naturalizing. Of the c. 14,200 vascular plant species native (i.e. both endemic and non-endemic) to Europe, 3,383 have become naturalized somewhere in the world; this is three times more than expected by chance (van Kleunen et al., 2015). Furthermore, five of the ten globally most widely naturalized species are native to Europe (Pyšek et al., 2017). The European flora is, therefore, suitable for studying drivers of differences between observed and expected naturalized ranges.

Current knowledge of which region- and plant-related characteristics influence the extent to which species fill their expected naturalized ranges is poor and mainly derived from the search of the socio-economic drivers of niche/range filling of a few invasive alien species (usually < 50; e.g. Hill et al., 2017; Petitpierre et al., 2012; Strubbe et al., 2013, 2015). However, the drivers of niche/range filling have never been explored for naturalized species (a much

broader group of species than invasives; Richardon et al., 2000); yet the processes driving the ability to survive and reproduce are likely to differ from the processes driving the spread and impact of invasive species (e.g. Abellán et al., 2017). Moreover, the drivers of the difference between observed and expected naturalized regional floras (and not between observed and expected distributions of a few individual species) have never been explored.

In this study, we focused on 1,485 plant species endemic to Europe, an unprecedented number for this type of study, which allowed us to analyse in more detail what explains variation in the differences between observed and expected values. We determined both their current observed distribution in 931 non-European regions using the Global Naturalized Alien Flora database (GloNAF; van Kleunen et al., 2019) and their expected distribution using biogeoclimatic ensemble species distribution models (SDMs). We also explored which socio-economic factors explain why some regions have a recorded naturalized flora of European origin close to their expected naturalized flora, while other regions have not. Finally, we examined whether species distribution patterns, economic uses and functional traits explain why some species fill a larger portion of their expected range than others do.

## 2 | MATERIALS AND METHODS

### 2.1 | Expected naturalized ranges

#### 2.1.1 | Species selection

Many plant species native to Europe are also native to other continents such as Asia and Africa, for which there is usually a lower density (temporal and spatial) of species occurrence records (Meyer et al., 2016). Therefore, we focused on vascular plant species whose native range is restricted to Europe (European endemics), as this allows us to better capture the full realized niche of the species in their native range. The physiography of the European continent was defined as bordered by the Arctic Ocean to the north (Iceland and Norwegian Islands were included, Greenland was excluded), the Atlantic Ocean to the west (the British and Irish Isles were included and the Macaronesian archipelagos were excluded), the Ural Mountains, the Ural River, and the Caspian Sea to the east, and the Caucasus and the Mediterranean Sea to the south (Mediterranean islands were included, Anatolia was excluded).

The database Endemic vascular plants in Europe (EvaplantE; Hobohm, 2014), which comprises a list of > 6,200 endemic taxa, was used as a baseline for species selection. Scientific names were standardized based on a working list of all plant species [The Plant List (TPL): <http://www.theplantlist.org/> version 12-10-2018]. This taxonomic standardization was done with the package 'Taxonstand' (Cayuela et al., 2017) in R 4.0.2 (R Development Core Team, 2020). Standardized infraspecific taxa were excluded from the list, resulting in 4,965 species (Figure 1).

#### 2.1.2 | Compilation of species occurrence records

To comprehensively describe the distribution of the species in Europe (which in addition to the native range might for some species also include part of their alien range), we combined occurrence records from six sources (see Supporting Information Appendix S1 for details and references): the Global Biodiversity Information Facility (GBIF), the European Vegetation Archive (EVA; Chytrý et al., 2016), the EU-Forest dataset, the Atlas Florae Europaeae, the Plant Functional Diversity of Grasslands network (DIVGRASS) and the digital atlas of the German flora.

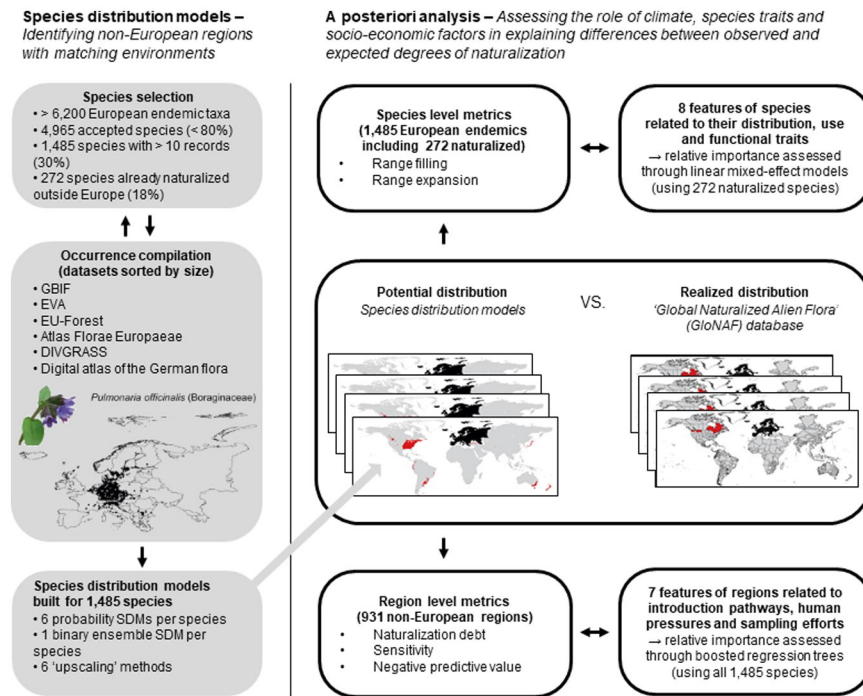
When several occurrence records from these different sources were duplicated for the same cell, only one occurrence record per species was kept to avoid pseudoreplication. Cell size was set to  $0.42^\circ \times 0.42^\circ$  (c. 50 km  $\times$  50 km at the equator) to approach the resolution of the source of occurrence records having the coarsest resolution (Atlas Florae Europaeae). Using a higher resolution (i.e. a smaller cell size) would have required downscaling of this source of occurrence records, which typically introduces spatial biases and uncertainty in the model predictions (Bombi & D'Amen, 2014). Moreover, this resolution appeared to be a good compromise to account for the fact that the selected environmental variables (climate, land use and soil type) determine invasion potential at different scales (10,000 to 200 km, 2,000 to 10 km, and 10 km to 10 m, respectively; Milbau et al., 2009).

Species with fewer than 10 deduplicated occurrence records were not further considered since the resulting SDM might be inaccurate. The final dataset comprised 135,189 occurrence records for a total of 1,485 European plant species, belonging to 327 genera and 67 families (Supporting Information Table S1), that is, on average 91 occurrence records per species with a maximum of 1,382 occurrence records for *Achillea ptarmica* (Asteraceae). We had enough occurrence records to build an SDM (i.e. 10 occurrences) for 272 European species among the 407 already naturalized species and for 1,213 species currently not known to be naturalized anywhere.

#### 2.1.3 | Environmental predictors of expected ranges

We defined six environmental variables to model and project species expected ranges. These variables were related to climate, land use, and soil physico-chemical properties, which are commonly recognized to shape the distribution of plants (Gurevitch et al., 2006). Annual mean temperature ( $^\circ\text{C}$ ), annual precipitation (mm) and precipitation seasonality (yearly coefficient of variation) representing the period 1979–2013 were provided at a 30-arcsec resolution by the CHELSA (Climatologies at high resolution for the earth's land surface areas) climate database (Karger et al., 2017). Worldwide correlations between these variables and the others described below did not exceed the threshold of  $|r| = .70$  (Supporting Information Figure S1) beyond which collinearity begins to severely distort model estimations and subsequent predictions (Dormann et al., 2013). However,





**FIGURE 1** Workflow of this study used to determine which features of both plants and recipient regions explain possible discrepancies between observed and expected extents of naturalization. GBIF = Global Biodiversity Information Facility; EVA = European Vegetation Archive; DIVGRASS = Plant Functional Diversity of Grasslands network; SDM = species distribution model [Colour figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

these variables reflecting trends in average climate conditions were significantly correlated with climate extremes to which plants are recognized to be highly responsive (Zimmermann et al., 2009). The percentage of each grid cell with primary land cover based on the Harmonized Global Land Use models was also used (Chini et al., 2014). Organic carbon content (g per kg) and pH in the first 15 cm of soil were extracted at a 1-km resolution from the global gridded soil information database SoilGrids (Hengl et al., 2014). Environmental variables were aggregated (using the mean value) to the resolution of  $0.42^\circ \times 0.42^\circ$ .

### 2.1.4 | Species distribution modelling

Comparing the realized niche within the native versus alien ranges can lead to three cases: (a) in the alien range, the species uses a similar or smaller realized niche than in the native range, for example, because of new competitors, herbivores or pathogens, or dispersal limitation; (b) the species occupies a realized niche very different from the one in the native area but within the species' initial fundamental niche, for example, because of new biotic interactions, multiple sites of introduction, niche differentiation or different environmental conditions; or (c) the realized niche extends outside of the species' initial fundamental niche, for example, because of rapid niche evolution, enemy-release or new positive interactions (see Gallien et al., 2010 for further details). As options (b) and (c) remain impossible to predict for a large number of species, we only considered option (a) in this study.

The expected distribution of the 1,485 European plant species was modelled by statistically relating the environmental predictors to the distribution data in Europe. Six SDM methods

including generalized additive models, generalized linear models, generalized boosting trees, maximum entropy, multivariate adaptive regression splines and random forest were used. All of these methods require presence and absence or pseudo-absence/background data (a random subset of the available environmental conditions in the area, i.e. Europe in our case), whose selection can significantly affect predictions if not made adequately. Consequently, we performed a preliminary analysis to identify the most appropriate set of pseudo-absences for each SDM method (Supporting Information Appendix S2). The predictive performance of SDM in Europe was assessed by measuring the area under the receiver operating characteristic (ROC) curve (AUC) and the true skill statistics (TSS; Allouche et al., 2006). AUC values can range from 0 to 1, an AUC of 1 indicating a perfect fit, an AUC of .5 meaning that predictions from SDM do not differ from random, and an AUC of 0 meaning the SDM is always incorrect. TSS ranges from -1 to 1, where 1 indicates perfect agreement and 0 indicates a random prediction.

To combine the predictive capability of the six SDMs, their projections were subsequently aggregated into an average or consensus projection. To ensure the quality of the ensemble SDM, we only kept the projections for which the accuracies estimated by AUC and TSS were higher than .8 and .6, respectively, and assembled the selected SDMs using a committee-average approach with each SDM weighted proportional to its TSS evaluation. Probability maps obtained from the ensemble forecasting projections were then transformed into binary suitable/non-suitable maps using the threshold maximizing the TSS to ensure the most accurate predictions, since it is based on both sensitivity and specificity. The entire species distribution modelling workflow was performed within the 'biomod2' R platform (Thuiller et al., 2009).

## 2.2 | Recorded naturalized ranges

The current observed naturalized range of the European flora was obtained through the GloNAF database version 1.2, a recently compiled database of c. 14,000 naturalized alien plant species covering 1,029 regions worldwide, corresponding to administrative regions such as countries, states and provinces, and also including 381 islands (van Kleunen et al., 2019). The size of a region ranged from 0.03 km<sup>2</sup> (Tauna islet of the Gambier archipelago, French Polynesia) to 2,486,952 km<sup>2</sup> (the Republic of the Sudan plus South Sudan). A total of 407 European endemic species were found as naturalized in at least one of the non-European GloNAF regions.

We explored various definitions of suitability by analysing what proportion of a region should be suitable for a species to become naturalized in this region. The following definitions were used: at least one suitable cell, 1, 5, 20, 50 or 100% suitable cells. For each region, species were classified into: true positives (TP), that is, species that are both observed and predicted as being successfully naturalized in the region; false positives (FP), that is, species that have not become naturalized, but are predicted as successfully naturalized; false negatives (FN), that is, species that are observed as successfully naturalized but not predicted to be naturalized; and true negatives (TN), that is, species that are neither predicted nor observed to be naturalized in the region. Then, the predicted naturalized species pool was compared to the observed naturalized pool by calculating: (a) the naturalization debt in the broad sense hereafter referred to as 'naturalization debt' ( $FP/(TP + FP)$ ), that is, the proportion of predicted naturalizations that are not yet observed either because species are not yet introduced to the focal area (introduction debt) or because they were introduced, but are not naturalized yet (naturalization debt in the strict sense, i.e. sensu Rouget et al., 2016), which can take several decades or centuries for some taxa; (b) the assemblage sensitivity ( $TP/(TP + FN)$ ), that is, the proportion of observed naturalizations that are correctly predicted; (c) the negative predictive value ( $TN/(TN + FN)$ ), that is, the proportion of species

that were not predicted to be naturalized and have not become naturalized in the region.

## 2.3 | Socio-economic drivers of naturalization patterns

Seven factors were extracted to explain naturalization debt, assemblage sensitivity and the negative predictive value (Table 1). These factors were selected because they provided insights into introduction pathways, human pressures and inventory effort: airport density ('Airports'), seaport density ('Seaports'), number of international treaties relevant to invasive alien species ('Treaties'), human population density ('Population'), proportion of croplands ('Cropland'), human development index ('Development') and species inventory effort ('Inventory'; for details, see Supporting Information Appendix S3). These socio-economic factors were independent of the surface area of the regions. They were available for 526 non-European regions including 17 islands or archipelagos and 509 mainland areas. Regional factors were tested for multicollinearity by computing a correlation matrix based on Pearson's  $r$ . Cross-correlations did not exceed  $r = .51$  (between airport and seaport density), which is below the threshold of .70 (Supporting Information Figure S3; Dormann et al., 2013).

Identification of the drivers of naturalization patterns was based on boosted regression trees (BRTs), a machine-learning method that combines a large number of relatively simple tree models to optimize predictive performance (Elith et al., 2008). The quality of BRT fits was controlled with the coefficient of determination of the regression between estimated and observed values ( $r^2$ ) and the standard error (SE). Results were interpreted by looking at the relative influence of the seven regional socio-economic factors on the predictive models (calculated based on the number of times a factor is selected in the model, weighted by its improvement to the overall model) and by considering the partial dependence of the predictions on each factor after accounting for the average effect of the other factors.

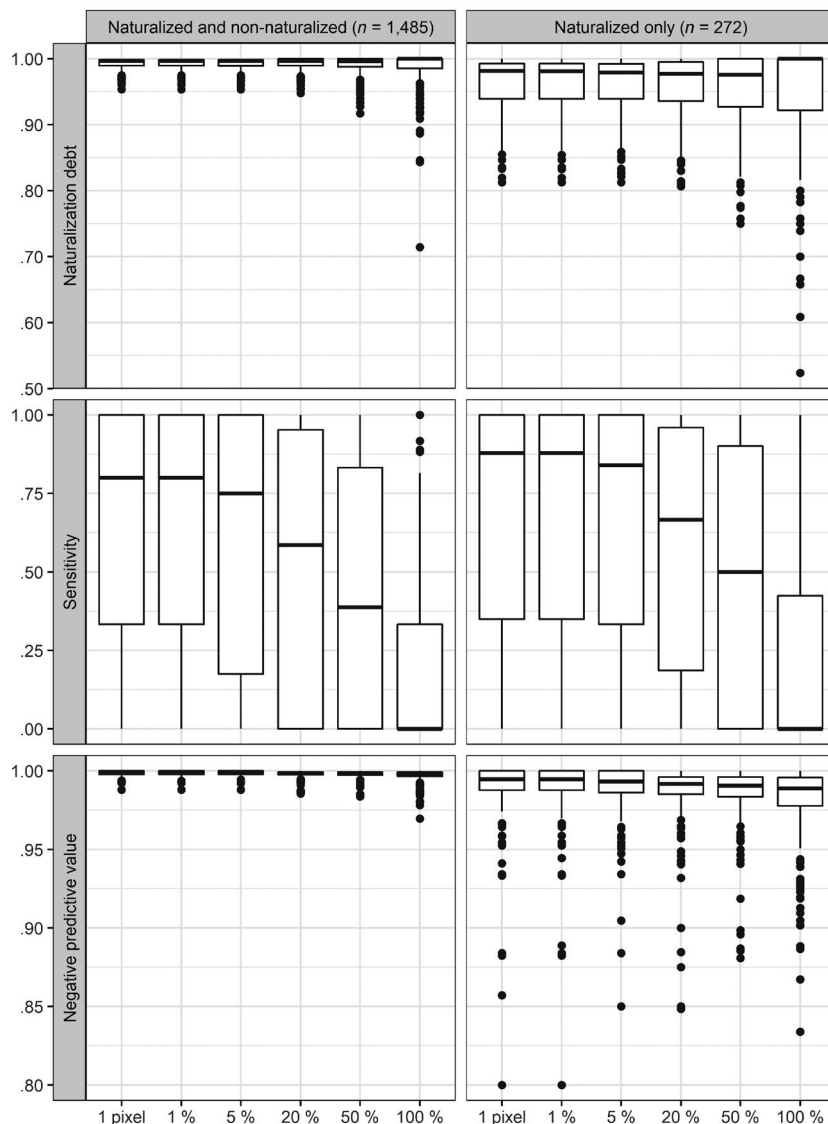
**TABLE 1** Summary of the seven socio-economic factors used to assess the drivers of naturalization success of the European vascular plants including the year and spatial resolution at which the original data were collected/calculated, the range of values for the regions used in analyses, and which transformation was applied to these values prior to analyses. Minimum and maximum values are those, after transformation, over the 526 non-European regions for which all socio-economic factors were available

Variable	Year	Resolution	Unit	Min	Max	Transformation
<i>Introduction pathways</i>						
Airports	2019	–	#/100,000 km <sup>2</sup>	0	11	log
Seaports	2014	–	#/100,000 km <sup>2</sup>	0	15	log
Treaties	2016	–	#	6	30	–
<i>Human pressures</i>						
Population	2000	1 km	Inhabitants/km <sup>2</sup>	0	10	log
Cropland	<2007	5 arcmin	%	0	95	–
Development	2015	5 arcmin	Dimensionless	.27	.93	–
<i>Sampling effort</i>						
Inventory	2015–2016	110 km	%	0	249	–

## 2.4 | Plant features explaining naturalization patterns

To assess which species-level features affect naturalization success, we first compared European species recorded as naturalized outside Europe ('naturalized') and those not recorded as naturalized outside Europe ('non-naturalized'). Eight features including two distributional indices, the economic use of the species and four functional traits were selected: (a) species prevalence in Europe measured as the number of  $0.42^\circ \times 0.42^\circ$  cells occupied; (b) the surface area of the species' expected range outside Europe. These distributional indices were used because more widespread species are usually hypothesized to have a greater potential to spread and a higher invasion success (Gallien et al., 2019); (c) whether the species is used as an ornamental plant or (d) has other economic relevance (e.g. food plant, medicines, materials) according to the World Economic Plants (WEP) database (Wiersema & León, 2013) since the ornamental and economic relevance of a species is thought to be a key predictor of its probability of being

introduced and its naturalization success (van Kleunen et al., 2018, 2020); (e) the growth form (i.e. graminoid, non-graminoid herb, shrub or tree), a major determinant of invasiveness (Pyšek & Richardson, 2007); (f) mean specific leaf area (SLA), that is, the ratio of leaf area to leaf dry mass (expressed in  $\text{m}^2/\text{g}$ ); (g) mean plant height (in m); and (h) mean seed mass (in g). These last three functional traits represent key axes of plant ecological strategies following the leaf-height-seed (LHS) scheme of Westoby (1998). Correlation between LHS traits was  $|r| < .27$ . Functional traits were extracted from the TRY database (Kattge et al., 2020). Linear mixed-effect models (LMMs) were fitted using a phylogenetic generalized least square approach (PGLS) to compare features of naturalized and non-naturalized species while controlling for between-species phylogenetic distances (for details on the phylogeny used, see Supporting Information Appendix S4). As we did not have data on all features for each species (see Figure 6), we ran separate LMMs for each of the eight features. LMMs were performed using the *glsl()* function of the R package 'nlme' (Pinheiro et al., 2019).



**FIGURE 2** Comparison of predicted and observed naturalized floras in 931 non-European regions based on various upscaling methods used to transform local naturalization predictions into regional naturalization predictions. The left column comprises our full set of European plants (either already naturalized or not yet naturalized outside of Europe) and the right column comprises only the subset of those plants already naturalized outside of Europe. Naturalization debt refers to the proportion of predicted naturalizations that are not yet observed, sensitivity to the proportion of correctly predicted successful naturalizations and negative predictive value to the proportion of correctly predicted unsuccessful naturalizations. A region was considered suitable if it contained at least one suitable pixel, 1% suitable cells, 5, 20, 50 and 100%



We used LMMs accounting for phylogeny to test whether plant-feature values explain: (a) the relative extent of range filling, that is, the proportion of suitable regions where a species has become naturalized; and (b) the relative extent of range expansion, that is, the proportion of unsuitable regions where a species has nevertheless become naturalized. The indices of range filling and expansion that we used are similar to the indices used by Petitpierre et al. (2012). However, we calculated them in the geographical space instead of in the environmental space because occurrence data on naturalized plants are not evenly comprehensive across regions (which would have been needed to offer a reliable fit of the niches), and when an alien species is reported, we do not always know whether or not it reproduces in the wild (Figure 1).

### 3 | RESULTS

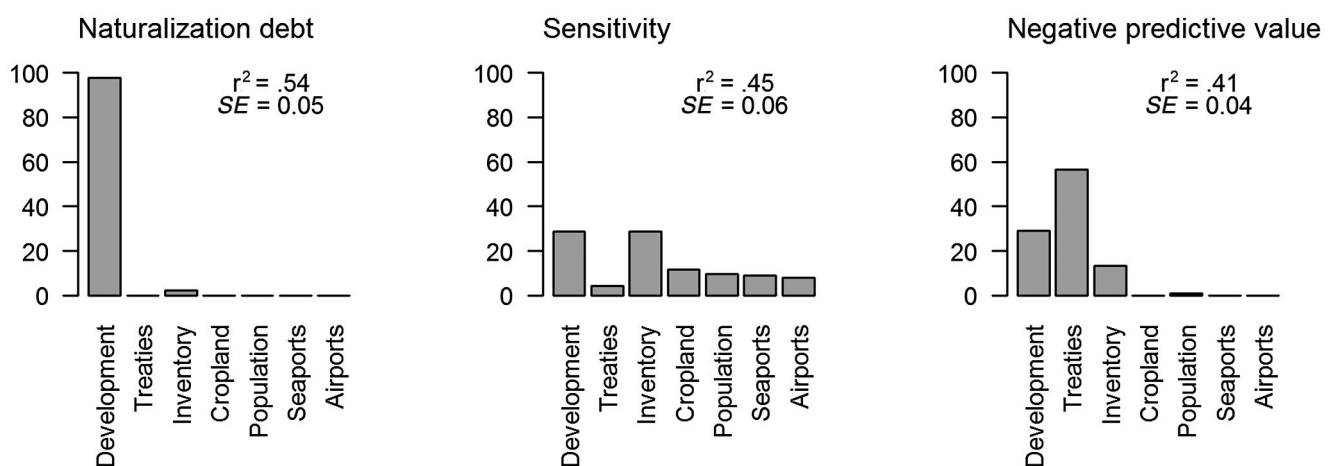
#### 3.1 | Socio-economic drivers of naturalization patterns

European endemic vascular plant species have naturalized in 319 non-European regions. The naturalization debt (i.e. the proportion of species that are not yet naturalized in a particular region but likely to become so, if introduced, given their environmental envelope) ranged from 95 to 100% (mean = 99%) among the 931 non-European regions according to the threshold of one pixel and from 71 to 100% (mean = 99% as well) according to the threshold of 100% of suitable cells (Figure 2). This means that less than 5% and less than 29% of species for which a region is suitable according to the lowest and highest thresholds, respectively, have actually been recorded there. When restricting the analysis to 272 European species already naturalized outside of Europe, the naturalization debt decreased but

remained relatively high with values in the range 81–100% (mean = 96%) with the former threshold and 52–100% (95%) with the latter threshold.

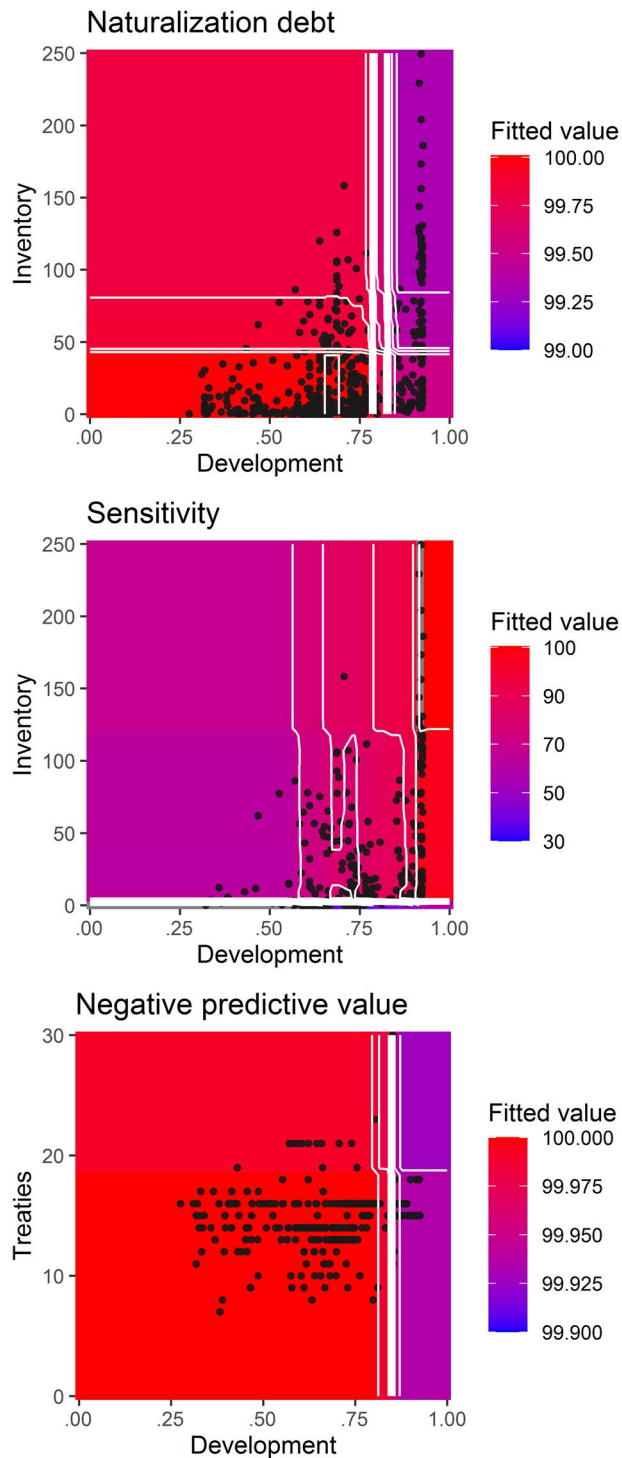
The proportion of observed naturalizations that are correctly predicted based on environmental matching (sensitivity) ranged from 0 to 100% for all thresholds and both species sets. However, the average value of sensitivity was two percentage points lower for the full set of species (19–64%) than for the subset of already naturalized species (21–66%). The average proportion of unsuccessful naturalizations in unsuitable environments (negative predictive value) was between 98.8 and 100% (mean = 99.9%) with the lowest threshold, and between 97.0 and 100% (mean = 99.7%) with the highest threshold for the full set of species. When considering already naturalized species only, the negative predictive value decreased substantially to the range 80.0–100% (mean = 99.4%) in the former case and 83.4–100% (mean = 98.8%) in the latter case. The threshold of 1% of suitable cells offered the best solution for maximizing sensitivity while still having a high negative predictive value when comparing observed and expected naturalized floras. Therefore, we used predictions from this threshold in the subsequent analyses.

Naturalization debt was lower in more developed regions (human development index > .7; Figures 3 and 4) with moist and cool climates, including coastal North America, Japan, temperate Australia and New Zealand (Figure 5). Sensitivity increased progressively with the level of development and inventory effort (Figures 3 and 4), and reached maximum values in north-east America, the South American Atlantic Forest and the southern Andes, Turkey, the east-central African mountains, South Africa, Japan, temperate Australia and New Zealand (Figure 5). The negative predictive value was lower in more developed regions (human development index > .7), with more treaties relevant to biological



**FIGURE 3** Relative influence of seven socio-economic factors in explaining differences between observed and expected European naturalized floras calculated by boosted regression trees. Naturalization debt refers to the proportion of predicted naturalizations that are not yet observed, sensitivity to the proportion of correctly predicted successful naturalizations and negative predictive value to the proportion of correctly predicted unsuccessful naturalizations. Socio-economic factors are sorted by mean relative influence over the three metrics. These results are for the upscaling method '1%' that is, a region was considered suitable for a species if it contained at least 1% suitable cells

invasions (number of treaties > 20; Figures 3 and 4), including regions in North America, the southern tip of South America and Africa, most of northern Asia excluding Mongolia, temperate Australia and New Zealand (Figure 5).



**FIGURE 4** Joint partial dependence plots of interactions fitted by boosted regression trees between the most contributing regional socio-economic factors and differences between observed and expected European naturalized floras. Naturalization debt refers to the proportion of predicted naturalizations that are not yet observed, sensitivity to the proportion of correctly predicted successful naturalizations and negative predictive value to the proportion of correctly predicted unsuccessful naturalizations. Black dots represent empirical data points. These results are for the upscaling method '1%' that is, a region was considered suitable for a species if it contained at least 1% suitable cells [Colour figure can be viewed at [wileyonlinelibrary.com](http://wileyonlinelibrary.com)]

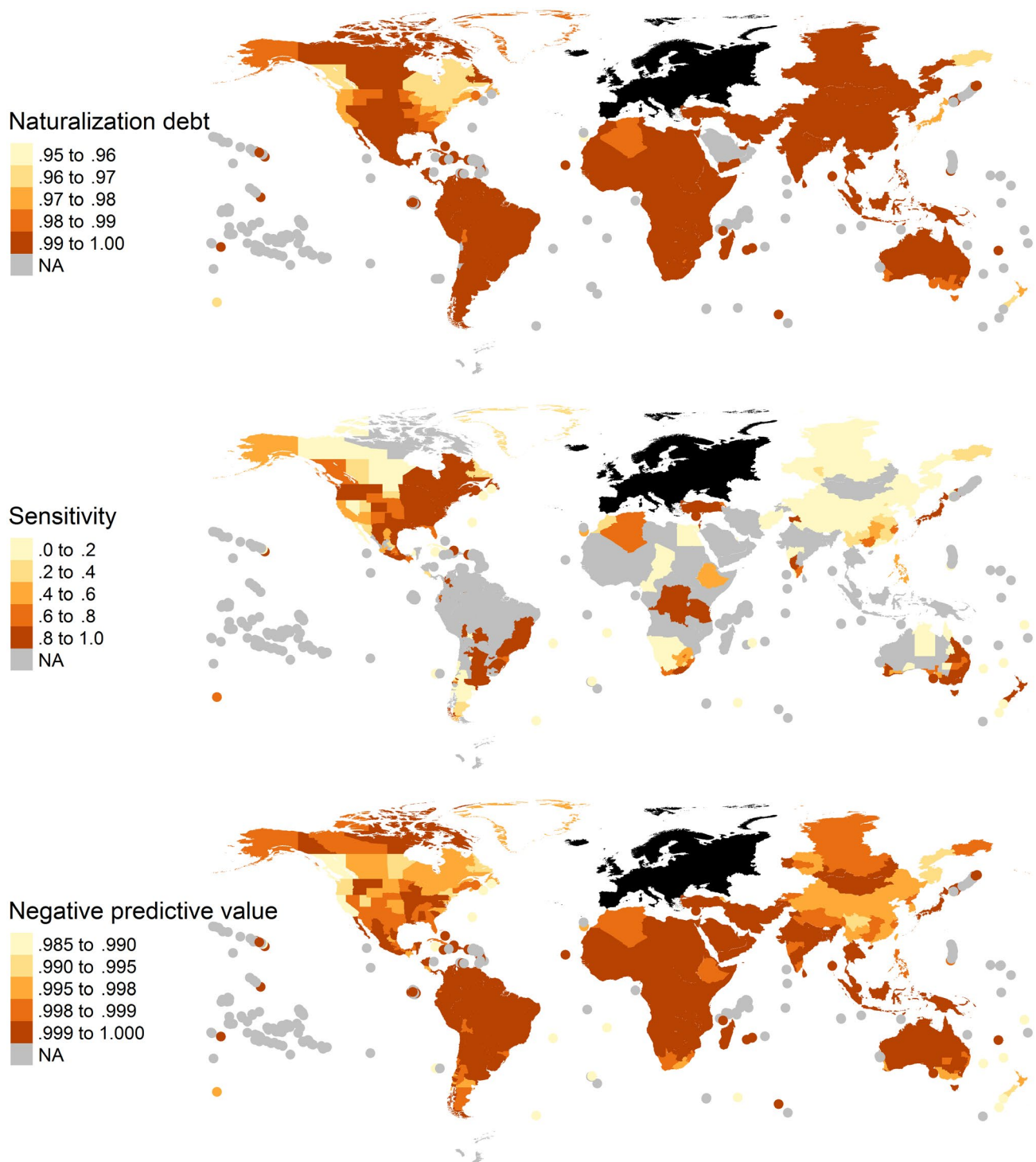
### 3.2 | Plant features explaining naturalization patterns

Compared to plant species that are currently not known to be naturalized somewhere else, those that are naturalized tend to be more widespread in Europe, to have a wider expected range outside Europe, and to have an ornamental or other economic use (Figure 6). There was no significant difference in functional traits between naturalized and non-naturalized species.

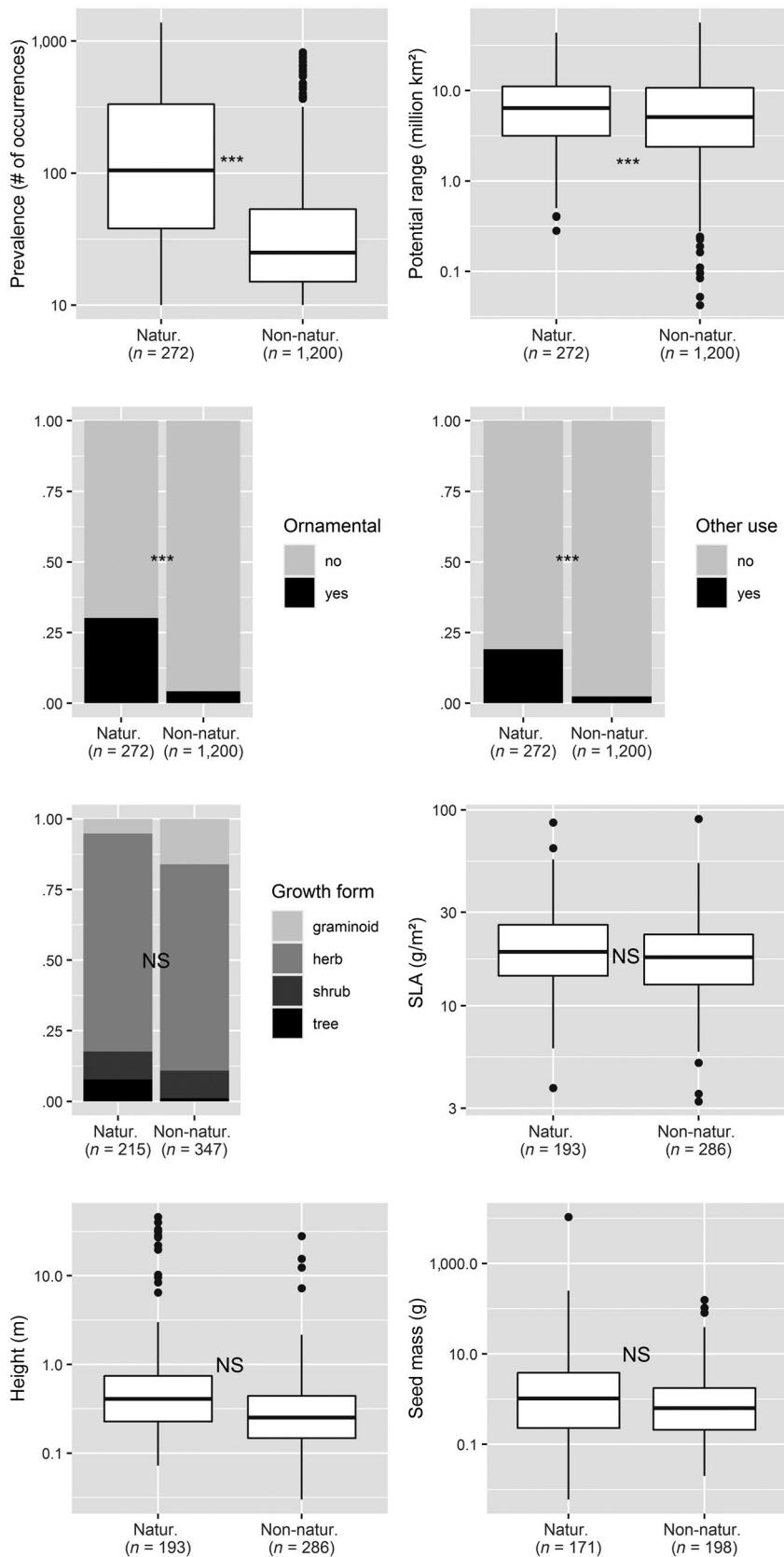
On average, species were found to be naturalized in 4.2% of their potentially suitable regions outside Europe (Supporting Information Figure S4). Thirty of the species occupied less than 0.1% of suitable regions and 31 species occupied more than 10% of suitable regions. The species with the most complete range filling was *Syringa vulgaris* (Oleaceae), native to the Balkan Peninsula and Romania, being currently naturalized in 70% of its suitable regions (Supporting Information Table S2). The average range expansion into regions estimated to be unsuitable was 0.4%. 118 species (43%) exhibited range expansion above 0.1%, but only one species had a range expansion above 10%: *Verbascum virgatum* (Scrophulariaceae), native to south-western Europe and southern England, and currently naturalized in 10.5% of the regions that were predicted to be unsuitable for it. We found that species performing best at filling their expected range have an ornamental or other economic relevance, a higher SLA, and a narrower expected range than species with a less complete range filling (Table 2). Furthermore, species observed as naturalized beyond their expected range tend to have an ornamental or other economic use and to be less widespread in their native range than species with no range expansion (Table 2).

## 4 | DISCUSSION

The global geographical pattern of richness in naturalized plant species from Europe is non-random as supported by our results, originating from a complex interplay of at least four mechanisms: (a) environmental matching between the native range of individual species and their expected alien range, which largely controls unsuccessful naturalization; (b) the alteration of the environment in the alien ranges by socio-economic activities (after accounting for sampling effort); (c) introduction pressure associated with the economic



**FIGURE 5** Global maps of the differences between observed and expected European naturalized floras. Regions with no Global Naturalized Alien Flora (GloNAF) data are displayed in white and Europe in black. Naturalization debt refers to the proportion of predicted naturalizations that are not yet observed, sensitivity to the proportion of correctly predicted successful naturalizations (there are many NA values as many regions have no observed naturalizations by European endemic plants) and negative predictive value to the proportion of correctly predicted unsuccessful naturalizations. These results are for the upscaling method '1%' that is, a region was considered suitable for a species if it contained at least 1% suitable cells [Colour figure can be viewed at [wileyonlinelibrary.com](https://onlinelibrary.wiley.com)]



**FIGURE 6** Differences in distributional, economic and functional features between European vascular plants already recorded as naturalized outside Europe ('Natur.') and those not recorded as naturalized outside Europe ('Non-natur.'), 1,472 species in total. 'Prevalence' refers to the number of  $0.42^\circ \times 0.42^\circ$  cells in Europe occupied; 'Potential range' to the surface area of the species' expected range outside Europe; 'Ornamental' and 'Other use' indicate whether species are used as an ornamental plant or has other economic relevance (e.g. food plant, medicines, materials), respectively; 'Growth form' reflects the morphology of a plant, especially its physiological adaptation to the environment; 'SLA' for 'specific leaf area' is the ratio of leaf area to leaf dry mass; 'Height' and 'Seed mass' are the plant height and seed mass, respectively. Significance levels of the relationships fitted with linear mixed-effect models accounting for between-species phylogenetic distances are indicated by asterisks ( $.05 < p\text{-value} \leq .01$ : \*;  $.01 < p\text{-value} \leq .001$ : \*\*;  $p\text{-value} < .001$ : \*\*\*)

**TABLE 2** Regression coefficients of the relationship between distributional, economic and functional features of naturalized plants from Europe and their range filling (proportion of suitable regions where species are observed as naturalized) and range expansion (proportion of unsuitable regions where species are observed as naturalized). Continuous variables were standardized to allow comparison. Significance of the relationships fitted with linear mixed-effect models accounting for between-species phylogenetic distances is indicated in bold and significance levels are indicated by asterisks

	Range filling	Range expansion	n
Prevalence in Europe	.13	-.40*	272
Potential range outside Europe	-.25**	.08	272
Ornamental	.22***	.14*	272
Other us	.21***	.22***	272
Growth form	.02	.01	193
Specific leaf area (SLA)	.27*	-.01	117
Height	.05	.01	175
Seed mass	.02	-.01	156

\*.05 < p-value ≤ .01.; \*\*.01 < p-value ≤ .001.; \*\*\*p-value < .001.

use of the species; and (d) differences in functional traits (e.g. SLA), which cause some species to more completely fill their expected alien ranges than others.

#### 4.1 | Accounting for environmental matching to decipher patterns of naturalization

We modelled environmental tolerance of species and then the influence of extrinsic socio-economic drivers (Figure 1), instead of both together as done in previous studies (Dawson et al., 2017; Essl et al., 2015). This allowed us to separate the effect of environmental matching in explaining the global pattern of richness in naturalized plant species from Europe. We elucidated that the environment alone is able to correctly predict up to two thirds of successful naturalizations (i.e. sensitivity) and 99.9% of unsuccessful naturalizations (i.e. negative predictive value), depending on the method used to compare local expected distributions with regional observed distributions ('upscaling method'; Figure 2). Based on a similar approach, Bellard et al. (2013) showed that even the so-called 'world's 100 worst invasive alien species' have a quite restricted expected range, covering mainly Europe and the areas along the Atlantic coast of North America. Thus, it clearly appears that knowing the available suitable environmental space of species is critical to assess the factors that determine their naturalization success.

Our approach indicated that non-European regions are currently occupied by less than 5% of the endemic European plants for which the area, or parts of it, would be suitable (Figure 2). Therefore, all regions have an enormous naturalization debt. This debt suggests that many endemic European plants have not been introduced outside of Europe yet or that they still have not overcome the biotic barriers provided by herbivores, pathogens and native competitors or the lack of mutualists. Nevertheless, the large environmentally suitable ranges outside of Europe confirm that European plants have considerable potential for naturalization outside their native range (Pyšek et al., 2017; van Kleunen et al., 2015), once they have overcome dispersal barriers and biotic resistance. Our results, therefore,

indicate that it is unlikely that the number of new naturalizations will soon have reached its peak.

SDMs are increasingly used as a basis to implement biosecurity policies, for example, by drawing up watch lists or lists of undesirable species (e.g. Padayachee et al., 2019). As biosecurity measures are usually taken at a regional scale while SDMs predict expected distributions at smaller scales, it is critical to determine which portion of a region should be suitable for an alien species to become naturalized, and hence, of concern. Our retrospective approach comparing SDM outputs with the current naturalized flora from Europe in 931 non-European regions showed that a threshold in the order of 1% of a focal region offers a good trade-off between maximizing correctly predicted successful and unsuccessful naturalizations (Figure 2). Nevertheless, it may be preferable to choose a lower threshold with the precautionary principle, and this threshold might change when using a different spatial resolution.

#### 4.2 | Socio-economic drivers explaining differences from expected patterns of naturalization

Our results revealed that naturalization debt mostly decreases with the level of development (Figures 3 and 4), which means that a higher proportion of species with the ability to become naturalized are observed as such in more developed regions. In these regions, the distribution of naturalized species is typically closer to an equilibrium with the environment. This result is in line with Pyšek et al. (2010) and Dawson et al. (2017), who found that human impact (represented by proxies like wealth and per capita GDP, respectively) plays a leading role in driving the global patterns of animal and plant naturalizations. One explanation is that economic wealth is often associated with alien species introductions and with profound alteration of natural habitats, which benefit alien species establishment and growth (MacDougall & Turkington, 2005; Seebens et al., 2015).

Surprisingly, unlike other studies (Dawson et al., 2017; Tingley et al., 2018), proxies of international transport (i.e. density of airports and seaports in our study) did not explain naturalization debt (Figures 3 and 4). Nevertheless, we found that regions with the



lowest naturalization debt are all either coastal or island regions (coastal North America, Algeria, Japan, temperate Australia and New Zealand; Figure 5). As a result, although naturalization debt seems little influenced by the density of seaports, it is likely that the volume of maritime traffic, or at least the presence of seaports, does play a significant role by increasing the rates of species introductions. One might argue that this pattern also reflects that coastal regions are usually more strongly developed than more inland regions as a consequence of the increasing demand for infrastructures to sustain residential, commercial and tourist activities (Supporting Information Figure S2), which offers more opportunities for introduction and naturalization of alien plants in coastal regions (Gallardo et al., 2015). We found that naturalization debt was higher for the full set of 1,485 European species (either already naturalized or not yet naturalized) than for the subset of 272 species already naturalized, which are more likely to have overcome dispersal barriers (Figure 2). This suggests that introduction pathways influence naturalization debt although, unfortunately, it remains difficult to assess by how much.

In addition to the influence of development and potentially of introduction pathways on naturalized range filling, sensitivity was also found to be affected by the magnitude of inventory effort (Figures 3 and 4). Specifically, regions with the highest quality records of native plant richness also tend to have more records of alien plants. This is also corroborated by Dawson et al. (2017), and indicates that the geographical coverage of reported plant naturalizations remains unevenly exhaustive across regions and thus that our findings might be significantly influenced by biases in recording effort. In contrast, the negative predictive value decreased in regions with more treaties relevant to invasive alien species (Figures 3 and 4). This most likely does not indicate that treaties promote invasions, but that it is more likely that a region adopts a treaty when it has many invasive species. The effectiveness of these relatively recent treaties in preventing new invasions can only be assessed in the future.

Unexpectedly, human population density and the proportion of cropland had a relatively low contribution to explaining the global pattern of naturalization. In contrast to this, high human population densities are thought to influence the likelihood of an introduced species becoming naturalized and spreading through human impacts on the environment (Essl et al., 2011; Pyšek et al., 2010). Similarly, intensive agriculture (associated with historical sowing or planting of now naturalized species and associated weed-seed contaminants) is widely recognized to be a major cause of new introductions of naturalized and invasive species (Bellard et al., 2016; Perrings et al., 2005; Seebens et al., 2015). A potential explanation for the low contributions of population density and the proportion of cropland to driving patterns of naturalization could be its redundancy with the proportion of primary land cover used as an environmental variable in SDMs since urbanization and agriculture are largely responsible for land conversion (Chini et al., 2014). Although this redundancy makes it difficult to isolate the effects of human population density and agriculture, we believe

the introduction of primary land cover into our SDMs was critical in this study to distinguish plants that are drivers and passengers of human-induced disturbances (MacDougall & Turkington, 2005). Another possible cause for the low contribution of agriculture is the ancient human colonization history of Europeans who used to transport cultivated plants across continents while other regions only recently opened up to international movements of plants (di Castri, 1989). In this perspective, most of the plants used in agriculture and associated weeds endemic to Europe with the potential to become naturalized might have already spread widely (Monnet et al., 2020).

### 4.3 | Plant features explaining differences from expected patterns of naturalization

The average score of alien geographical range filling of naturalized plants from Europe was 4.2%. This might appear to be very low compared to percentages of environmental niche filling reported in other studies (Hill et al., 2017; Liu et al., 2020; Petitpierre et al., 2012; Strubbe et al., 2013, 2015). However, it should be noted that it is easier to fill a niche than to fill all locations that have that niche. None of our species had a range filling above the commonly used threshold of 90%. Similarly, no invasive birds showed more than 90% of their native niche filled in the invasive range (Strubbe et al., 2013). However, no less than 52% of invasive plants (Petitpierre et al., 2012), 32% of 22 insects (Hill et al., 2017) and 10% of vertebrates (Strubbe et al., 2015) had a niche filling above 90%. Moreover, only one naturalized plant species from Europe out of 272 had a range expansion above the commonly used threshold of 10%, while it was the case for the niche expansion of 55% of invasive insects (Hill et al., 2017), 29% of birds (Strubbe et al., 2013), 17% of vertebrates (Strubbe et al., 2015) and 14% of invasive plants (Petitpierre et al., 2012).

We found that SLA was positively correlated with range filling of naturalized plants from Europe (Table 2). According to the LHS scheme of Westoby (1998), SLA is a proxy for rapid growth and an acquisitive resource-use strategy, which would be typical of resource-rich and/or disturbed environments, both of which have become more common in recent centuries due to human activities. In the competitor/stress-tolerator/ruderal (CSR) framework of the universal adaptive strategy theory (Grime, 1977), rapid growth is a key characteristic of strong competitors, suggesting that strong competitors for light and nutrients are also filling their naturalized ranges faster. Indeed, Guo et al. (2018) recently showed that species with high scores on the competitor axis were more likely to naturalize. Thus, it seems very consistent that having a high SLA represents a key strategy in regions subject to intensive development associated with rapid, deep and widespread anthropogenic perturbations. SLA should, therefore, also indicate potential for faster expansion towards the equilibrium range in alien regions. This corroborates previous results showing that SLA is higher for naturalized and invasive species compared to non-naturalized or native species (Pyšek & Richardson, 2007; Pyšek et al., 2015; van Kleunen et al., 2010).

Interestingly, seed mass was not significantly related to range filling (Table 2). This trait can have contrasting effects on patterns of naturalization since large seeds have greater nutrient reserves, which increases the chances of becoming naturalized, but small seeds are beneficial for long-distance dispersal and therefore favour spread (Crawley et al., 1996; Moodley et al., 2013). This suggests that it is not a species' dispersal ability or the ability to settle in new regions that mainly drive naturalization but the ability to be transported by humans for aesthetic or economic values. Our results show that species used by humans are indeed more frequently naturalized (Figure 6), and perform better at filling (e.g. *Syringa vulgaris*) and expanding (e.g. *Verbascum virgatum*) their potential range than species not used by humans (Table 2). Ornamental horticulture, agriculture and forestry are recognized as major pathways of alien plant introduction. For instance, it has been estimated that at least 75 and 93%, respectively, of the naturalized alien plants worldwide are grown in domestic and botanical gardens (van Kleunen et al., 2018), and plants with a known economic importance are 18 times more likely to be naturalized somewhere in the world (van Kleunen et al., 2020). Possible reasons include: economic importance increases the number of releases and/or the number of individuals released (i.e. propagule pressure), while human selection favours traits related to invasiveness (e.g. ease to grow and reproduce, early and long flowering period, low susceptibility to insect pests or pathogens).

Moreover, we found that the same set of distributional features explains whether or not species are naturalized and the extent of naturalized range filling and geographical expansion of species range (range filling and expansion were significantly correlated;  $r = .61$ ,  $p$ -value  $< .001$ ), but in opposite directions. Indeed, naturalized species tend to be more widespread in their native range and to have wider expected alien ranges than non-naturalized species, that is, they are typically generalist species (Gallien et al., 2019). In contrast, naturalized species with the most complete range filling and broadest range extension tend to have more limited expected alien ranges and to be less widespread in their native range, respectively. This result needs to be interpreted with caution since it can simply arise from a numerical artefact as it may be easier to fill a small expected range than a large one. Another possible cause could be that SDMs tend to be more accurate for specialist species than for generalists (Connor et al., 2018). Furthermore, range filling depends on opportunities to disperse, so species with a wider expected alien range may basically have less chance to fill it. Finally, more narrowly distributed endemics can surprisingly have a greater range expansion if their observed distribution in the native range results from biogeographical (e.g. the Mediterranean Basin, the Alps in Europe) or from biological interactions rather than from climatic barriers. In that case SDMs would fail to fit their fundamental climatic envelope due to niche truncation (Bush et al., 2018). It could also be that their current distribution matches their climatic niche, but that this distribution is not correctly reflected in data sources, maybe because they mostly inhabit under-sampled regions. Consequently, the projected expected alien ranges may underestimate the true potential ranges.

## 5 | CONCLUSIONS

Our findings indicate that the environment largely controls the expected naturalized range of European plants. We demonstrated that increased anthropogenic disturbance associated with human development as well as increased probability of introductions associated with international exchange primarily explain the extent to which species spread within their expected naturalized range. We showed that plants selected for ornamental and other economic purposes perform better at filling and expanding their range, most likely because of increased introduction pressure and a preference for cultivating plants with traits that also make them more likely to naturalize. We revealed that species with functional traits indicative of rapid growth and acquisitive resource use tend to fill their range more completely than species with slower responses. These findings give a new overall picture of the drivers of naturalization that can help plan future studies on the macroecology of alien species as well as in designing future biosecurity plans. As harmful invasive species emerge from the pool of naturalized species, the analysis of which species could naturalize is a first step in identifying species that could ultimately become invasive in a region.

## ACKNOWLEDGMENTS

Robin Pouteau was supported by the National Natural Science Foundation of China (grant number 31901176) and Taizhou University (2018YQ001). Carsten Meyer acknowledges funding by the Volkswagen Foundation through a Freigeist Fellowship (A118199), and additional support by iDiv, funded by the German Research Foundation (DFG-FZT 118, 202548816). Petr Pyšek and Jan Pergl were supported by the Czech Science Foundation (EXPRO grant 19-28807X) and the Czech Academy of Sciences (long-term research development project RVO 67985939). Idoia Biurrun and Juan Antonio Campos were supported by the Basque Government (IT936-16). Gianmaria Bonari, Milan Chytrý, Ilona Knollová, Anna Kuzemko and Corrado Marcenò were supported by the Czech Science Foundation (19-28491X). Rosario G. Gavilán was supported by Madrid Government project REMEDINAL (P2018/EMT-4338). Jens-Christian Svenning considers this work a contribution to his VILLUM Investigator project 'Biodiversity Dynamics in a Changing World' funded by VILLUM FONDEN (16549). Mark van Kleunen was supported by the German Research Foundation DFG (264740629). The authors appreciate the helpful comments by three anonymous referees and the Handling Editor, Melodie A. McGeoch.

## AUTHOR CONTRIBUTIONS

Robin Pouteau and Mark van Kleunen conceived the idea for this manuscript with input from Wilfried Thuiller. All authors provided the data, Robin Pouteau performed the analysis and wrote the first draft of the manuscript with help from Wilfried Thuiller and Mark van Kleunen, and all authors contributed substantially to revisions.

## DATA AVAILABILITY STATEMENT

The data used to build the species distribution models (occurrence records and environmental variables) came from openly accessible repositories cited in the manuscript (for details on the EVA database, see <http://euroveg.org/eva-database-obtaining-data>). All 1,485 generated habitat suitability raster files are available at <https://dataverse.ird.fr/dataset.xhtml?persistentId=doi:10.23708/RNGS8Z>

## ORCID

Robin Pouteau  <https://orcid.org/0000-0003-3090-6551>  
 Caroline Brunel  <https://orcid.org/0000-0002-0705-5165>  
 Barry J. Conn  <https://orcid.org/0000-0002-7156-4802>  
 Aleksandr L. Ebel  <https://orcid.org/0000-0002-7889-4580>  
 Holger Kreft  <https://orcid.org/0000-0003-4471-8236>  
 Carsten Meyer  <https://orcid.org/0000-0003-3927-5856>  
 Jan Pergl  <https://orcid.org/0000-0002-0045-1974>  
 Petr Pyšek  <https://orcid.org/0000-0001-8500-442X>  
 Patrick Weigelt  <https://orcid.org/0000-0002-2485-3708>  
 Svetlana Aćić  <https://orcid.org/0000-0001-6553-3797>  
 Emiliano Agrillo  <https://orcid.org/0000-0003-2346-8346>  
 Ariel Bergamini  <https://orcid.org/0000-0001-8816-1420>  
 Christian Berg  <https://orcid.org/0000-0002-0587-3316>  
 Erwin Bergmeier  <https://orcid.org/0000-0002-6118-4611>  
 Idoia Biurrun  <https://orcid.org/0000-0002-1454-0433>  
 Steffen Boch  <https://orcid.org/0000-0003-2814-5343>  
 Gianmaria Bonari  <https://orcid.org/0000-0002-5574-6067>  
 Zoltán Botta-Dukát  <https://orcid.org/0000-0002-9544-3474>  
 Helge Bruehlheide  <https://orcid.org/0000-0003-3135-0356>  
 Juan Antonio Campos  <https://orcid.org/0000-0001-5992-2753>  
 Andraž Čarni  <https://orcid.org/0000-0002-8909-4298>  
 Laura Casella  <https://orcid.org/0000-0003-2550-3010>  
 Maria Laura Carranza  <https://orcid.org/0000-0001-5753-890X>  
 Milan Chytrý  <https://orcid.org/0000-0002-8122-3075>  
 Michele De Sanctis  <https://orcid.org/0000-0002-7280-6199>  
 Jürgen Dengler  <https://orcid.org/0000-0003-3221-660X>  
 Jörg Ewald  <https://orcid.org/0000-0002-2758-9324>  
 Rosario G. Gavilán  <https://orcid.org/0000-0002-1022-445X>  
 Ute Jandt  <https://orcid.org/0000-0002-3177-3669>  
 Florian Jansen  <https://orcid.org/0000-0002-0331-5185>  
 Borja Jiménez-Alfaro  <https://orcid.org/0000-0001-6601-9597>  
 Anna Kuzemko  <https://orcid.org/0000-0002-9425-2756>  
 Tatiana Lysenko  <https://orcid.org/0000-0001-6688-1590>  
 Corrado Marcenò  <https://orcid.org/0000-0003-4361-5200>  
 Vasilij Martynenko  <https://orcid.org/0000-0002-9071-3789>  
 Jesper Erenskjold Moeslund  <https://orcid.org/0000-0001-8591-7149>  
 Ricarda Pätzsch  <https://orcid.org/0000-0002-3349-0910>  
 Remigiusz Pielech  <https://orcid.org/0000-0001-8879-3305>  
 Pavel Shirokikh  <https://orcid.org/0000-0003-1864-4878>  
 Jozef Šibík  <https://orcid.org/0000-0002-5949-862X>  
 Angela Stanisci  <https://orcid.org/0000-0002-5302-0932>  
 Zvezdana Stančić  <https://orcid.org/0000-0002-6124-811X>  
 Jens-Christian Svenning  <https://orcid.org/0000-0002-3415-0862>

Grzegorz Swacha  <https://orcid.org/0000-0002-6380-2954>  
 Pavel Dan Turtureanu  <https://orcid.org/0000-0002-7422-3106>  
 Milan Valachovič  <https://orcid.org/0000-0001-5296-5240>  
 Kiril Vassilev  <https://orcid.org/0000-0003-4376-5575>  
 Mark van Kleunen  <https://orcid.org/0000-0002-2861-3701>

## REFERENCES

- Abellán, P., Tella, J. L., Carrete, M., Cardador, L., & Anadón, J. D. (2017). Climate matching drives spread rate but not establishment success in recent unintentional bird introductions. *Proceedings of the National Academy of Sciences USA*, 114, 9385–9390. <https://doi.org/10.1073/pnas.1704815114>
- Allouche, O., Tsoar, A., & Kadmon, R. (2006). Assessing the accuracy of species distribution models: Prevalence, kappa and the true skill statistic (TSS). *Journal of Applied Ecology*, 46, 1223–1232. <https://doi.org/10.1111/j.1365-2664.2006.01214.x>
- Bellard, C., Leroy, B., Thuiller, W., Rysman, J.-F., & Courchamp, F. (2016). Major drivers of invasion risks throughout the world. *Ecosphere*, 7, e01241. <https://doi.org/10.1002/ecs2.1241>
- Bellard, C., Thuiller, W., Leroy, B., Genovesi, P., Bakkenes, M., & Courchamp, F. (2013). Will climate change promote future invasions? *Global Change Biology*, 19, 3740–3748. <https://doi.org/10.1111/gcb.12344>
- Blackburn, T. M., Pyšek, P., Bacher, S., Carlton, J. T., Duncan, R. P., Jarošík, V., Wilson, J. R. U., & Richardson, D. M. (2011). A proposed unified framework for biological invasions. *Trends in Ecology and Evolution*, 26, 333–339. <https://doi.org/10.1016/j.tree.2011.03.023>
- Bombi, P., & D'Amen, M. (2014). Scaling down distribution maps from atlas data: A test of different approaches with virtual species. *Journal of Biogeography*, 39, 640–651. <https://doi.org/10.1111/j.1365-2699.2011.02627.x>
- Bush, A., Catullo, R. A., Mokany, K., Thornhill, A. H., Miller, J. T., & Ferrier, S. (2018). Truncation of thermal tolerance niches among Australian plants. *Global Ecology and Biogeography*, 27, 22–31. <https://doi.org/10.1111/geb.12637>
- Cayuela, L., Stein, A., & Oksanen, J. (2017). *Taxonstand: taxonomic standardization of plant species names v.2.1*. R Foundation for Statistical Computing. <https://cran.r-project.org/web/packages/Taxonstand/index.html>. Accessed on 21 March 2020.
- Chini, L. P., Hurr, G. C., & Frolking, S. (2014). *Harmonized global land use for years 1500–2100, V1*. Data set. Oak Ridge National Laboratory Distributed Active Archive Center, USA. <http://daac.ornl.gov>. Accessed on 21 March 2020.
- Chytrý, M., Hennekens, S. M., Jiménez-Alfaro, B., Knollová, I., Dengler, J., Jansen, F., Landucci, F., Schaminée, J. H. J., Aćić, S., Agrillo, E., Ambarli, D., Angelini, P., Apostolova, I., Attorre, F., Berg, C., Bergmeier, E., Biurrun, I., Botta-Dukát, Z., Brisse, H., ... Yamalov, S. (2016). European vegetation archive (EVA): An integrated database of European vegetation plots. *Applied Vegetation Science*, 19, 173–180. <https://doi.org/10.1111/avsc.12191>
- Connor, T., Hull, V., Viña, A., Shortridge, A., Tang, Y., Zhang, J., Wang, F., & Liu, J. (2018). Effects of grain size and niche breadth on species distribution modeling. *Ecography*, 41, 1270–1282. <https://doi.org/10.1111/ecog.03416>
- Crawley, M. J., Harvey, P. H., & Purvis, A. (1996). Comparative ecology of the native and alien floras of the British Isles. *Philosophical Transactions of the Royal Society B: Biological Sciences*, 351, 1251–1259.
- Dawson, W., Moser, D., van Kleunen, M., Kreft, H., Pergl, J., Pyšek, P., Weigelt, P., Winter, M., Lenzner, B., Blackburn, T. M., Dyer, E. E., Cassey, P., Scrivens, S. L., Economo, E. P., Guénard, B., Capinha, C., Seebens, H., García-Díaz, P., Nentwig, W., ... 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, R. (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). John Wiley.
- Dormann, C. F., Elith, J., Bacher, S., Buchmann, C., Carl, G., Carré, G., Marquéz, J. R. G., Gruber, B., Lafourcade, B., Leitão, P. J., Münkemüller, T., McClean, C., Osborne, P. E., Reineking, B., Schröder, B., Skidmore, A. K., Zurell, D., & Lautenbach, S. (2013). Collinearity: A review of methods to deal with it and a simulation study evaluating their performance. *Ecography*, 36, 27–46. <https://doi.org/10.1111/j.1600-0587.2012.07348.x>
- Elith, J., Leathwick, J. R., & Hastie, T. (2008). A working guide to boosted regression trees. *Journal of Animal Ecology*, 77, 802–813. <https://doi.org/10.1111/j.1365-2656.2008.01390.x>
- Essl, F., Dullinger, S., Rabitsch, W., Hulme, P. E., Hülber, K., Jarošík, V., Kleinbauer, I., Krausmann, F., Kühn, I., Nentwig, W., & Vilà, M. (2011). Socioeconomic legacy yields and invasion debt. *Proceedings of the National Academy of Sciences USA*, 108, 203–207.
- Essl, F., Dawson, W., Kreft, H., Pergl, J., Pyšek, P., van Kleunen, M., Weigelt, P., Mang, T., Dullinger, S., Lenzner, B., Moser, D., Maurel, N., Seebens, H., Stein, A., Weber, E., Chatelain, C., Inderjit, Genovesi, P., Kartesz, J., ... Winter, M. (2019). Drivers of the relative richness of naturalized and invasive plant species on Earth. *AoB PLANTS*, 11, plz051. <https://doi.org/10.1093/aobpla/plz051>
- Essl, F., Dullinger, S., Moser, D., Steinbauer, K., & Mang, T. (2015). Macroecology of global bryophyte invasions at different invasion stages. *Ecography*, 38, 488–498. <https://doi.org/10.1111/ecog.00905>
- Feng, Y., Maurel, N., Wang, Z., Ning, L., Yu, F.-H., & van Kleunen, M. (2016). Introduction history, climatic suitability, native range size, species traits and their interactions explain establishment of Chinese woody species in Europe. *Global Ecology and Biogeography*, 25, 1356–1366. <https://doi.org/10.1111/geb.12497>
- Gallardo, B., Zieritz, A., & Aldridge, D. C. (2015). The importance of the human footprint in shaping the global distribution of terrestrial, freshwater and marine invaders. *PLoS ONE*, 10, e0125801. <https://doi.org/10.1371/journal.pone.0125801>
- Gallien, L., Münkemüller, T., Albert, C. H., Boulangeat, I., & Thuiller, W. (2010). Predicting potential distributions of invasive species: Where to go from here? *Diversity and Distributions*, 16, 331–342.
- Gallien, L., Thornhill, A. H., Zurell, D., Miller, J. T., & Richardson, D. M. (2019). Global predictors of alien plant establishment success: Combining niche and trait proxies. *Proceedings of the Royal Society B: Biological Sciences*, 286, 20182477.
- Grime, J. P. (1977). Evidence for the existence of three primary strategies in plants and its relevance to ecological and evolutionary theory. *The American Naturalist*, 111, 1169–1194. <https://doi.org/10.1086/283244>
- Guo, W.-Y., van Kleunen, M., Winter, M., Weigelt, P., Stein, A., Pierce, S., Pergl, J., Moser, D., Maurel, N., Lenzner, B., Kreft, H., Essl, F., Dawson, W., & Pyšek, P. (2018). The role of adaptive strategies in plant naturalization. *Ecology Letters*, 21, 1380–1389. <https://doi.org/10.1111/ele.13104>
- Gurevitch, J., Scheiner, S. M., & Fox, G. A. (2006). *The ecology of plants* (2nd ed.). Sinauer.
- Hengl, T., de Jesus, J. M., MacMillan, R. A., Batjes, N. H., Heuvelink, G. B. M., Ribeiro, E., Samuel-Rosa, A., Kempen, B., Leenaars, J. G. B., Walsh, M. G., & Gonzalez, M. R. (2014). SoilGrids1km—Global soil information based on automated mapping. *PLoS ONE*, 9, e105992. <https://doi.org/10.1371/journal.pone.0105992>
- Hill, M. P., Gallardo, B., & Terblanche, J. S. (2017). A global assessment of climatic niche shifts and human influence in insect invasions. *Global Ecology and Biogeography*, 26, 679–689. <https://doi.org/10.1111/geb.12578>
- Hobohm, C. (2014). *Endemism in vascular plants*. Plant and vegetation 9 Springer.
- Hulme, P. (2009). Trade, transport and trouble: Managing invasive species pathways in an era of globalization. *Journal of Applied Ecology*, 46, 10–18. <https://doi.org/10.1111/j.1365-2664.2008.01600.x>
- IPBES (Intergovernmental Science-Policy Platform on Biodiversity and Ecosystem Services). (2019). *Global assessment report on biodiversity and ecosystem services of the Intergovernmental Science-Policy Platform on Biodiversity and Ecosystem Services* (E. S. Brondizio, J. Settele, S. Díaz & H. T. Ngo [Eds.]). Bonn: IPBES Secretariat.
- Karger, D. N., Conrad, O., Böhrer, J., Kawohl, T., Kreft, H., Soria-Auza, R. W., Zimmermann, N. E., Linder, H. P., & Kessler, M. (2017). Climatologies at high resolution for the earth's land surface areas. *Scientific Data*, 4, 170122. <https://doi.org/10.1038/sdata.2017.122>
- Kattge, J., Bönsch, G., Díaz, S., Lavorel, S., Prentice, I. C., Leadley, P., Tautenhahn, S., Werner, G. D. A., Aakala, T., Abedi, M., Acosta, A. T. R., Adamidis, G. C., Adamson, K., Aiba, M., Albert, C. H., Alcántara, J. M., Alcázar, C. C., Aleixo, I., Ali, H., ... Wirth, C. (2020). TRY plant trait database—enhanced coverage and open access. *Global Change Biology*, 26, 119–188. <https://doi.org/10.1111/gcb.14904>
- Liu, C., Wolter, C., Xian, W., & Jeschke, J. M. (2020). Most invasive species largely conserve their climatic niche. *Proceedings of the National Academy of Sciences USA*, 117, 23643–23651. <https://doi.org/10.1073/pnas.2004289117>
- MacDougall, A. S., & Turkington, R. (2005). Are invasive species the drivers or passengers of change in degraded ecosystems? *Ecology*, 86, 42–55.
- Meyer, C., Weigelt, P., & Kreft, H. (2016). Multidimensional biases, gaps and uncertainties in global plant occurrence information. *Ecology Letters*, 19, 992–1006. <https://doi.org/10.1111/ele.12624>
- Milbau, A., Stout, J. C., Graae, B. J., & Nijs, I. (2009). A hierarchical framework for integrating invasibility experiments incorporating different factors and spatial scales. *Biological Invasions*, 11, 941–950. <https://doi.org/10.1007/s10530-008-9306-2>
- Monnet, A. C., Vorontsova, M. S., Govaerts, R. H. A., Svenning, J.-C., & Sandel, B. (2020). Historical legacies and ecological determinants of grass naturalizations worldwide. *Ecography*, 43, 1373–1385. <https://doi.org/10.1111/ecog.04609>
- Moodley, D., Geerts, S., Richardson, D. M., & Wilson, J. R. U. (2013). Different traits determine introduction, naturalization and invasion success in woody plants: Proteaceae as a test case. *PLoS ONE*, 8, e75078. <https://doi.org/10.1371/journal.pone.0075078>
- Padayachee, A. L., Proches, S., & Wilson, J. R. U. (2019). Prioritising potential incursions for contingency planning: Pathways, species, and sites in Durban (eThekweni), South Africa as an example. *Neobiota*, 47, 1–21. <https://doi.org/10.3897/neobiota.47.31959>
- Perrings, C., Dehnen-Schmutz, K., Touza, J., & Williamson, M. (2005). How to manage biological invasions under globalization. *Trends in Ecology Evolution*, 20, 212–215. <https://doi.org/10.1016/j.tree.2005.02.011>
- Petitpierre, B., Kueffer, C., Broennimann, O., Randin, C., Daehler, C., & Guisan, A. (2012). Climatic niche shifts are rare among terrestrial plant invaders. *Science*, 335, 1344–1348. <https://doi.org/10.1126/science.1215933>
- Pinheiro, J., Bates, D., & R Core Team. (2019). *nlme: Linear and non-linear mixed effects models v.3.1-141*. R Foundation for Statistical Computing. <https://cran.r-project.org/web/packages/nlme/index.html>. Accessed on 21 March 2020.
- Pyšek, P., Jarošík, V., Hulme, P. E., Kuhn, I., Wild, J., Arianoutsou, M., Bacher, S., Chiron, F., Didžiulis, V., Essl, F., Genovesi, P., Gherardi, F., Hejda, M., Kark, S., Lambdon, P. W., Desprez-Loustau, M.-L., Nentwig, W., Pergl, J., Poboljsaj, K., ... Winter, M. (2010). Disentangling the role of environmental and human pressures on biological invasions across Europe. *Proceedings of the National Academy of Sciences USA*, 107, 12157–12162. <https://doi.org/10.1073/pnas.1002314107>
- Pyšek, P., Manceur, A. M., Alba, C., McGregor, K. F., Pergl, J., Štajerová, K., Chytrý, M., Danihelka, J., Kartesz, J., Klimešová, J., Lučanová, M., Moravcová, L., Nishino, M., Sádlo, J., Suda, J., Tichý, L., & Kühn, I. (2015). Naturalization of central European plants in North America: Species traits, habitats, propagule pressure, residence time. *Ecology*, 96, 762–774. <https://doi.org/10.1890/14-1005.1>



- Pyšek, P., Pergl, J., Essl, F., Lenzner, B., Dawson, W., Kreft, H., Weigelt, P., Winter, M., Kartesz, J., Nishino, M., Antonova, L. A., Barcelona, J. F., Cabelas, F. J., Cárdenas, D., Cárdenas-Toro, J., Castaño, N., Chacón, E., Chatelain, C., Dullinger, S., ... 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, 203–274. <https://doi.org/10.23855/preslia.2017.203>
- Pyšek, P., & Richardson, D. M. (2007). Traits associated with invasiveness in alien plants: where do we stand? In W. Nentwig (Ed.), *Biological invasions* (pp. 97–125). Springer-Verlag.
- R Development Core Team (2020). *R: A language and environment for statistical computing*. R Foundation for Statistical Computing.
- Richardson, D. M., & Pyšek, P. (2012). Naturalization of introduced plants: Ecological drivers of biogeographical patterns. *New Phytologist*, 196, 383–396. <https://doi.org/10.1111/j.1469-8137.2012.04292.x>
- Richardson, D. M., Pyšek, P., Rejmanek, M., Barbour, M. G., Panetta, F. D., & West, C. J. (2000). Naturalization and invasion of alien plants: Concepts and definitions. *Diversity and Distributions*, 6, 93–107. <https://doi.org/10.1046/j.1472-4642.2000.00083.x>
- 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, 445–456. <https://doi.org/10.1111/ddi.12408>
- Seebens, H., Blackburn, T. M., Dyer, E. E., Genovesi, P., Hulme, P. E., Jeschke, J. M., Pagad, S., Pyšek, P., Winter, M., Arianoutsou, M., Bacher, S., Blasius, B., Brundu, G., Capinha, C., Celesti-Grampow, L., Dawson, W., Dullinger, S., Fuentes, N., Jäger, H., ... Essl, F. (2017). No saturation in the accumulation of alien species worldwide. *Nature Communications*, 8, 14435. <https://doi.org/10.1038/ncomms14435>
- Seebens, H., Essl, F., Dawson, W., Fuentes, N., Moser, D., Pergl, J., Pyšek, P., van Kleunen, M., Weber, E., Winter, M., & Blasius, B. (2015). Global trade will accelerate plant invasions in emerging economies under climate change. *Global Change Biology*, 21, 4128–4140. <https://doi.org/10.1111/gcb.13021>
- Stohlgren, T. J., Barnett, D. T., Jarnevich, C. S., Flather, C., & Kartesz, J. (2008). The myth of plant species saturation. *Ecology Letters*, 11, 313–322. <https://doi.org/10.1111/j.1461-0248.2008.01153.x>
- Strubbe, D., Beauchard, O., & Matthysen, E. (2015). Niche conservatism among non-native vertebrates in Europe and North America. *Ecography*, 38, 321–329. <https://doi.org/10.1111/ecog.00632>
- Strubbe, D., Broennimann, O., Chiron, F., & Matthysen, E. (2013). Niche conservatism in non-native birds in Europe: Niche unfilling rather than niche expansion. *Global Ecology and Biogeography*, 22, 962–970. <https://doi.org/10.1111/geb.12050>
- Thuiller, W., Lafourcade, B., Engler, R., & Araújo, M. B. (2009). BIOMOD—a platform for ensemble forecasting of species distributions. *Ecography*, 32, 369–373. <https://doi.org/10.1111/j.1600-0587.2008.05742.x>
- Tingley, R., García-Díaz, P., Rocha Arantes, C. R., & Cassey, P. (2018). Integrating transport pressure data and species distribution models to estimate invasion risk for alien stowaways. *Ecography*, 41, 635–646. <https://doi.org/10.1111/ecog.02841>
- van Kleunen, M., Dawson, W., Essl, F., Pergl, J., Winter, M., Weber, E., Kreft, H., Weigelt, P., Kartesz, J., Nishino, M., Antonova, L. A., Barcelona, J. F., Cabezas, F. J., Cárdenas, D., Cárdenas-Toro, J., Castaño, N., Chacón, E., Chatelain, C., Ebel, A. L., ... Pyšek, P. (2015). Global exchange and accumulation of non-native plants. *Nature*, 525, 100–103. <https://doi.org/10.1038/nature14910>
- van Kleunen, M., Essl, F., Pergl, J., Brundu, G., Carboni, M., Dullinger, S., Early, R., González-Moreno, P., Groom, Q. J., Hulme, P. E., Kueffer, C., Kühn, I., Máguas, C., Maurel, N., Novoa, A., Parepa, M., Pyšek, P., Seebens, H., Tanner, R., ... Dehnen-Schmutz, K. (2018). The changing role of ornamental horticulture in alien plant invasions. *Biological Reviews*, 93, 1421–1437. <https://doi.org/10.1111/brv.12402>
- van Kleunen, M., Pyšek, P., Dawson, W., Essl, F., Kreft, H., Pergl, J., Weigelt, P., Stein, A., Dullinger, S., König, C., & Lenzner, B. (2019). The Global Naturalized Alien Flora (GloNAF) database. *Ecology*, 100, e02542.
- van Kleunen, M., Weber, E., & Fisher, M. (2010). A meta-analysis of trait differences between invasive and non-invasive plant species. *Ecology Letters*, 13, 235–245. <https://doi.org/10.1111/j.1461-0248.2009.01418.x>
- van Kleunen, M., Xu, X., Yang, Q., Maurel, N., Zhang, Z., Dawson, W., Essl, F., Kreft, H., Pergl, J., Pyšek, P., Weigelt, P., Moser, D., Lenzner, B., & Frisette, T. S. (2020). Economic use of plants is key to their naturalization success. *Nature Communications*, 11, 3201. <https://doi.org/10.1038/s41467-020-16982-3>
- Westoby, M. (1998). A leaf-height-seed (LHS) plant ecology strategy scheme. *Plant and Soil*, 199, 213–227.
- Wiersma, J. H., & León, B. (2013). *World economic plants: A standard reference*. CRC Press. Interactive database available online at <https://npgsweb.ars-grin.gov/gringlobal/taxon/taxonomysearcheco.aspx>. Accessed on 21 March 2020.
- Xu, W.-B., Svenning, J.-C., Chen, G.-K., Zhang, M.-G., Huang, J.-H., Chen, B., Ordóñez, A., & Ma, K.-P. (2019). Human activities have opposing effects on distributions of narrow-ranged and widespread plant species in China. *Proceedings of the National Academy of Sciences USA*, 116, 26674–26681. <https://doi.org/10.1073/pnas.1911851116>
- Zimmermann, N. E., Yoccoz, N. G., Edwards, T. C., Meier, E. S., Thuiller, W., Guisan, A., Schmatz, D. R., & Pearman, P. B. (2009). Climatic extremes improve predictions of spatial patterns of tree species. *Proceedings of the National Academy of Sciences USA*, 106, 19723–19728. <https://doi.org/10.1073/pnas.0901643106>

## BIOSKETCH

This work is a collaboration of two international working groups: Global Naturalized Alien Flora (GloNAF) and European Vegetation Archive (EVA), together with external collaborators.

**Robin Pouteau** is a plant ecologist at IRD (French Research Institute for Sustainable Development) with a strong focus on the impact of biological invasions on biodiversity. This work is part of his postdoctoral research at Taizhou University.

## SUPPORTING INFORMATION

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

**How to cite this article:** Pouteau R, Thuiller W, Hobohm C, et al. Climate and socio-economic factors explain differences between observed and expected naturalization patterns of European plants around the world. *Global Ecol Biogeogr*. 2021;30:1514–1531. <https://doi.org/10.1111/geb.13316>