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Plant diversity drives global patterns of insect invasions

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During the last two centuries, thousands of insect species have been transported (largely inadvertently) and established outside of their native ranges worldwide, some with catastrophic ecological and economic impacts. Global variation in numbers of invading species depends on geographic variation in propagule pressure and heterogeneity of environmental resistance to invasions. Elton's diversity-invasibility hypothesis, proposed over sixty years ago, has been widely explored for plants but little is known on how biodiversity affects insect invasions. Here we use species inventories from 44 land areas, ranging from small oceanic islands to entire continents in various world regions, to show that numbers of established insect species are primarily driven by diversity of plants, with both native and non-native plant species richness being the strongest predictor of insect invasions. We find that at large spatial scales, plant diversity directly explains variation in non-native insect species richness among world regions, while geographic factors such as land area, climate and insularity largely affect insect invasions indirectly via their effects on local plant richness.

Insects display an enormous diversity of life histories and exist in virtually every terrestrial environment across all world regions. Given their immense diversity, it comes as no surprise that insects outnumber all other taxa of non-native animals worldwide¹. Many non-native insect species are notorious, with catastrophic impacts on agriculture, human health and natural ecosystems².

Even though most world biomes are affected by biological invasions, certain regions appear to be more prone to invasions than others. Geographic variation observed in numbers of established non-native species^{3,4} can be attributed both to variation in historical propagule pressure and to habitat characteristics that make certain areas more prone to invasions, i.e. "invasibility"⁵. The concept of biotic resistance describes the effects of community characteristics to promote or inhibit establishment of non-native species.

Both theoretical and experimental studies of invasive plants support the diversity resistance hypothesis, which posits that diverse communities are highly competitive and readily resist invasion⁶. But there is a scale dependency in such relationships; at larger spatial scales (areas > 10 km²) there tends to be an inverse relationship between plant diversity and resistance to plant invasions⁷. Less is known about effects of local diversity on resistance to insect invasions. At small spatial scales, most studies have found that insect abundance and diversity are negatively related to plant diversity though a few have found the opposite^{8–10}. At larger spatial scales, geographic variation in numbers of insect invasions can be substantial¹ but relatively little information exists about how invasibility is related to plant diversity¹¹.

Here we investigate global variation in numbers of invading insect species, seeking explanations for observed patterns. In particular, we explore the role of plant diversity in explaining historical insect invasions. We assemble an extensive data set on numbers of native and non-native insect species in 44 global regions, both mainland and island, and apply structural equation modeling to tease apart the roles of propagule pressure and habitat invasibility. We find that the dominant driver of both native and non-native insect diversity is plant diversity.

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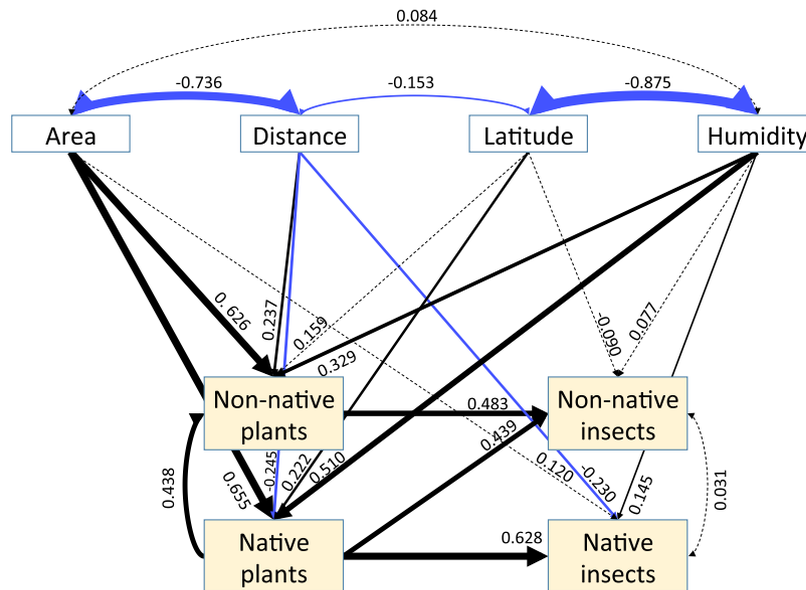


Figure 1. Fit of the reduced structural equation model predicting native and non-native plant and insect species richness. Regression parameter estimates are shown next to arrows; black arrows indicate positive estimates, blue arrows indicate negative estimates and weight of each arrow is proportional to the estimated value. Dashed arrows correspond to non-significant relationships. Distance represents insularity and is measured by distance to the mainland.

Since propagule pressure cannot be directly measured, we evaluated the following potential proxies: occupants (i.e., resident human population), gross domestic product (GDP) and distance to mainland (for continental regions this was set at zero). Potential proxies of invasibility considered were latitude, land area, yearly mean temperature, yearly mean humidity, net primary productivity, number of native plant species and number of naturalized alien plant species.

Results

Pairwise correlations indicate that native and non-native insect species richness are mutually correlated with each other and with most proxies for propagule pressure and invasibility (Supplementary Fig. S1). Given, the high collinearity of these explanatory factors and possibility for complex networks of causality, we applied structural equation modeling (SEM) to investigate these relationships¹².

In the full SEM model (Supplementary Fig. S2), the strongest determinant of non-native insect species richness was native plant richness, though non-native plant richness had a nearly equal effect. Number of human occupants did not have a significant effect on non-native insect richness but both occupants and GDP had strong influences on non-native plant species. Covariances among land area, distance, GDP, occupants and net productivity were generally high which caused an overall low concordance of the model with the data and obscured our ability to differentiate among the effects of these factors.

Following an iterative procedure (see Materials and Methods), a reduced model (Fig. 1) with good concordance with the data was identified. In the reduced model, where occupants, GDP, temperature and productivity are removed, the strongest relationships were the positive effects of land area on non-native and native plant richness. The next strongest relationship was the positive effect of native plant richness on native insects. Again, non-native insect richness was primarily determined by non-native plant richness, followed closely by native plants. Though land area and humidity had strong positive effects on both native and non-native plant richness, they did not have substantial direct effects on either native or non-native insects.

Discussion

Overall, these results indicate the dominance of plant diversity as a driver of insect invasions. While direct competition plays a key role in plant community assembly¹³, it appears to play a lesser role in insects¹⁴. Herbivory is the dominant life history among insect species, including non-native species¹⁵ and most non-herbivorous species use herbivores as hosts. Furthermore, most insect herbivores are either monophagous or oligophagous¹⁶. Consequently, insect community assembly is typically structured around host plant phylogeny¹⁷. The ability of phytophagous insects to colonize new regions is likely dependent upon locating hosts, and consequently regions supporting more diverse plant communities offer greater opportunities for herbivore colonization.

Land area, latitude, climate, and insularity are all well known drivers of global biodiversity^{18,19}. However, our analysis indicates that none of these factors directly affect non-native insect diversity (Fig. 1). These factors more strongly affect plant diversity and thus it appears that their effects on insect diversity mostly operate indirectly via native and non-native plant diversity. Several different mechanisms have been proposed to explain latitudinal gradients of diversity²⁰.

The role of non-native plant diversity as a driver of insect invasions may be related to the phenomenon of “invasional meltdown” in which initial invasions promote subsequent invasions²¹. While invasional meltdown is often considered to occur as a result of invasion-driven disturbance, plant invasions may simply create additional niches for non-native insects to exploit. For example, in Europe 46% of non-native herbivores are limited to non-native host plants²². Many insect herbivores that feed on non-native hosts also utilize native hosts and thus can adversely affect native flora. An important applied implication of these results is that biosecurity efforts aimed at limiting plant invasions may have added benefits in reducing potentially damaging insect invasions as well. Some caution should be used in interpreting the non-native plant/insect invasion association since part of the relationship observed here could have arisen from economic factors (e.g. trade) functioning as drivers of both types of invasions.

Findings reported here of positive effects of plant richness on insect invasions stand in contrast to analyses at smaller spatial scales indicating plant invasions lead to decreased insect diversity^{23,24}. Our results also differ from previous findings that insect abundance and diversity tend to be negatively related to richness of all plant species at small spatial scales^{9,10}. These previous studies are inconsistent with the positive effect of plant diversity on insect invasions observed here but this inconsistency may be analogous to the “invasion paradox” observed in plant invasions in which there is scale dependency in the effect of diversity on biotic resistance to invasions; at large spatial scales invasibility to plants is positively associated with native plant richness, but at smaller scales invasibility generally decreases with native plant richness^{7,25}.

Similar to our report, Hawkins and Porter²⁶ found that native insect herbivore richness was positively associated with plant species richness at large spatial scales. However, they concluded that this did not reflect a direct causal relationship; instead they hypothesized that both plant and insect richness were independently driven by primary productivity and moisture availability. However our analysis indicates the opposite, both native and alien insect diversity are directly determined by plant diversity and the effects of climate and productivity are indirect, operating via plant diversity. Just why our study and theirs have reached contrary conclusions is not yet clear, though differences in the spatial scale of analysis is likely to be a contributing factor, suggesting that investigations of scale effects would be a useful further avenue to explore.

At large spatial scales, plant diversity may increase the number of available niches for insect herbivores and thus promote invasions; this phenomenon is referred to as the “facilitation effect” in the infectious disease literature²⁷. But at smaller spatial scales, plant diversity may impair the ability of insects to locate hosts; this is referred to as the “dilution effect”²⁸ and may cause a negative association between host richness and invasion success. Such opposing effects of facilitation and dilution may explain scale-dependent influences of plant diversity on habitat invasibility to insect invasions.

While the 44 regions analyzed here comprise a diverse sample of land areas from many parts of the world, they are not a random sample and we acknowledge the possibility that unknown bias associated with the areas we have used may affect our conclusions. This analysis utilized all known exhaustive inventories of native and non-native insect species; however, these are most readily available from economically developed countries in the northern hemisphere and selected islands in the southern hemisphere²⁹. Future work, especially on scaling effects, would benefit from a broader data set that includes inventories from less well-investigated regions. The development of a global register of introduced and invasive species is a useful step to promote the availability of such information³⁰.

Materials and Methods

Total numbers of established native and non-native of insect species, as well as numbers of native and non-native naturalized vascular plant species, were derived from a series of species inventories obtained from 44 land areas ranging from small islands to continents (Supplementary Methods online, Supplementary Table S1). For these same regions, we also assembled proxy data on habitat invasibility (land area, latitude, mean annual temperature, mean annual humidity and net primary productivity) and propagule pressure (distance from mainland, occupants and GDP). All data can be viewed in Supplementary Table S2. Numerous pathways facilitate insect invasions so we acknowledge that these proxies do not capture geographical variation in all forms of propagule pressure.

Relationships among propagule pressure proxies, invasibility proxies, native and non-native naturalized plant species richness and native and non-native insect species richness in each of the 44 land areas were quantified using structural equation modeling (SEM). The SEM approach can reveal causal relationships among multiple variables in complex networks (particularly when some of these variables are highly collinear)¹². An *a priori* structural equation model network was identified to represent hypothesized dependencies between predictor and response variables based on logical causal relationships. The full model (Supplementary Fig. 2) included all logical causal connections between invasibility and propagule pressure proxies with native and non-native plant and insect species richness. The model was evaluated using standard SEM procedures¹² implemented with the lavaan package in the R language. All variables (species richness and environmental data) were log-transformed and standardized to stabilize the analyses except latitude was not log-transformed but transformed as absolute values. A maximum likelihood procedure in the lavaan package provides a chi-square statistic that can be used to test the hypothesis of model - data consistency. A value of P (chi-square test) below the standard critical value of 0.05 indicates a significant deviation between observed and model implied covariances. Larger P -values indicate greater model concordance with data and values of $P > 0.05$ are indicative of overall model adequacy^{12,31}.

The full model (Supplementary Fig. 2) provided poor concordance with the data (minimum test statistic = 44.518, $df = 15$, $P = 0.000$). Subsequently, all non-significant connections ($P(>|z|) > 0.5$) were removed from the full model and the revised model was evaluated. Next, each connection that was removed in the previous step was re-evaluated using a modification index (MI), which represents the reduction of the chi-square value associated with removing the connection. Those connections with $MI > 3.84$ (95% percentile of the chi-square distribution with $df = 1$) were retained and checked again in the next step. These steps were repeated until the overall

P-value of the chi-square test was maximized. In cases where inclusion of a specific connection had a negligible effect on the model *P*-value, we also compared AIC values among candidate models and selected the model with the smallest AIC. The final reduced model shown in Fig. 1 provided good concordance with the data (minimum test statistic = 1.692, *df* = 6, *P* = 0.946).

Data and materials availability. The authors declare that all data supporting the findings of this study are available within the article and its Supplementary Information files.

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Author Contributions

A.M.L., T.Y., A.R., S.A., S.L.C., E.G.B. and P.P. contributed to the conception of the study, assembly of data and drafting of the manuscript. A.M.L. and T.Y. performed statistical analyses.

Additional Information

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