Plant extinctions and introductions lead to phylogenetic and taxonomic homogenization of the European flora

Marten Wintera,b,1, Oliver Schweigerb, Stefan Klotzb, Wolfgang Nentwigc, Pavlos Andriopoulosd, Margarita Arianoutsoufd, Corina Basnoud, Pinelopi Delipetrouf, Viktoras Didžiulisg, Martin Hejdh, Philip E. Hulmei, Philip W. Lambdona, Jan Pergil, Petr Pyšekh,k, David B. Royli, and Ingolf Künn

aDepartment of Community Ecology, UFZ-Helmholtz Centre for Environmental Research, D-06120 Halle (Saale), Germany; bEcology and Evolution Unit, Department of Biology, University of Fribourg, CH-1700 Fribourg, Switzerland; cInstitute of Ecology and Evolution, University of Bern, CH-3012 Bern, Switzerland; dDepartment of Ecology and Systematics, Faculty of Biology, School of Sciences, University of Athens, 15784 Athens, Greece; eCenter for Ecological Research and Forestry Applications, Universitat Autònoma de Barcelona, 08193 Bellatera, Spain; fDepartment of Botany, Faculty of Biology, School of Sciences, University of Athens, 15784 Athens, Greece; gCoastal Research and Planning Institute, Klaipeda University, LT 92294 Klaipeda, Lithuania; hInstitute of Botany, Academy of Sciences of the Czech Republic, CZ-25243 Průhonice, Czech Republic; iBio-Protection Research Centre, Lincoln University, Canterbury, New Zealand; Kew Herbarium, Royal Botanic Gardens, Richmond, Surrey TW9 3AB, United Kingdom; jNatural Environment Research Council Centre for Ecology and Hydrology Wallingford, Crownmarsh Gifford, Wallingford OX10 88B, United Kingdom

Edited by Harold A. Mooney, Stanford University, Stanford, CA, and approved October 28, 2009 (received for review June 26, 2009)

Human activities have altered the composition of biotas through two fundamental processes: native extinctions and alien introductions. Both processes affect the taxonomic (i.e., species identity) and phylogenetic (i.e., species evolutionary history) structure of species assemblages. However, it is not known what the relative magnitude of these effects is at large spatial scales. Here we analyze the large-scale effects of plant extinctions and introductions on taxonomic and phylogenetic diversity of floras across Europe, using data from 23 regions. Considering both native losses and alien additions in concert reveals that plant invasions since AD 1500 exceeded extinctions, resulting in (i) increased taxonomic diversity (i.e., species richness) but decreased phylogenetic diversity within European regions, and (ii) increased taxonomic and phylogenetic similarity among European regions. Those extinct species were phylogenetically and taxonomically unique and typical of individual regions, and extinctions usually were not continent-wide and therefore led to differentiation. By contrast, because introduced alien species tended to be closely related to native species, the floristic differentiation due to species extinction was lessened by taxonomic and phylogenetic homogenization effects. This was especially due to species that are alien to a region but native to other parts of Europe. As a result, floras of many European regions have partly lost and will continue to lose their uniqueness. The results suggest that biodiversity needs to be assessed in terms of both species taxonomic and phylogenetic identity, but the latter is rarely used as a metric of the biodiversity dynamics.

Globalization is progressively altering the composition of biotas worldwide (1–3). The interplay of two fundamental processes — extinctions of native species and introductions and successful establishment of alien species (sensu 4; hereafter referred to as invasion) — has been known to reduce the distinctiveness of species communities. Global species extinctions lead to a continuous decrease of overall species richness (i.e., γ-diversity) (5). However, at the scale of continents, regions, and countries, invasions exceed local extinctions and result in an increase in local or regional species richness (i.e., α-diversity) (5–9).

Changes in species composition driven by the combined effects of invasions and extinctions can result in decreasing (i.e., homogenization), increasing (i.e., differentiation), or unchanged compositional turnover of species (i.e., β-diversity) (10, 11) and traits between and within continents (12, 13). The higher magnitude of invasions compared with extinctions is known to lead to decreased β-diversity between regions at continental (14–17) and regional scales (18–21).

Previous evidence of biotic homogenization at continental and regional scales has largely examined impacts on the taxonomic structure of species assemblages, yet extinctions and invasions might also affect phylogenetic structure (22). The phylogenetic structure of a species assemblage represents the evolutionary history of its members and reflects the diversity of genetic and thus morphologic, physiologic, and behavioral characteristics (23). High phylogenetic diversity within and across communities may enable rapid adaptation to changing environmental conditions across both ecologic and evolutionary time scales (24, 25).

Research on the relative effects of extinctions and invasions upon the structure (α-diversity) and spatial distribution (β-diversity) of phylogenetic patterns may thus enhance our understanding of how evolutionary and ecologic factors contribute to general diversity patterns (26, 27).

The composition of species invasions and extinctions (28) is not randomly distributed among plant taxa but reflects the response of specific life-history traits to natural and human-induced environmental change (29, 30). Extinctions usually befall specialized endemic or rare species, often from species-poor families, which form distinct parts of biotas (31, 28). Hence, their loss should result in a pronounced decrease of phylogenetic and taxonomic β-diversity within and between regions to which these species are unique (32, 33). Successful invaders are often ecologic generalists with wide distributions (34), often belonging to species-rich families (35). Hence, the gain of such common species should also decrease phylogenetic and taxonomic β-diversity within and between regions.

Recent studies of the effect of alien species on the phylogenetic composition of plant communities revealed ambiguous results; species composition can be modified by the addition of either (i) more distantly related species than expected by chance


The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

Freely available online through the PNAS open access option.

1To whom correspondence should be addressed. E-mail: marten.winter@ufz.de.

This article contains supporting information online at www.pnas.org/cgi/content/full/0907088106/DCSupplemental.
(phylogenetic overdispersion) (36) or (ii) more closely related species (phylogenetic clustering) (37); and see ref. 38 for review.

Generally, very little is known about the relative effects of extinctions and invasions on phylogenetic relationships among species at large spatial scales (27). On the basis of a comprehensive dataset of original (before extinctions and introductions) and current (after extinctions and introductions) floras across several European regions, here we use phylogenetic information to assess the consequences of species extinctions and introductions on taxonomic and phylogenetic diversity.

Results and Discussion

Since AD 1500, the processes of species extinctions and invasions, acting in concert, resulted in a net increase in overall European plant species richness (γ-diversity) (Table 1). This increase is due to 69 extinctions of European plants and 1,621 invasions of plants from outside Europe (Fig. S1 and Table S1) and is also accompanied by increased phylogenetic γ-diversity. The increased species richness at the European level is reflected by higher species richness in all European regions (α-diversity), where species invasions also exceeded extinctions (Table S1). However, the prevalence of invasions over extinctions decreased rather than increased phylogenetic α-diversity over the same period; an additional decrease was caused by extinctions (both \( P < 0.001 \); Table 1, Table S1, and Table S2).

Despite increased species richness at the European and regional level, extinctions and invasions in concert resulted in decreased β-diversity among species (\( \beta_{\text{tax}} \)) and β-diversity among phylogenetic lineages (\( \beta_{\text{phy}} \)) of European regions (Table 1), indicating that European floras became phylogenetically and taxonomically impoverished. A decrease of phylogenetic richness with increasing species richness was previously reported on a national scale (39).

However, considering extinctions and invasions separately revealed contrasting patterns. Invasions have generally led to taxonomic and phylogenetic homogenization, whereas species extinctions result in differentiation. Species extinctions increased \( \beta_{\text{tax}} \) and \( \beta_{\text{phy}} \) (Fig. 1B), although \( \beta_{\text{tax}} \) and \( \beta_{\text{phy}} \) of extinct species were higher than those of extant native species (\( P < 0.001 \)). Species from distantly related taxonomic branches have become extinct in different regions. By contrast, invasions decreased \( \beta_{\text{tax}} \) and \( \beta_{\text{phy}} \) of European floras (Fig. 1C), although \( \beta_{\text{tax}} \) and \( \beta_{\text{phy}} \) of alien species was higher than those of native species (both \( P < 0.001 \)). Hence, although alien floras are taxonomically and phylogenetically diverse, the constituent species are either native to other European regions or, if introduced from outside Europe, tend to be closely related to native European species.

We reason that many species that have become extinct or have been introduced in a particular region are components of the current native floras of other European regions, because (i) the loss of taxonomically more diverse native species (extinctions) led to increased \( \beta_{\text{tax}} \) and the gain of more diverse aliens led to decreased \( \beta_{\text{tax}} \), and (ii) invasions of phylogenetically more diverse alien species decreased \( \beta_{\text{phy}} \) and the loss of more diverse extirpate species increased \( \beta_{\text{phy}} \) among floras. This is because extinctions of plant species in Europe are mainly regional rather than continent-wide extinctions. Moreover, ca. 53% of plant invasions in European floras are due to species exchange among European regions (40). Such a high percentage is likely to increase taxonomic and phylogenetic similarities among European floras.

Generally, the negative effect of aliens on \( \beta_{\text{tax}} \) and \( \beta_{\text{phy}} \) is approximately seven times the magnitude of the positive effect of species losses (compare \( \beta_{\text{tax}} \) values, Fig. 1B and C). Therefore, the taxonomic and phylogenetic homogenization among European floras due to species invasions masks any regional differentiation due to species extinctions. These patterns are consistent with those found for North America, where species extinctions only played a minor role in defining compositional patterns for state floras, owing to the low numbers of extinct species (41). Although extinction processes are much slower than invasions, it is unlikely that accounting for lag effects in extinctions (8) would change the pattern dramatically.

With compositional changes due to extinctions and invasions, \( \beta_{\text{tax}} \) increased linearly with increasing \( \beta_{\text{phy}} \) (Fig. 1A; \( R^2 = 0.86, P < 0.001 \)). However, this relationship is not inevitable (27); the high correlation indicates that species determining this pattern are probably widespread and closely related to extant native species. The weaker relationship between \( \beta_{\text{tax}} \) and \( \beta_{\text{phy}} \) derived from the effect of extinctions indicates that there were more phylogenetically unique species among extinct native plants. In contrast to the main patterns, among some regions we did observe occasions when species introductions led to taxonomic and phylogenetic differentiation, and in some circumstances extinctions led to increased homogenization (as species that are unique to only one region are lost). However, we did not find any systematic trend in these patterns, and they were not related to geographic distance, species richness, or other specific attributes of the floras (e.g., level of endemism).

As with α-diversity, two scenarios whereby \( \beta_{\text{tax}} \) and \( \beta_{\text{phy}} \) show contrasting patterns (homogenization vs. differentiation) are possible (27): (i) a high \( \beta_{\text{tax}} \) (differentiation) and a low \( \beta_{\text{phy}} \) (homogenization) could be observed with a high proportion of resident endemic species or if the different communities consist of close congeners; (ii) a low \( \beta_{\text{tax}} \) (homogenization) and a high \( \beta_{\text{phy}} \) (differentiation) is very unlikely, because a high species overlap will always generate a high phylogenetic overlap (27).

Effects of scale dependencies are known for biodiversity patterns in general (42) and invasion processes in particular (43, 44). Phylogenetic structures could also be scale dependent (45). It has been argued that large-scale patterns of phylogenetic clustering reflect biogeographic rather than ecologic processes (38). Moreover, the perception of taxonomic homogenization is dependent on the spatial scale at which samples are gathered and

| Table 1. Taxonomic and phylogenetic diversity components across Europe |
|--------------------------|--------------------------|--------------------------|
| **Level**                | **Diversity**            |
|                          | \( \gamma \)              | \( \beta \)              | \( \alpha \)              |
| **Taxonomic level**      |                          |                          |
| Original flora           | 10,928                   | 0.5432 ± 0.1622          | 2,194 ± 1194             |
| Current total flora      | 12,624                   | 0.5155 ± 0.1490          | 2,664 ± 1215             |
| **Phylogenetic level**   |                          |                          |
| Original flora           | 1.3797                   | 1.4493 ± 0.1049          | 1.4772 ± 0.0202          |
| Current total flora      | 1.3942                   | 1.4462 ± 0.0951          | 1.4725 ± 0.0189          |

\( \gamma \) applies to overall European diversity, whereas \( \beta \) and \( \alpha \) are mean values (± SD) across European regions. Taxonomic diversities are based on species numbers (\( \gamma \) and \( \alpha \)) and \( \beta_{\text{tax}} \) values. Phylogenetic diversities are based on \( \Delta^{*} \) (\( \gamma \) and \( \alpha \)) and \( \beta_{\text{phy}} \) values.
increases with increasing sample area (46). Similarly, phylogenetic homogenization may also be scale dependent because closely related alien and native species may not cooccur in the same plant communities (47, 48). Thus, using regions may overestimate the degree of homogenization experienced by local plant communities. Furthermore, political boundaries will often encompass various biogeographic regions, further increasing the similarity across these sample regions (49).

In conclusion, we have shown that increasing species numbers in European regional floras over the last 5 centuries have been accompanied by a decrease of phylogenetic and taxonomic uniqueness. To restrict indicators of conservation priorities or ecosystem health simply to species richness can thus be misleading and does not capture the important effects of taxonomic or phylogenetic distinctiveness (22, 54, 55). Even if native diversity is considered separately, the decreasing numbers of native species can have more serious consequences for the phylogenetic diversity than can be inferred from species richness alone. We show that by combining taxonomic and phylogenetic information on \( \alpha \)- and \( \beta \)-diversity, we gain new insight into changes in biodiversity patterns at different levels, which are likely to be relevant for ecosystem processes (37, 27, 39). Phylogenetic diversity reflects the evolutionary history of a community, which may also reflect its functional diversity (23, 26). Hence, diminished phylogenetic and taxonomic information could decrease the capacity of species assemblages to respond to environmental changes and therefore threaten ecosystem functioning (39). Considering the rate of species invasions into Europe (40, 56), the above trends are likely to continue.

**Methods**

**Species Data.** Species occurrence data and their status (alien, native, or extinct) were collected for 23 European countries or regions (e.g., the Baltic States are represented as one region; Table S1; all countries/regions are hereafter referred to as regions). Because extinction and invasion rates of island ecosystems are much higher than those of mainland regions (52), we excluded island data and worked only with mainland data for which the geographic definition of Flora Europaea (57) allowed us to distinguish between mainland and island data (see details in Fig. S1): Greece (Crete, Karpathos, Kasos, Gavdos, and those Aegean Islands outside of Europe defined by Flora Europaea), Spain (Balearic Islands), France (Corse), Italy (Sardinia, Sicily), and the Malta Archipelago.

Lists of native species (see ref. 4 for definition) for European regions were derived from the European Science Foundation European Documentation System database of Flora Europaea (57; http://rbg-web.rbge.org.uk/FE/html). Data on extinct plant species were taken from national Red Lists of alien species per country were compiled from the DAISIE database (58; www.europe-aliens.org). An alien species is defined as one introduced to the region as a result of human activities and successfully naturalized (i.e., forming reproducing populations in the wild) (sensu 4). Only plant species introduced after AD 1500 were considered (4), which are hereafter referred to as “aliens.”

We only considered taxa at the species level. In Europe there are 12,624 species, of which 10,928 are native, 537 extinct, and 3,353 alien. Some species were
assigned to different categories in different regions [e.g., 1,726 species are native to one region but are alien to another (see ref. 40), or 468 native species are extinct in one region but occur in other regions (Table S1)].

**Taxonomic β-Diversity.** We used the Morisita-Horn dissimilarity index (MH) to calculate \( \beta_{tax} \) among floras. The index is computed as:

\[
\beta_{tax} = \text{MH}_{jk} = 1 - \frac{2 \sum_{i} x_{ij} x_{ik}}{(\lambda_{j} + \lambda_{k}) \sum_{i} x_{ij} \sum_{i} x_{ik}}
\]

with \( \lambda_{j} = \left( \sum_{i} x_{ij} \right)^{2} \) and \( \lambda_{k} = \left( \sum_{i} x_{ik} \right)^{2} \),

where \( x_{ij} \) and \( x_{ik} \) represent the frequencies of species \( i \) in regions \( j \) and \( k \), respectively. The index ranges from 0 (total identity between two samples and low β-diversity) to 1 (absolute dissimilarity of these samples and high β-diversity). This index is less sensitive to species richness and sample size than most other indices (59). Because the index needs abundance data, we defined pseudo-abundance of one for each species.

**Phylogenetic β-Diversity.** \( \beta_{phy} \) was defined as β-diversity with a temporal dimension measured as phylogenetic distance between communities according to branch lengths (27). The online software tool PhyloMatic (60) was used to construct a supertree using species and genus data of all species considered. In the absence of phylogenetic branch lengths for the whole tree, we calculated pseudobranch lengths to weight the height of the nodes according to their position in the tree (61). We assigned a relative height (with tips at 0 and root at 1) to each node and then calculated branch lengths as the difference between the heights of two nodes. This ensures that the total branch length from root to any tip is constant. Grafen’s method sets node height from the tip proportional to the number of descendant terminal nodes (taxa) minus 1 (61). To calculate the branch length we used the function `brlen` of the R-package ape (62).

The \( \beta_{phy} \) between two floras was assessed using the PhyloSor index (37). To facilitate the comparison of \( \beta_{phy} \) and \( \beta_{tax} \), \( \beta_{phy} \) was also computed as dissimilarity:

\[
\beta_{phy} = 1 - \text{PhyloSor}_{jk} = \frac{BL_{jk}}{(BL_{j} + BL_{k})} \frac{1}{2}
\]

where \( BL_{jk} \) is the branch length common to communities \( j \) and \( k \), and \( BL_{j} \) and \( BL_{k} \) are the total branch lengths of communities \( j \) and \( k \), respectively. \( \beta_{phy} \) ranges from 0 (both communities are composed of the same taxa) to 1 (two communities share no taxa).

To disentangle the effect of the loss and gain of species on the β-diversity between regions at the species (\( \beta_{tax} \)) and phylogenetic (\( \beta_{phy} \)) level, we separated the processes of species extinctions and invasions. Because comprehensive dates of plant extinctions do not exist, we arbitrary defined that all extinctions occurred after AD 1500. Thus, we defined the “original flora” (before AD 1500) as all extant and extinct native species, the “current native flora” as only extant native species, and the “current total flora” as extant natives and alien species. Introducing these three categories, we calculate different effects of different floral elements on \( \beta_{tax} \) and \( \beta_{phy} \): (i) the combined effect as the difference between the β-diversities of original and current flora \( \Delta_{ii} \) (current total flora) \( \beta_{original} \) and \( \beta_{current total} \); and (ii) the effect of extinct alien species as the difference between β-diversities of original and current native flora \( \beta_{original native} \) and \( \beta_{current native total} \); and (iii) the effect of alien species as the difference between β-diversities of total current flora and current native flora \( \beta_{current total native} \) and \( \beta_{current non native} \). Furthermore, we calculated separately the \( \beta_{tax} \) and \( \beta_{phy} \) of extinct, alien, and native species among regions.

**Alpha Diversity.** Beside taxonomic α-diversity in terms of species numbers (Table S2), we also assessed phylogenetic α-diversity of the original and current floras by using Warwick’s average taxonomic distinctness (\( \Delta_{i} \)) (63). \( \Delta_{i} \) was originally developed on taxonomic relationships but can be adapted to phylogenetic information (64). The index was calculated as:

\[
\Delta_{i} = \frac{\sum_{j} B_{j}}{s(s-1)}
\]

where \( B_{j} \) is built from the distance matrix of species based on branch lengths, and \( s \) is the number of species. The index is based on the sum of branch lengths between species (as provided by the distance matrix) and can be interpreted as the mean distance between two randomly chosen species independent of their distance from the root of the tree. \( \Delta_{i} \) is mathematically unbiased by species richness; that is, it does not automatically increase with sample size and reflects the phylogenetic structure of a subset from a phylogenetic tree best, unlike the majority of other available phylogenetic diversity indices (64). Smaller \( \Delta_{i} \) values indicate that, on average, the assemblage of species is phylogenetically more closely related and less distinct. Analogous to \( \beta_{tax} \) and \( \beta_{phy} \), we calculated (i) combined effect \( \Delta_{i} \) (original flora) \( \beta_{original} \) and \( \beta_{current total} \); (ii) extinct effect \( \Delta_{i} \) (current native total) \( \beta_{current native total} \) and \( \beta_{current native total} \); and (iii) alien effect \( \Delta_{i} \) (current total flora) \( \beta_{current total} \) and \( \beta_{current total} \). Significant differences between all relationships of β-diversity or \( \Delta_{i} \) were assessed using Fisher’s paired comparisons design test (65).

**ACKNOWLEDGMENTS.** We thank Stefan Michalski and Emmanuel Paradis for providing phylogenetic and technical expertise; and the Delivering Alien Invasive Species Inventories for Europe (DAISIE) consortium for providing the data. This study was partly funded by the European Union through the FP 6 projects DAISIE (contract number SSPI-CT-2003–511202) and ALARM (GEC05-CT-2004-00675). M.H., J.P., and P.P. were also supported by Grants AVG60050516 (from the Academy of Sciences of the Czech Republic), MSM0021620828, and LC06073 (both from the Ministry of Education, Youth and Sports of the Czech Republic).
