The global distribution of bamboos: assessing correlates of introduction and invasion

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Abstract. There is a long history of species being moved around the world by humans. These introduced species can provide substantial benefits, but they can also have undesirable consequences. We explore the importance of human activities on the processes of species dissemination and potential invasions using the Poaceae subfamily Bambusoideae ('bamboos'), a group that contains taxa that are widely utilised and that are often perceived as weedy. We (1) compiled an inventory of bamboo species and their current distributions; (2) determined which species have been introduced and become invasive outside their native ranges; and (3) explored correlates of introduction and invasion. Distribution data were collated from Kew's GrassBase, the Global Biodiversity Information Facility and other online herbarium information sources. Our list comprised 1662 species in 121 genera, of which 232 (14 %) have been introduced beyond their native ranges. Twelve (0.7 % of species) were found to be invasive. A non-random selection of bamboos have been introduced and become invasive outside their native ranges. Asiatic species in particular have been widely introduced. There was a clear over-representation of introduced species in the genera Bambusa and Phyllostachys which also contain most of the listed invasive species. The introduction of species also correlated with certain traits: taxa with larger culm dimensions were significantly more likely to have been moved to new areas; and those with many cultivars had a higher rate of dissemination and invasion. It is difficult to determine whether the patterns of introduction and invasion are due simply to differences in propagule pressure, or whether humans have deliberately selected inherently invasive taxa. In general, we suggest that human usage is a stronger driver of introductions and invasions in bamboos than in other taxa that have been well studied. It is likely that as bamboos are used more widely, the number and impact of invasions will increase unless environmental risks are carefully managed.

Keywords: Bamboo; Bambusoideae; biological invasions; cultivars; introduced species; invasive species; Poaceae.

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Introduction

Human-mediated dissemination of species has intensified over the past three centuries with the increase of global traffic (Meyerson and Mooney 2007; Ricciardi 2007). Some introduced species naturalize (reproduce consistently) in their new ranges and some naturalized species invade (spread from sites of introduction). This has created a global-scale natural experiment in biogeography (Bardsley and Edward-Jones 2006; Richardson 2006; Richardson et al. 2011a; Richardson et al. 2011c; Yoshida et al. 2007). Considerable efforts have been made by invasion scientists to understand the key drivers of invasion, and to determine whether generalisations can be made on how some species manage to overcome barriers associated with different stages of the introduction-naturalization-invasion continuum (Blackburn et al. 2011; Kueffer et al. 2013; Moodley et al. 2016; Richardson and Pyšek 2012). However, as introduced taxa often represents a non-random selection of all taxa, there is some ‘taxonomic selectivity’ in which taxa become invasive (McKinney and Lockwood 1999).

Biological invasions are, by definition, the result of human-mediated dispersal and can only be understood in the context of human activities. The movement of species is often influenced by their direct value to humans (McKinney and Lockwood 1999), in particular as introduced species have been essential to the development of all contemporary human societies (Prance and Nesbitt 2005). With intentional plant introductions, morphological traits have been shown to be important in facilitating the introduction and invasion of species (Pyšek and Richardson 2007). Certain traits may be of high value to humans at the introduction stage and thus influence the initial movement of these species into new ranges. For example, Proteaceae with showy flowers and Cactaceae with other traits valued for ornamentation were found to be overrepresented among introduced species in these families (Moodley et al. 2013; Novoa et al. 2015). For both these families, traits that enabled greater ability to spread were found to be more important for invasion success post-introduction. Traits underlying invasion success can also be highly taxon or context specific. In many woody plant taxa, such as Acacia, Pinus and Proteaceae, seedbank size and longevity are associated with invasion success (Grotkopp et al. 2002; Moodley et al. 2013; Richardson and Kluge 2008), while in Cactaceae growth form is an important determinant of invasion success. (Novoa et al. 2015).

We focused on bamboos, a large subfamily of the grasses (Poaceae: Bambusoideae; 1662 species in 121 genera). Bamboos have a range of functional forms distributed over numerous biogeographic regions, including dwarf herbaceous species found in temperate climates and giant tropical woody species that can grow up to 20 m tall (Bystriakova et al. 2004). It is estimated that 2.5 billion people are directly involved with the production and consumption of bamboo (Scurlock et al. 2000). The main economic value of bamboo lies in the utility of the hardened culm, which serves many of the same functions as timber (Chung and Yu 2002; Scurlock et al. 2000). What makes bamboo a particularly interesting group beyond timber functions, however, is the versatility of uses and the utilisation of all plant parts. Leaves are used for fodder, shoots for human consumption, culms for biomass, construction, textiles, musical instruments and many bamboos are used in horticulture (Hunter 2003). This has led to many species being intentionally moved outside of their native ranges (Cook and Dias 2006; Townsend 2013).

Over the past few decades, bamboos have seen an upsurge in popularity, largely driven by a perception of certain species as wonder plants or miracle crops, i.e. plants that are believed to be especially valuable in meeting current economic, environmental and social needs (Hoogendoorn and Benton 2014; Liese and Köhl 2015). Various authors have argued that commercially grown bamboos are more sustainable and renewable than current forestry crops (Bansal and Zoolagud 2002; Song et al. 2011). Modern processing techniques have also transformed the range of products that can be made from bamboo. Therefore, the rate at which species are being introduced and cultivated in new ranges has increased; especially cultivation of bamboos in response to an increased global demand for timber products (Hunter 2003; INBAR 2003).

Most research on bamboos has focused on aspects of commercial cultivation and uses such as methods for maximizing yields and on providing economic valuations of plantings in different contexts. To date, we are not aware of any comprehensive studies on the invasion ecology of bamboos, despite their reputation for being a group that contains highly ‘invasive’ species (Buckingham et al. 2011; Space and Flynn 2000). Many species possess weedy attributes, such as fast growth rates, clonal reproduction and the formation of long-lived monospecific stands (Lima et al. 2012). Bamboos can dramatically alter ecosystem dynamics through competitive exclusion and expansion of patches that form from clonal reproduction. A growing number of papers address some of these issues (Blundell et al. 2003; Kobayashi et al. 2015; Kudo et al. 2011; Lima et al. 2012; Rother et al. 2016; Suzuki 2015; Yang et al. 2015).

While there has been a long history of bamboo introductions, little is known about which species have been moved where, and the outcomes of these movements. The aims of this paper were to (1) compile an inventory...
of all bamboo species and their current global distribution; (2) determine which species have been introduced and which have become invasive outside of their native ranges; and (3) explore correlates of introduction and invasion.

We expected that certain correlates, both biological (i.e. taxonomy, phylogeny, plant traits) and social (i.e. introduction effort, the utility of species), will have resulted in taxonomic selectivity in introduction effort (Table 1).

**Methods**

**Inventory of species and distribution**

Establishing inventories of taxa, their distribution and cases of invasions are fundamentally important in the field of invasion science and the lack of such information can hinder management efforts (McGeoch et al. 2012). To document the dissemination of bamboos, we required up-to-date taxonomic lists and distribution data.

The identification of bamboos is notoriously problematic (reviewed by Kellogg 2015). Due to the rarity of flowering cycles (7 to more than 120 years in woody species; Janzen 1976), species identification often relies heavily on vegetative material, but most species have few, if any, reliable diagnostic vegetative features. Consequently, there are major discrepancies between the classification of bamboos and species lists. Significant improvements have been made by specialist groups such as the Bamboo Phylogeny Group (2012) and, more generally, by GrassBase, an on-going international initiative to collate taxonomic data on the family Poaceae at the Royal Botanical Gardens, Kew, UK. GrassBase includes a list of all bamboo species, their distributions and trait data (Clayton et al. 2015; Vorontsova et al. 2015). We verified and updated the accepted taxa in GrassBase both as one of us has specialist experience in grass taxonomy (MSV) and by collaborating with a bamboo taxonomy specialist (Lynn G. Clark, Iowa State University). We also included recent literature on new species and other changes in classification published up to September 2015 (Kellogg 2015). [see Supporting Information—Table S1 for full species list].

An extensive search was undertaken between June 2014 and January 2015 to document the introduction of bamboos to areas outside of their native ranges. This included searches of the Web of Science and other platforms of academic and grey literature. Most information was retrieved from online databases specialising in global herbarium records and/or non-native species records, namely the Global Biodiversity Information Facility (GBIF), Kew’s GrassBase, the Global Compendium of Weeds (GCW), Pacific Island Ecosystems at Risk (PIER), Delivering Alien Invasive Species Inventories for Europe (DAISIE), Invasive Species Specialist Group (ISSG) and CABI’s Invasive Species Compendium (CABI-ISC), but independent literature searches also provided useful data [see Supporting Information—Fig. S1]. GBIF provided the greatest amount of data on the locality of species with over 84,000 entries for ‘Bambusoideae’ species. Of these, around 29% of records had sufficient ancillary data for our purposes (of the 71% that did not, 8% lacked a scientific name, 21% a country and 71% a locality).

When pooled with the other databases, 179 species names did not match our accepted species list. Unknown names were removed; synonyms and spelling errors were updated or corrected accordingly and kept in the final database [see Supporting Information—Table S2]. We discarded records on the basis of names that we could not resolve using these criteria. The final list for analyses included over 27,000 entries. Names of geographic regions were defined based on the International Organization for Standardization for country codes and regions (ISO 31661-1 standard; with the exception of a few island regions which were independently defined, such as Hawaii and the Galapagos Islands).

**Dissemination and status**

We categorized the presence of a species in a given country or region as native or non-native (or introduced) based on distribution data from Kew’s GrassBase and cross-referenced with Ohrnberger (1999). These two data sources provide a complete inventory of the taxonomy and distribution of bamboos that was needed to establish native and introduced ranges. We defined these categories using the compendium of concepts in invasion science proposed by Richardson et al. (2011). Species were listed as ‘non-native’ or ‘introduced’ when their presence in a region is due to human activity. Note that our records do not distinguish between successful introductions (where species have established and are still present today) and failed introductions (where species no longer occur in that region)—they simply reflect the presence of a species in a given region at some point in time. We classified a subset of ‘non-native’ species as ‘invasive’. Invasive species are ‘naturalized plants that produce reproductive offspring often in large numbers at a considerable distance from parent plants...’ (Richardson et al. 2011b). Records of bamboos being listed as invasive were found either through the databases mentioned above, or through an independent literature search. References for invasions came from a combination of peer-reviewed literature and official government reports, which were then cross-checked to validate claims that species were ‘invasive’ following the
Table 1 Features correlated with the introduction and invasion status of bamboos.

| Correlate/ Measurement | Expectation | Result | Consequence | Figure/table in this paper |
|------------------------|-------------|--------|-------------|---------------------------|
| Taxonomy (genera)      | Introduced species will tend to come from certain genera | The genera *Bambusa*, *Phyllostachys*, *Semiarundinaria*, *Shibataea*, and *Thysostachys* had a significant proportion of species that have been introduced; and *Bambusa*, *Phyllostachys* and *Pleioblastus* had a significant proportion of species that were invasive (both relative to other genera) | The pool of introduced species is a very particular subset of all bamboos, so need to be careful about assessing traits linked to invasiveness only on introduced taxa | Fig. 4 |
| Phylogeny              | There will be a non-random assortment of which species are introduced across the phylogeny | Only culm height showed significant phylogenetic signal, other variables including status were not | | See Fig. S2 |
| Lineage (neotropical woody, etc.) | Taxa from particular biogeographical regions are more likely to become introduced (even if phylogeny and introduction history are taken into account) | Temperate bamboos have had a high rate of species introduced compared with other lineages. Both temperate and paleotropical woody bamboos contain invasive species, but neither had a significant number compared with the other | Bamboos from other parts of the world are likely to have significant potential for utilisation in the future. Region of origin could be an important correlate of risk | Table 2 |
| Number of countries/ regions a species have been introduced to | Species of bamboo that have been introduced to many ranges will have a higher likelihood of becoming invasive | The number of countries a species has been introduced to was strongly (positively) correlated with the likelihood of it being invasive | Risk and impacts caused by non-native bamboos are a function of propagule pressure | See text for details |
| Number of cultivars    | Species with a greater number of cultivars will be more likely to have been introduced than species with fewer cultivars | Introduced species tended to have more cultivars | There has been a possible selection for species that show high levels of phenotypic variation, this can potentially be linked to a greater ability to adapt and so become invasive. On the other hand, more efforts may have simply been made to develop cultivars for common species | See text for details |
|                       | Species with many cultivars will have a higher likelihood of becoming invasive | Greater number of cultivars was an important determinant of invasion | Invasiveness has been selected for during breeding and cultivation practices | |
| Culm form              | Woody lineages will have a higher proportion of introduced species than herbaceous | Woody bamboos are preferred for introduction | As herbaceous species have had much lower rates of introduction, there has been a bias in the natural experiment. | Table 2 |
| Culm dimensions (diameter and height) | Introduced species will on average have greater culm dimensions than non-introduced species | There is an affinity for species to be introduced that have greater culm dimensions | Smaller bamboos will be less likely to have been introduced. | Fig. 5 |
| Rhizome form (running or clumping species) | Introduced bamboo species with running rhizomes are more likely to become invasive, although there is no prior expectation as to how this might affect which species are introduced | Rhizome form was not an indicator of invasive species. However, we did find more running type bamboos have been introduced (although this is correlated with temperate species which have had a bias for introduction) | Control and regulation of bamboos should consider both running and clumping forms | Table 2 |
criteria of Richardson et al. (2011b) [see Supporting Information—Table S3].

To conceptualize and display the flows of introduced and invasive species between and within different biogeographic regions around the world, we used circo visualization from the R package ‘circlize’ (Gu et al. 2014).

Correlates of introduction and invasion

Morphological traits: To determine whether particular traits were related with the introduction status and invasion success of bamboos, we collated trait data from GrassBase. The dataset included 14 trait categories (culms, culm-sheaths, leaves, ligule, etc.). However, only culm dimensions (diameter and height) and underground rhizome system (runner or clumper) were consistently recorded (data on other traits were not available for more than half of the species). These traits were chosen as they were considered relevant to the study and data were available for many of the species.

Different culm properties provide different benefits—thicker-walled culms yield more biomass, greater diameter can produce stronger culms, etc. (Chung and Yu 2002; Scurlock et al. 2000). To determine whether introduced and/or invasive species had taller and/or wider culms than non-introduced species, we used linear models with log-transformed culm dimension (height or diameter) as a response variable and introduction status as the predictor variable. We also included lineage affiliation (paleotropical woody, neotropical woody, temperate woody and herbaceous) as an additional predictor as these have been identified as genetically distinct groups within bamboos that have particular growth forms associated with each (Kelchner et al. 2013). We also tested the differences in culm form of woody versus herbaceous groups in a number of introduced species compared with non-introduced species, and the number of invasive compared with non-invasive species using Fisher’s exact tests. All statistical tests were conducted in R (R Core Team 2015).

Underground rhizome type was also considered a relevant trait for invasion success, as it is often used as a means of separating invasive from non-invasive bamboos (Hamilton 2010; Royal Horticultural Society 2015). There are two forms: running (leptomorph) and clumping (pachymorph). Although sub-forms exist within these categories, for simplicity we only used these two broad categories. Running species are considered to have a greater ability to spread rapidly and are generally considered more invasive than clumping species (Buckingham et al. 2014). To test the difference in number of running and clumping species in the groups of introduced compared with non-introduced, and the number of invasive compared with non-invasive species, we used Fisher’s exact tests.

Taxonomic, geographic and phylogenetic patterns. The exchange of species and the rates of invasion are rarely random, but often have distinct patterns that are influenced by a number of factors, some human-mediated and others related to the evolutionary history of species. Within particular groups this can lead to ‘taxonomic selectivity’. In the case of bamboo, forestry and horticulture have been the main drivers of introductions, and this has led to the preferential selection of taxa. To test whether introductions and invasions have been random, we used Fisher’s exact test to analyse differences between numbers of introduced compared with non-introduced species, and the number of invasive compared with non-invasive species across genera, lineages (i.e. neotropical woody), and introduced countries.

If certain bamboo traits are important to invasion success, and if these traits reflect evolutionary history, then we would expect the phylogeny to indicate ‘taxonomic selectivity’, with only certain lineages becoming invasive. Much work has been done on reviewing this phenomenon to improve the prediction of extinctions. Studies have found that extinctions within taxonomic groups in birds, mammals and plants tend not to be randomly distributed across phylogenies but are concentrated in particular high-risk clades (Fritz and Purvis 2010; McKinney and Lockwood 1999). This is arguably due to phylogenetically conserved life-history traits or ecology (Fritz and Purvis 2010; Purvis 2008; Schwartz and Simberloff 2001; Thomas 2008). There is evidence to suggest this is also true with invasiveness across taxa (Lockwood 1999; Lockwood et al. 2001; Lockwood and McKinney 2001; Novoa et al. 2015; Yessoufou et al. 2016). We explore this for bamboos by testing the phylogenetic signal of status (introduced/invasive) and other correlates of introduction and invasion. To do this we collated genetic data for one chloroplast gene region (maturase K; matK) for all taxa with available data in the online GenBank repository (ncbi.nlm.nih.gov) for phylogeny reconstruction. Where possible, GenBank accessions denoted as ‘voucher’ specimens were used. Our final dataset comprised 124 taxa (including two non-bamboo grass species Bromus interruptus & Trisetum spicatum as outgroup taxa). DNA sequence data were combined and aligned in the BioEdit version 7.0.5.3 (Hall 2006) and were edited manually. Flanking regions were trimmed to avoid excessive missing data. Our final DNA alignment consisted of 860 characters and contained three gaps ranging between 1 and 6 base pairs. A Bayesian inference phylogeny was reconstructed using Mr Bayes v 3.2 (Ronquist and Huelsenbeck 2003), jModelTest2.13 (Darriba et al.,...
and the Akaike information criterion (Akaike, 1973) determined the best fit model for our data as the GTR + I + G model. The Bayesian model was run for 1.5 million generations sampling every 1000th generation and a consensus tree was built, discarding the first 25% of trees as burn-in. Posterior probabilities (PP) were calculated using a majority rule consensus method to assess tree topology support.

To test whether continuous traits (culm dimensions) are phylogenetically clustered or over-dispersed, we used Blomberg’s K statistic with a null hypothesis of Brownian Motion Model (Blomberg et al. 2003). We also tested for phylogenetic signal of other variables, i.e., introduction and invasion frequency (the number of countries a species has been introduced to or become invasive), and propagule pressure (using the frequency of cultivars as a proxy; see below) using Pagel’s λ (lambda) which uses transformation of the branch lengths assuming Brownian motion (Pagel 1999). Both analyses were done using the R packages ‘phytools’ and function Phylosig.R (Revell, 2012)

Species traits, status and cultivar diversity per species were mapped onto the phylogeny to visualise patterns using the R package ‘adephylo’ (Jombart et al. 2010) [see Supporting Information—Fig. S2]. We used the D statistic (Fritz and Purvis 2010) to test for phylogenetic signal and strength of binary traits. This method tests whether traits are randomly assigned across the phylogeny tips (when D equals 0), and whether they are clustered (D equals 1) under a Brownian threshold model. We carried out two tests: one for introduction status (introduced/not introduced) across the whole phylogeny; in the second, we used a tree trimmed to include only introduced bamboos and tested invasion status (invasive/not invasive). This was done using the R package Caper with function phylo.d (Orme et al. 2012).

Introduction effort and utility: Many species of bamboo have had cultivars developed for improving their utility and value. We suggest that cultivar diversity associated with species could provide a proxy and quantitative means to measure their popularity and utility. Cultivars are cultivated plant varieties that are developed through selective breeding, genetic manipulations such as polyploidy and hybridization. They are often distinctive, uniform and stable and retain key characteristics when propagated (Brickell et al., 2009). Cultivar diversity likely corresponds with propagation frequency and will, therefore, be an important determinant of the probability of introduction, as well as invasion success.

As there is no officially accredited list of bamboo cultivars, we used the list compiled by Ohrnberger (1999) based on the 1995 International Code of Nomenclature for Cultivated Plants (ICNCP). To assess the relationship between introduction status and the number of cultivars developed we used a generalized linear model with a Poisson error structure with number of cultivars as the response variable and status as a predictor variable. As a proxy of introduction effort, we used the number of regions into which a species has been introduced. We tested for this using a generalized linear model with a Poisson error structure with the number of regions a species has been introduced to as a predictor variable and the number of regions a species is invasive in as a response variable.

Results

Inventory of species and distribution

Our final list of bamboo species contained 1662 species representing 121 genera, with native species distributed across 122 countries and distinct islands/regions.

Dissemination and status

Two hundred and thirty-two species (14% of the species in the subfamily) are known to have been introduced outside of their native ranges, with about 5.2% (12 species) of these introduced species becoming invasive (Fig. 1). However, some regions of the world were markedly over- or under-represented in terms of the number of introduced species (Fig. 2). There were also cases of unknown or disputed native ranges possibly due to a combination of a high degree of introductions and/or lack of reliable records (11 species across 60 countries and regions). Asiatic species have been most widely exported, with Oceania, North America and Europe being the predominant recipients (Fig. 1). All the species reported as invasive are Asiatic. Although South America has a rich native bamboo flora, most movements of these species have been within the continent. We found no evidence of invasive alien bamboos originating from this region. The range of invasive species is shown in Fig. 3.

Correlates of introduction and invasion

Morphological traits: We found all three trait characteristics tested (rhizome form, culm height and culm diameter) to be significantly associated with different stages along the introduction-naturalization-invasion continuum.

For rhizome forms, a significantly higher proportion of introduced species had runner rhizomes (leptomorphs) than clumping rhizomes (pachymorphs), but there was no significant difference in rhizome form for invasive species (Table 2).

For culm dimensions, there were significant differences between lineages ($F(3,791) = 89.65; P < 0.001$);
we, therefore, included lineage affiliation in the analyses below. We found that the average culm diameter for introduced bamboos was significantly greater than for non-introduced bamboos ($R^2 = 0.2687, F(5, 786) = 57.75, P < 0.001$). There was no significant difference in diameter between introduced and invasive species of bamboos in general. Within the paleotropical woody group, species were found to have wider culms relative to other groups. Culm height was greater in the group of introduced species ($P < 0.001$) and for the invasive group ($P = 0.015$), compared with the non-introduced group of species. All woody groups were found to be significantly taller than the herbaceous group ($R^2 = 0.5039, F(5, 937) = 190.4, P < 0.001$).

Taxonomic, geographic and phylogenetic patterns: At the lineage level, temperate and paleotropical woody bamboo species have been introduced to significantly more countries/regions compared with other groups (Table 2). Herbaceous species had a low proportion of introduced species. Both temperate and paleotropical woody bamboos contained invasive species, yet only temperate woody taxa had a significant proportion of introduced species that have become invasive. At the genus level, there was a significantly (Fisher’s exact test; $P < 0.05$) high proportion of introduced species that belonged to the genera *Arundinaria* (100 %), *Thyrocastachys* (100 %), *Semiarundinaria* (71.4 %), *Phyllostachys* (63 %), *Shibateae* (57.1 %), *Himalayacalamus* (50 %) and *Bambusa* (25.6 %) (Fig. 4). *Phyllostachys* ($n = 5$) and *Pseudosasa* ($n = 2$) were significant in the number of invasive species, with the remaining invasive species belonging to *Bambusa* ($n = 3$), *Dendrocalamus* ($n = 1$) and *Pleioblastus* ($n = 1$).

With respect to phylogenetic signal, our retrieved phylogeny showed low resolution due to the conservative nature of the *matK* gene. Nevertheless, major and well-supported clades corresponded well with higher-level bamboo taxonomy (e.g. subtribe) and known biogeography. Of the continuous traits tested, culm height ($K = 0.097, P = 0.014$) had a significant phylogenetic signal using Blomberg’s K statistic; but using Pagel’s $\lambda$ both culm height ($\lambda = 0.251, P < 0.001$) and culm diameter ($\lambda = 0.418, P < 0.001$) were significant. For our binary status traits, we found a random pattern for introduction status ($D = 0.96, P_{rand} = 0.273, P_{BM} = 0.00$) and for invasion status ($D = 1.24, P_{rand} = 0.77, P_{BM} = 0.00$).

Introduction effort and utility: We found strong evidence that cultivar diversity was associated with introduction status. Species with more cultivars were significantly more likely to have been introduced ($b = 3.56 \pm 0.277, P < 0.001$) and have become invasive ($b = 5.89 \pm 0.313, P < 0.001$). Compared with introduced species, invasive species had a greater number of cultivars ($b = 2.32 \pm 0.181, P < 0.001$), and non-introduced species had significantly fewer cultivars ($b = -3.56 \pm 0.298, P < 0.001$). Further, we found that the number of regions a species was invasive to be positively and significantly correlated with the number of regions to which a species has been introduced (Poisson GLM: $b = 1.02 \pm 0.090, P < 0.001$).
Discussion

Bamboo species have had a long history of introductions and are now commonly found around the world (Figs 1A and 2) but only a few (12) species are invasive (Fig. 3). As predicted, the movement of bamboos is, however, far from complete and the selection and distribution of species has not been random. We identified three main factors that appear to have influenced patterns of introduction and invasion: introduction effort, propagation of species and selection of traits. Each of these is discussed below and we conclude with an assessment of the current extent of bamboo invasion and expansion of some taxa in their native ranges.

Figure 2. Number of bamboo species found in 52 countries and islands with the highest bamboo richness. Regions with less than 15 species were excluded (135 regions) from the figure. Shading indicates the status of bamboo species in that region (native/introduced/invasive). Significance was calculated using Fisher’s exact tests between numbers of introduced compared with non-introduced species and numbers of invasive compared with non-introduced species across countries.
Table 2: The effect of biogeographic lineage, culm form and underground rhizome form on whether taxa tended to be introduced or become invasive. Each group was tested independently to determine whether species in a particular group or with particular features have been introduced and become invasive significantly more often than other bamboo species. This was done using a Fisher’s exact test comparing the number of introduced versus non-introduced species, and invasive versus non-invasive.

| All       | Status       | Introduced | Invasive       |
|-----------|--------------|------------|----------------|
|           | N | N | %   | P  | N | N | %   | P  |
| Biogeographic lineage |              |            |                |                |            |            |                |                |
| Temperate woody | 500 | 101 | 20.2 (16.8–24.0) | 0.0067 | 8 | 2 | (0.9–3.8) | 0.022 |
| Paleotropical woody | 450 | 72 | 16.0 (12.7–19.7) | 0.0088 | 4 | 1 | (0.3–2.7) | 1.00 |
| Neotropical woody | 300 | 32 | 11.0 (7.9–15.0) | 0.813 | 0 | – | – | 0.0460 |
| Herbaceous | 114 | 8 | 7.0 (3.1–13.4) | 0.0005 | 0 | – | – | 0.615 |
| Culm form |              |            |                |                |            |            |                |                |
| Woody | 1293 | 202 | 16.4 (14.4–18.5) | 0.0067 | 12 | 1.1 (0.6–1.9) | 0.615 |
| Herbaceous | 114 | 7 | 7.0 (3.1–13.4) | 0.0067 | 0 | – | – | 0.615 |
| Underground rhizome form |              |            |                |                |            |            |                |                |
| Running | 331 | 71 | 21.4 (16.9–26.4) | 0.0018 | 8 | 1.6 (0.4–4.1) | 0.24 |
| Clumping | 860 | 116 | 13.5 (11.2–16.0) | 0.0018 | 4 | 0.7 (0.2–1.6) | 0.24 |

Introduction effort

Introduction effort, or propagule pressure, has consistently been linked with successful invasions as greater numbers of propagules and more frequent introductions mean higher probabilities of invasion (Colautti et al. 2006; Lockwood et al. 2005; Von Holle and Simberloff 2005). The positive correlation of propagule pressure and invasion success has been observed in many taxa including birds (Duncan 1997; Veltman et al. 1996), mammals (Crowell 1973; Forsyth et al. 2004) and aquatic species (Colautti 2005; Duggan et al. 2006). This is notable with intentional introductions, such as the case with many ornamental (Dehnen-Schmutz and Touza 2008) and cultivated agricultural (Pysek et al. 2009a, b) plants. We found a clear link between introduction effort and invasiveness in bamboos. Although it was not possible to measure propagule pressure directly, species that had been more widely disseminated were much more likely to have become invasive.

Historical activities in the native range have also played an important role in influencing introduction effort. For example, the local propagation and use of native species may increase the chance of a species becoming established after introductions (Forcella and Wood 1984, Lockwood et al. 2005, Pysek et al. 2009a, b). Woody bamboos, in particular, have long been used as a harvested forest resource in regions where they are native (Lobovikov et al. 2007). We found that woody bamboos from Asia have been introduced much more often than species from other regions, and all invasive bamboos are native to Asia. This may be explained by an extensive history of active cultivation of woody bamboos around the continent which has promoted the movement of a subset of species (Scurlock et al. 2000; Yuming et al. 2004; Yuming and Chaomao 2010). Notably in China, bamboo
has been widely used for millennia (Li and Kobayashi 2004). Bamboos have shaped the history of this region and they are now an ingrained cultural and economic aspect of many Asian societies. This would have profoundly influenced the way bamboos from this region have been distributed to other parts of the world.

By comparison, the exploitation of bamboo resources in South and Central America, regions also rich in native bamboo species (roughly 32% species; 530 species), has been historically limited to local and small-scale usage as a forest resource, and, to a lesser extent, as a cultivated crop (Londoño 1998). The number of exported...
species (or propagation with regards to cultivars) has been low compared with Asiatic species, with the movements being mostly within the continent (Fig. 1A). If these patterns continue, it is likely that future introductions will continue to come from Asia, although there might be significant untapped potential in bamboos from the Americas (Li and Kobayashi 2004).

We found strong selection bias, and, therefore, taxonomic selectivity, for the mostly Asian genera *Bambusa* and *Phyllostachys*. Both genera harbour a high number of invasive species (relative to other bamboo genera) and have been extensively introduced around the world (Fig. 4). *Phyllostachys* is a highly utilized temperate woody genus (59 species) from Asia, mostly central China. More than 50 % of species in this genus have been moved outside of their native ranges (the highest proportion of any bamboo genus), and six species are listed as invasive. *Bambusa*, a paleotropical woody genus, is also highly utilized and is the second largest bamboo genus (149 species). At least 25 % of species in the genus have been introduced to areas outside their natural ranges, and three species have become invasive. Of these, *B. vulgaris* is the most widely distributed species (123 countries); indeed it deserves the title of 'the most common bamboo in the world' (Ferrelly 1984). The introduction of *B. vulgaris* to many tropical islands in the Pacific and the Caribbean by early shipping trade routes has left a legacy of naturalized populations (Pacific Island Ecosystems at Risk 2011; O'Connor et al. 2000; Rashford 1995).

### Propagation of species

The fact that some bamboo taxa have been introduced much more widely than others is similar to the patterns observed in other plant groups where there has been a clear bias for species with traits associated with human-usage (Moodley et al. 2013; Novoa et al. 2015). Species suited for ornamental and agricultural purposes have a higher degree of introduction effort. The horticulture trade in particular has been consistently identified as a major introduction pathway for invasive plants (Dehnen-Schmutz and Touza 2008). Aspects of the industry have been found to be good indicators of risk. For example, increased market availability of species and lower prices of seeds were found to increase the invasion success of species traded in the British horticultural market (Dehnen-Schmutz and Touza 2008).

Drew et al. (2010) argued that the horticultural industry is driven by a demand for novel and exotic species, but that there is also a demand for more robust (i.e. with higher stress tolerance) plants for easy maintenance. As the development of cultivars has helped the industry meet some of these demands, cultivar diversity likely reflects the utility (and market demand) of species for horticulture or cultivation. In the case of bamboos, where

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**Figure 5.** Culm diameter (mm) and culm height (cm) of bamboo species (error bars indicate 95 % confidence intervals) across lineages, and grouped by status. Shading indicates the number of species at each point, with lighter yellow representing less species and darker red shades representing many species. Numbers at the top of each plot indicate the number of species (in which data were available) for the corresponding status group.
there has been a consistent and long history of propagation and distribution of plants for horticulture (ornamental plants, landscape improvement, erosion control, etc.) and agroforestry (construction material, crafts, paper pulp, fuel), we expected that the movement of bamboos would be partially influenced by popularity of certain species (Lobovikov et al. 2007; Rashford 1995). We found that greater cultivar diversity of species was strongly correlated with the frequency of introductions, and even more so with invasions. We also noted that our list of cultivars were all species of Asian origin, providing further support for the view that historical cultivation of species in this region has been a key determinant for their global export.

Although we did not measure the market preferences directly, cultivar diversity also likely reflects aspects of demand and can help reveal insights into the market preference for certain species. Species that are more widely traded and utilised will have had more efforts made to develop cultivars and vice versa, supporting the notion that market preferences are a key driver of introduction effort with bamboos, as is the case with other economically valuable plant taxa. As far as we know, the link between cultivar development and utility of a species with respect to increasing the probability of introduction and invasions has not been explored for other plant groups.

Selection of traits

Horticulture directly facilitates the movement of species, but it also provokes the selection of certain traits that can increase establishment and the invasion potential of propagules once introduced (Anderson et al. 2006; Dehnen-Schmutz and Touza 2008; Kowarik 2003; Mack 2000; Martínez-Ghersa and Ghersa 2006). Linking traits to the success of invasive species has been a strong focus of invasion science and many studies have revealed generalities across many taxonomic groups. Production of large numbers of seeds, fast growth rates and large plant size are some examples of traits positively associated with invasiveness (Cadotte and Lovett-Doust 2001; Pysek and Richardson 2007; Van Kleunen et al. 2010).

We found that traits likely related to economic benefits are important in bamboos. Culm attributes were associated with the status of species—whether they had been introduced and were invasive; in particular there was an over-representation of introduced and invasive species with greater dimensions. This may be because the culm is a valuable aspect of the plant, and there has been an incentive to select for bigger bamboos to increase production of woody biomass and in general produce larger poles (Kleinhenz and Midmore 2001). However, culm traits did not explain why Asiatic species have been more introduced (and become invasive) than bamboos from other parts of the world. We found that neotropical woody bamboos (of South and Central American origin) were similar to woody bamboo groups in terms of size. Other traits that are important for bamboo as a construction material, which we were unable to test, include culm wall thickness, culm flexibility and internode length.

We expected that the type of clonal growth in bamboos would be an important determinant of invasiveness because bamboos rarely proliferate sexually. It is often suggested in the literature that species that produce long rhizomes (i.e. runner species) are more aggressive than species that produce short rhizomes (RHS 2015). However, we found that both running and clumping species have become invasive. Therefore, the pattern of clonal growth did not clearly separate invasive from non-invasive species and other factors such as human usage, propagule pressure and residency time, need to be considered in any discussion of invasiveness in bamboos.

Species belonging to the genus Phyllostachys are most often referenced regarding their ability to spread widely due to fast growth rates and extensive sympodial systems of rhizomes, features which can lead to the formation of monocultures (Isagi and Torii 1997; Suzuki and Nakatsubo 2001). The formation of dense stands can result in a decline in biodiversity through the exclusion of native species (Huai et al. 2010; Okutomi et al. 1996; ShangBin et al. 2013; Yang et al. 2008). Phyllostachys species have also been shown to invade on a more localised scale, such as in horticultural garden settings (Royal Horticultural Society 2015). In the United States, Phyllostachys species (typical examples being P. aurea, P. aureosulcata, and P. edulis) are distributed and planted as popular ornamental and garden screening plants. However, perhaps due to lack of management and knowledge in maintaining the underground rhizome system, there are reports of populations that have escaped and become naturalized to the extent that they have been shown to occupy 71,588 acres of forests in the US (Miller et al. 2008). Phyllostachys can also cause a nuisance in urban areas (Connecticut Invasive Plants Council 2011; Joint Standing Committee Hearings 2013). Reported issues in urban areas include structural damage to property from emerging shoots, colonization of gardens and neighbouring land, difficulty and high costs of removing populations due to robust root systems (Joint Standing Committee Hearings 2013). There have been moves to regulate, at the county and state level, the planting and sale of running species (Joint Standing Committee Hearings 2013). With increasing examples of issues surrounding the planting of Phyllostachys species,
it is likely that other temperate bamboos with similar growth habits and uses will cause similar problems.

**Expansion in the native range**

Aspects of the native range have been found to influence the invasiveness of species. For example, species originating from regions with high phylogenetic diversity are more likely to be successful invaders, perhaps because they have more competitive traits (Fridley and Sax 2014). All invasive bamboos originated from Asia, but there was no evidence of a significant phylogenetic signal indicating a particular lineage or clade of bamboo that may be a source for invasive species. This suggests that other factors such as human-mediated usage are more important in explaining invasiveness. However, the corollary of the above observation is that areas with low species richness are likely to be highly invisible (Fridley and Sax 2014). In terms of recipient regions, we did find that the majority (8 of 12) of the areas where bamboo invasions were recorded were islands (areas of low general native plant diversity and specifically low native bamboo diversity).

Another important factor associated with phylogenetic diversity and invasiveness was the size of the range of species. Species with larger native ranges tend to have greater invasion success, because they possess traits that have facilitated establishment over a wide range of environmental conditions (e.g. Moodley et al. 2013; Novoa et al. 2014; Pysek et al. 2009a, b). Range size has also been manipulated by human-usage, as many species have been moved and cultivated beyond the extent of their native provenance. We were unable to account for native range size as delimiting ranges for bamboos was difficult, especially in Asia where there has been extensive exchange and cultivation of species over millennia (Lobovikov 2005; Yuming et al. 2004). We found many records for the movement of Asiatic species to other continents, but much less information on within-continent movements. For example, Moso bamboo (*Phyllostachys edulis* syn. *P. pubescens*), one species of about 583 native to China, has become widespread (both through natural spread and cultivation) and is estimated to make up 80% of bamboo cover (5 million ha) across the country (Bowyer et al. 2014). Its distribution is still increasing, in part due to extensive plantings but also due to disturbances in mixed forests (Gagnon and Platt 2008) that have facilitated its increased abundance and dominance in some vegetation types (Huai et al. 2010; Rother et al. 2016; Song et al. 2015; ShangBin et al.; 2013; Tokuoka et al. 2015, Yang et al. 2008; Xu et al. 2015).

In general, expansion and weedy behaviour of plants in their native range has been shown to be a good indicator of invasive potential (e.g. Richardson and Bond 1991). As past introductions of bamboos have favoured a certain set of species from particular regions, there is significant potential for bamboos in other parts of the world such as South America to be utilised in the future. Such species that have been identified as being highly competitive and weedy in native regions have the potential to become invasive in new areas given the opportunity, and should be carefully evaluated for future introductions. Some examples of bamboos that are found to be weedy and have had impacts in their native ranges are *Pleioblastus arentostriatus* (syn. *P. chino*; Kobayashi et al. 1999; Tokuoka et al. 2015), *Fargesia nitida* (Wang et al. 2012) and *Sasa chartacea* (Tomimatsu et al. 2011) in East Asia. *Ochlandra travancorica* (Dutta and Reddy 2016) and *Melocanna baccifera* (Majumdar et al. 2015) from India, and *Guadua tagoara* (Rother et al. 2016) and *Guadua paraguayana* (Galvão et al. 2012) from South America have not been widely moved outside of their native ranges but, given the observed weedy tendencies of these species in their native ranges, they could pose risks if future introductions were to occur.

Without accurate records on the original ranges of many taxa, it is difficult to comment on the rate of spread and the extent of invasions. We suspect that invasions of some species may have gone unnoticed. This is due to scant information on the native provenance in some regions, and problems with identifying some bamboo species. This is the case where some species are widely dispersed at the continental level and are assumed to be native while they may well be introduced in parts of their current range.

**Extent of invasions**

Overall, we found few invasive species of bamboos (0.7% of taxa) despite the diversity, high rate of dissemination and utilization of various species globally; we had expected this number to be higher. The low number of invasive bamboos is in marked contrast with other taxa within the grass family, which have been noted for containing a high concentration of invasive species (studies estimate between 6 and 10%; Pysek 1998; Visser et al. 2016). Bamboos seem to be an exception in the group. Some of the most extensive invaders in the grass family are large-statured woody grasses, notably *Arundo donax* and *Phragmites australis* (D’Antonio et al. 2010; Lambert et al. 2010). These invasive woody grasses mostly rely on asexual means for spreading via the rhizome systems like many bamboos (Nadgauda et al. 1990). There is scope to investigate such mechanisms in explaining the ability of some large-statured woody grass species to be widespread invaders and why this appears not to be the general case with bamboos.
When compared with other plant taxa outside of the grass family, bamboos have a similarly low occurrence of invasive species; in the group of trees and shrubs it was found that between 0.5% and 0.7% of the global pool of species had become invasive (Richardson and Rejmánek 2011), and for the families of Proteaceae (Moodley et al. 2013), Araceae (Moodley et al. 2016) and Cactaceae (Novoa et al. 2015), 2%, 0.5% and 3% are invasive, respectively.

We discounted invasions in 26 regions (including those involving three additional species) as references could not be verified or were inaccessible. We suspect that the listing of some bamboos as invasive may be unwarranted (or inflated). This is the case with Dendrocalamus strictus, for which it was difficult to disentangle the rate of spread versus impacts, as there was not an explicit distinction in many references [see Supporting Information—Table S3]. In many cases, a long history of planting of bamboos gave the appearance of a prolific, spreading population, whereas the expansion of the population has in fact been minimal or non-existent (O'Connor et al. 2000). For this reason, it is important that standardized and measurable criteria be adopted for defining what ‘invasive’ means for bamboos.

**Conclusions**

Our results suggest that invasiveness in bamboo species is currently more a function of which species have been moved by humans and for what purposes than of inherent differences between species. Certain taxa, for historical and geographical reasons, have rarely been introduced. In particular, native South American bamboos have not yet been widely disseminated. Such taxa might hold promise for future utilisation, and could become invasive. By contrast, past introductions (especially from Asia) have radically rearranged the global distribution of some bamboo species, and new trends in the drivers of introductions are rapidly changing the dimensions in this natural experiment in biogeography. The emergence of large-scale bamboo plantations in new regions of the world represents a fascinating new stage in the bamboo story. There is an urgent need for science-based guidelines to minimize invasion risks.

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**Contributions by the Authors**

S.C., J.R.U.W and D.M.R conceived the idea. S.C compiled the data. V.V. contributed to analysing and visualizing data for final publication. J.J.L.R. assembled the phylogeny. M.V. provided the GrassBase database. S.C. led the writing of the manuscript with inputs from all co-authors.

**Conflicts of Interest**

None declared.

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**Supporting Information**

The following additional information is available in the online version of this article —

**Table S1.** List of Bambusoideae species (1660 species) based on accepted taxa from Kew’s GrassBase (http://www.kew.org/data/grasses-syn.html) with updates to include recent literature on new species and other changes in classification that have been published up until September 2016, described at the generic level in Kellogg (2015). Updates were contributed by bamboo taxonomy specialist Lynn G. Clark (Iowa State University) and grass taxonomy specialist, Maria S. Vorontsova. *232 species have been introduced outside of their native range (numbers following species indicate the number of introduced regions), †12 species are referenced as being invasive,?11 species have unknown or disputed native ranges. Note that the list does not include contemporary hybrids.

**Table S2.** List of named bamboo species that did not match our accepted species list and the changes made to include or exclude from the review database. Synonyms and spelling mistakes were updated accordingly, and unknown names were excluded.

**Table S3.** List of references for bamboo invasions and the locality (‘country/region’) of the reported invasion. The ‘database/report’ indicates where references were found.
All references were vetted for validity on invasion claim (see Richardson et al. 2011), ‘reference status’ indicates which reports were included or exclude in the review.

**Figure S1.** Species richness maps indicating the global geographic distribution of non-native bamboos by database source: (A) Global Biodiversity Information Facility-GBIF, (B) an independent search of literature, (C) Kew’s GrassBase, (D) Global Compendium of Weeds (GCW) (E) IUCN/SSC Invasive Species Specialist Group (ISSG), and (F) Invasive Species Compendium – CABI.

**Figure S2.** Phylogenetic tree of 122 bamboo taxa built using collated genetic data for one chloroplast gene region, matK (matK). All sequences were retrieved from the online GenBank repository (ncbi.nlm.nih.gov). Six variables are shown in columns alongside tree branches showing cultivars no. (number of cultivars), subspecies no. (number of subspecies), forms no. (number of forms), varieties no. (number of genetic varieties), culm diameter (max culm diameter) and culm height (max culm height) of corresponding species. Data in each column is scaled, where large black-filled circles indicate a higher quantity and white-filled circle indicate a smaller quantity of the particular variable associated with the given species relative to other taxa. Numbers after species names indicate the number of regions of introduction and red circles indicate invasive species.

**Literature Cited**

Anderson NO, Golatowitsch SM, Gomez N. 2006. Selection strategies to reduce the invasive potential in introduced plants. *Euphytica* **148**:203–216.

Akaike H. 1973. Akaike Information theory as an extension of the maximum likelihood principle. In: Petrov BN, Csaki F, eds. *Second international symposium on information theory*. Akademiai Kiado, Budapest, 267–281.

Bamboo Phylogeny Group. 2012. An updated tribal and subtribal classification for the bambusoideae (poaceae). In: Gielis J, Kiado, Budapest, 267–281.

Bansal AK, Zoolagud SS. 2002. Bamboo composites: Material of the structural industry. *Journal of Bamboo and Rattan* **10–12 April, Antwerp, Belgium, 3–27.

Bardsey D, Edward-Jones G. 2006. Stakeholders’ perceptions of the impacts of invasive exotic plant species in the Mediterranean region. *GeoJournal* **65**:199–210.

Blackburn TM, Pyšek P, Bacher S, Carlton JT, Duncan RP, Jarosík V, Wilson JRU, Richardson DM. 2011. A proposed unified framework for biological invasions. *Trends in Ecology & Evolution* **26**:333–339.

Blomberg SP, Garland T, Ives AR. 2003. Testing for phylogenetic signal in comparative data: behavioral traits are more labile. *Evolution* **57**:717–745.

Blundell AG, Scatena FN, Wentsel R, Sommers W. 2003. Ecorisk Assessment Using Indicators of Sustainability: Invasive Species in the Caribbean National Forest of Puerto Rico. *Journal of Forestry* **101**:14–19.

Bowyer J, Fernholz K, Frank M, Howe J, Bratkovich S, Pepke E. 2014. Bamboo products and their environmental impacts: revisited. Minneapolis, USA: Dovetail Partners, Inc. http://www.dovetailinc.org/report_pdfs/2014/dovetailbamboo0314.pdf.

Brickell CD, Alexander C, David JC, Hetterscheid WL, Leslie AC, Malecot V, Jin X, Cubey JI. 2009. International code of nomenclature for cultivated plants, 8th ed. *Scripta Horticulturae* **10**:ix–x, 1–184.

Brickell CD, Alexander C, David JC, Hetterscheid WL, Leslie AC, Malecot V, Jin X, Cubey JI. 2009. International code of nomenclature for cultivated plants, Vol. 10, 8th ed. *International Society for Horticultural Science*, 1–184.

Buckingham K, Jepson P, Wu L, Ramanuja Rao IV, Jiang S, Liese W, Lou Y, Fu M. 2011. The Potential of Bamboo is Constrained by Outmoded Policy Frames. *AMBIO* **40**:544–548.

Buckingham KC, Wu L, Lou Y. 2014. Can’t See the (Bamboo) Forest for the Trees: Examining Bamboo’s Fit Within International Forestry Institutions. *AMBIO* **43**:770–778.

Bystrickova N, Kapos V, Lysenko I. 2004. Bamboo biodiversity: Africa, Madagascar and the Americas. *UNEP/Earthprint* no. 19.

Cadotte MW, Lovett-Doust J. 2001. Ecological and taxonomic differences between native and introduced plants of southwestern Ontario. *Ecoscience* **8**:230–238.

Chung KF, Yu WK. 2002. Mechanical properties of structural bamboo for bamboo scaffolding. *Engineering Structures* **24**:429–442.

Clayton WD, Vorontsova MS, Harman KT, Williamson H. 2015. World Grass Species: Synonymy. http://www.kew.org/data/grasses-syn.html.

Colautti RI. 2005. Are characteristics of introduced salmonid fishes biased by propagule pressure?. *Canadian Journal of Fisheries and Aquatic Sciences* **62**:950–959.

Colautti RI, Grigorovich IA, MacIsaac HJ. 2006. Propagule pressure: a null model for biological invasions. *Biological Invasions* **8**:1023–1037.

Connecticut Invasive Plants Council. 2011. *Ninth annual report*. University of Connecticut. Connecticut Invasive Plant Working Group.

Cook GD, Dias L. 2006. It was no accident: deliberate plant introductions for invasive species control: a detailed examination of the horticultural industry. *The American Naturalist* **167**:s1–s4.

D’Antonio CM, Hughes RF, Tunison JT. 2010. Long-term impacts of invasive grasses and subsequent fire in seasonally dry Hawaiian woodlands. *Ecological Applications* **21**:1617–1628.

Damiña D, Taboada GL, Doalla R, Posada D. 2012. *jModelTest* 2: more models, new heuristics and parallel computing. *Nature Methods* **9**:772.

Dehnen-Schmutz K, Touza J. 2008. Plant invasions and ornamental horticulture: Pathway, propagule pressure and the legal framework. In: Teixeira da Silva J A, ed. *Floriculture, ornamental and plant biotechnology: advances and topical issues*. Isleworth: Global Science Books, 15–21.

Drew J, Anderson N, Andow D. 2010. Conundrums of a complex vector for invasive species control: a detailed examination of the horticultural industry. *Biological Invasions* **12**:2837–2851.

Duncan RP. 1997. The role of competition and introduction effort in the success of passeriform birds introduced to New Zealand. *The American Naturalist* **149**:903–915.
Fridley JD, Sax DF. 2014. The imbalance of nature: revisiting a Darwinian framework for invasion biology. Global Ecology and Biogeography 23:1157–1166.

Fritz SA, Purvis A. 2010. Selectivity in Mammalian Extinction Risk and Threat Types: a New Measure of Phylogenetic Signal Strength in Binary Traits. Conservation Biology 24:1042–1051.

Galvão F, Augustin CR, Curcio GR, Cosmo N, Koectra Z, Domanowski BP, Sawczuk AT. 2012. Guadua paraguayana’s impacts on a remaining of subtropical ombrophilous alluvial forest—a biogeochemistry approach. Floresta 42:355–368.

Gagnon PR, Platt WJ. 2008. Multiple disturbances accelerate clonal growth in a potentially monodominant bamboo. Ecology 89:612–618.

Gratopp E, Rejmanek M, Rost TL. 2002. Toward a causal explanation of plant invasiveness: seedling growth and life-history strategies of 29 pine (Pinus) Species. The American Naturalist 159:396–419.

Gu Z, Gu L, Eils R, Schlesner M, Brors B. 2014. circlize implements and enhances circular visualization in R. Bioinformatics 30 (19): 2811–2812.

Holl T. 2006. Bioedit Version 7.0. 5.3. Department of Microbiology, North Carolina State University [Online]. Available: http://www.ncbi.nlm.nih.gov/ Bioedit/bioedit.html (17 January 2017).

Hoogendoorn JC, Benton A. 2014. Bamboo and rattan production and the implications of globalization. In: Nikolaos W, Innes J, eds. Forests and globalization: challenges and opportunities for sustainable development. London: Routledge, 166–184.

Huay L, PeiXue L, HuTang D, Dan L, XianSheng Y. 2010. Impacts of Phyllostachys pubescens expansion on plant species diversity in Jigong Mountain and discussion of control measures. Journal of Xinyang Normal University Natural Science Edition 23:553–557.

Hunter I. 2003. Bamboo resources, uses and trade: the future?. Journal of Bamboo and Rattan 2:319–326.

INBAR. 2003. Greening red earth beijing: INBAR.

Isagi Y, Torii A. 1997. Range expansion and its mechanisms in a naturalized bamboo species, Phyllostachys pubescens, in Japan. Journal of Sustainable Forestry 6:127–141.

Janzen DH. 1976. Why bamboo wait so long to flower. Annual Review of Ecology and Systematics 7:347–391.

Joint Standing Committee Hearings. 2013. Connecticut General Assembly: Environment Committee. Part 6:1666 – 2009.

Jombart T, Balloux F, Dray S. 2010. Adeephy: new tools for investigating the phylogetic signal in biological traits. Bioinformatics 26(15):1907–1909.

Kelchner SA, Bamboo Phylogeny Group. 2013. Higher level phylogenetic relationships within the bamboos (Poaceae: Bambusoideae) based on five plastid markers. Molecular Phylogenetics and Evolution 67:404–41.

Kellogg EA. 2015. Poaceae. In: Kubitzki K, ed. The families and genera of vascular plants. Flowering plants. Monocots: Poaceae, Vol. 13. Cham: Springer.

Kleinhenz V, Midmore DJ. 2001. Aspects of bamboo agronomy. Advances in Agronomy 74:99–153.

Kobayashi T, Fukushima K, Hisamoto Y, Iinoue A. 2015. The species biology of bamboos in Japan: from gene to landscape. Plant Species Biology 30:42–44.

Kobayashi T, Saito A, Hori Y. 1999. Species diversity of the understory dominated by dwarf-bamboo Pleioblastus chinus Makino in a secondary forest with different numbers of years after the last mowing. Journal of the Japanese Society of Revelocity Technology 24:201–207.

Kowarik I. 2003. Human agency in biological invasions: secondary releases foster naturalisation and population expansion of alien plant species. Biological Invasions 5:293–312.

Kudo G, Amagai Y, Hoshino B, Kaneko M. 2011. Invasion of dwarf bamboo into alpine snow-meadows in northern Japan: pattern of expansion and impact on species diversity. Ecology and Evolution 1:85–96.

Kueffer C, Pysek P, Richardson DM. 2013. Integrative invasion science: model systems, multi-site studies, focused meta-analysis and invasion syndromes. New Phytologist 200:615–633.

Lambert AM, Dudley TL, Saltinston K. 2010. Ecology and Impacts of the Large-Statured Invasive Grasses Arundo donax and Phragmites australis in North America. Invasive Plant Science and Management 3:489–494.

Li ZH, Kobayashi M. 2004. Plantation future of bamboo in China. Journal of Forestry Research 15:233–242.

Liese W, Köhl M. 2015. Bamboo: the plant and its uses. Tropical forestry. Springer, Basel, Switzerland.

Lima RAF, Rother DC, Muler AE, Lepsch IF, Rodrigues RR. 2012. Bamboo overabundance alters forest structure and dynamics in the Atlantic Forest hotspot. Biological Conservation 147:32–39.

Lobovikov M. 2005. Bamboo forest trends. International Network for Bamboo and Rattan. Available: http://www.forest-trends.org/documents/files/doc_1123.pdf (17 January 2017).

Lobovikov M, Paudel S, Piazza M, Wu HR. 2007. World bamboo resources; a thematic study prepared in the framework of the global forest resources assessment 2005, Rome: Food and Agriculture Organization of The United Nations.

Lockwood JL. 1999. Using taxonomy to predict success among introduced avifauna: relative importance of transport and establishment. Conservation Biology 13:560–567.

Lockwood JL, Cassey P, Blackburn T. 2005. The role of propagule pressure in explaining species invasions. Trends in Ecology & Evolution 20:223–228.

Lockwood JL, McKinney ML (Eds). 2001. Biotic homogenization. New York: Springer US.

Lockwood JL, Simberloff D, McKinney ML, Von Holle B. 2001. How many, and which, plants will invade natural areas? Biological Invasions 3:1–8.

Londoño X. 1998. Evaluation of bamboo resources in Latin America. A summary of the final report of project no. 96-8300-01-4. Beijng: International Network for Bamboo and Rattan.

Mack RN. 2000. Cultivation fosters plant naturalization by reducing environmental stochasticity. Biological Invasions 2:111–122.
Martínez-Ghersa MA, Ghersa CM. 2006. The relationship of propagule pressure to invasion potential in plants. Euphytica 168: 87–96.

McGeoch MA, Spear D, Kleynhans EJ, Marais E. 2012. Uncertainty in invasive alien species listing. Ecological Applications 22: 959–971.

McKinney ML, Lockwood JL. 1999. Biotic homogenization: a few winners replacing many losers in the next mass extinction. Trends in Ecology & Evolution 14:450–453.

Majumdar K, Nath AJ, Gupta AK, Datta BK. 2015. Bamboo invasion: threat to primate conservation in North East India. Current Science 108:1969–1971.

Meyerson LA, Mooney HA. 2007. Invasive alien species in an era of globalization. Frontiers in Ecology and the Environment 5: 199–208.

Miller JH, Chambliss EB, Oswalt CM. 2008. Maps of occupation and estimates of acres covered by normative invasive plants in southern forests using SRS FIA data posted on March 15, 2008. Available: (http://www.invasive.org/fiamaps/) (13 January 2017).

Moodley D, Procheř Š, Wilson JRU. 2016. A global assessment of a large monocot family highlights the need for group-specific analyses of invasiveness. AoB Plants plw009v1-plw009.

Moodley D, Geerts S, Richardson DM, Wilson JRU. 2013. Different traits determine introduction, naturalization and invasion success in woody plants: Proteaceae as a test case. PLoS One 8: e75078.

Nadgauda RS, Parasharami VA, Mascarenhas AF. 1990. Precocious flowering and seeding behaviour in tissue-cultured bamboos. Nature 344:335–336.

Novoa A, Le Roux JJ, Robertson MP, Wilson JRU, Richardson DM. 2015. Introduced and invasive cactus species—a global review. AoB Plants 7:plu078.

O’Connor PJ Covich, AP, Scatena FN, Loope LL. 2000. Non-indigenous bamboo along headwater streams of the Luquillo Mountains, Puerto Rico: leaf fall, aquatic leaf decay and patterns of invasion. Journal of Tropical Ecology 16:499–516.

Ohrnberger D. 1999. The bamboos of the world: annotated nomenclature and literature of the species and the higher and lower taxa. Amsterdam: Elsevier Science.

Okutomi K, Shinoda S, Fukuda H. 1996. Causal analysis of the invasion of broad-leaved forest by bamboo in Japan. Journal of Vegetation Science 7:723–728.

Orme D, Freckleton R, Thomas G, Petzold T, Fritz S, Iacoa N, Pearse W. 2012. Caper: comparative analyses of phylogenetics and evolution in R package version 05. Available: https://cran.r-project.org/package=caper (17 January 2017).

Pacific Island Ecosystems at Risk (PIER). 2011. Bambusa spp. Institute of Pacific Islands Forestry. Available: http://www.hear.org/pier/species/bambusa_spp.htm (13 January 2017).

Pagel M. 1999. Inferring the historical patterns of biological evolution. Nature 401:877–884.

Prance G, Nesbitt M. 2005. The cultural history of plants. New York: Routledge, 24 February 2016.

Purvis A. 2008. Phylogenetic approaches to the study of extinction. Annual Review of Ecology, Evolution, and Systematics 39: 301–319.

Pyšek P. 1998. Is there a taxonomic pattern to plant invasions?. Oikos 82:282–294.

Pyšek P, Richardson DM. 2007. Traits associated with invasiveness in alien plants: where do we stand? In: Nentwig W, ed. Biological invasions. Berlin: Springer-Verlag, 97–125.

Pyšek P, Jarosík V, Pergl J, Randall R, Chytrý M, Kühn I, Tichý L, Danhelka J, Sádlo J. 2009a. The global invasion success of Central European plants is related to distribution characteristics in their native range and species traits. Diversity and Distributions 15:891–903.

Pyšek P, Krivánek M, Jarosík V. 2009b. Planting intensity, residence time, and species traits determine invasion success of alien woody species. Ecology 90:2734–2744.

Rashford JH. 1995. The past and present uses of bamboo in Jamaica. Economic Botany 49:395–405.

R Core Team. 2015. R: a language and environment for statistical computing. Vienna: R Foundation for Statistical Computing, 2012. Available: http://www.R-project.org (17 January 2017).

Revell LJ. 2012. phytools: an R package for phylogenetic comparative biology (and other things). Methods in Ecology and Evolution 3(2):217–223.

Ricciardi A. 2007. Are modern biological invasions an unprecedented form of global change?. Conservation Biology 21: 329–336.

Richardson DM. 2006. Pinus: a model group for unlocking the secrets of alien plant invasions? Persisla 78:375–388.

Richardson DM, Bond WJ. 1991. Determinants of plant distribution: Evidence from pine invasions. American Naturalist 137:639–668.

Richardson DM, Carruthers J, Hui C, Impson FAC, Richardson DM, Pyšek P, Le Roux JJ, Wilson JRU. 2011a. Human-mediated introductions of Australian acacias—a global experiment in biogeography. Diversity and Distributions 17:771–787.

Richardson DM, Kluge RL. 2008. Seed banks of invasive Australian Acacia species in South Africa: Role in invasiveness and options for management. Perspectives in Plant Ecology, Evolution and Systematics 10:161–177.

Richardson DM, Pyšek P. 2012. Naturalization of introduced plants: Ecological drivers of biogeographic patterns. New Phytologist 196:383–396.

Ritchie MD, Pyšek P, Carlton JT. 2011b. A compendium of essential concepts and terminology in invasion ecology. In: Richardson DM, ed. Fifty years of invasion ecology. The legacy of Charles Elton. Oxford: Wiley-Blackwell, 409–420.

Richardson DM, Rejmánek M. 2011c. Trees and shrubs as invasive alien species—a global review. Diversity and Distributions 17: 788–809.

Ronquist F, Huelsenbeck JP. 2003. MrBayes 3: Bayesian phylogenetic inference under mixed models. Bioinformatics 19: 1572–1574.

Rother DC, Rodrigues RR, Pizo MA. 2016. Bamboo thickets alter the demographic structure of Euterpe edulis population: a keystone, threatened palm species of the Atlantic forest. Acta Caeolologica 70:96–102.

Royal Horticultural Society (RHS). 2015. Bamboo. Royal Horticultural Society, United Kingdom. Available: https://www.rhs.org.uk/advice/profile?PID=79 (13 January 2017).

Scurlor JMO, Dayton DC, Hames B. 2000. Bamboo: an overlooked biomass resource? Biomass and Bioenergy 19:229–244.

Schwartz MW, Simberloff D. 2001. Taxon size predicts rates of rarity in vascular plants. Ecology Letters 4:464–469.
ShangBin B, GuoMa Z, YiXiang W, QianQian L, Juan C, YanYan C, Rui S. 2013. Plant species diversity and dynamics in forests invaded by Moso bamboo (*Phyllostachys edulis*) in Tianmu Mountain Nature Reserve. *Biodiversity Science* 21:285–295.

Song Q-N, Yang Q-P, Ouyang M, Long C-I, Chen F-S, Shi J-M. 2015. Changes in the hydrological functions of litter layer following *Phyllostachys edulis* expansion into evergreen broadleaved forest. *Chinese Journal of Ecology* 34:2281–2287.

Song X, Zhou G, Jiang H, Yu S, Fu J, Li W, Wang W, Ma Z, Peng C. 2011. Carbon sequestration by Chinese bamboo forests and their ecological benefits: assessment of potential, problems, and future challenges. *Environmental Reviews* 15:418–428.

Space JC, Flynn T. 2000. Observations on invasive plant species in *American Samoa*. Honolulu, HI: USDA Forest Service, Pacific Southwest Research Station, Institute of Pacific Islands Forestry.

Suzuki S. 2015. Chronological location analyses of giant bamboo (*Phyllostachys pubescens*) groves and their invasive expansion in a satoyama landscape area, western Japan. *Plant Species Biology* 30:63–71.

Suzuki T, Nakatsubo T. 2001. Impact of the bamboo *Phyllostachys bambusoides* on the light environment and plant communities on riverbanks. *Journal of Forest Research* 6:81–86.

Thomas GH. 2008. Phylogenetic distributions of British birds of conservation concern. *Proceedings of the Royal Society B* 275:2077–2083.

Tokouka Y, Ohigashi K, Watanabe K, Yamauchi H, Arai T, Nakagoshi N. 2015. Removal of competitive native plants by tree planting can accelerate the initial afforestation process: an experiment in an old field in Japan invaded by dwarf bamboo and kudzu. *Journal of Forest Research* 26:581–588.

Tomimatsu H, Yamagishi H, Tanaka I, Sato M, Kondo R, Konno Y. 2011. Consequences of forest fragmentation in an understory plant community: extensive range expansion of native dwarf bamboo. *Plant Species Biology* 26:13–12.

Townsend R. 2013. Bamboos at Kew. ISHS Acta Horticulturae 1003:1 I International Symposium on Genetic Resources of Bamboos and Palms and III International Symposium on Ornamental Palms, pp. 51–53.

Van Kleunen M, Weber E, Fischer M. 2010. A meta-analysis of trait differences between invasive and non-invasive plant species. *Ecology Letters* 13:235–245.

Veltman CJ, Nee S, Crawley MJ. 1996. Correlates of introduction success in exotic New Zealand birds. *The American Naturalist* 147:542–557.

Visser V, Wilson JRU, Fish L, Brown C, Cook GD, Richardson DM. 2016. Much more give than take: South Africa as a major donor but infrequent recipient of invasive non-native grasses. *Global Ecology and Biogeography* 25:679–692.

Von Holle B, Simberloff D. 2005. Ecological resistance to biological invasion overwhelmed by propagule pressure. *Ecology* 86:3212–3218.

Vorontsova MS, Clayton D, Simon BK. 2015. Grassroots e-floras in the Pooceae: growing GrassBase and GrassWorld. *PhytoKeys* 48:73–84.

Wang YJ, Shi XP, Peng Y, Zhong ZC, Tao JP. 2012. Effects of fine-scale pattern of dwarf bamboo on understory species diversity in *Abies faxoniana* forest, southwest, China. *Sains Malaysiana* 41:649–657.

Xu Q-F, Jiang P-K, Wu J-S, Zhou G-M, Shen R-F, Fuhrmann J. 2015. Bamboo invasion of native broadleaf forest modified soil microbial communities and diversity. *Biological Invasions* 17:433–444.

Yang SZ, Du QZ, Chen JX, Liu L. 2008. Effect of *Phyllostachys heterocycla* *var*. *pubescens* spreading on bird diversity. *Journal of Zhejiang Forest Science and Technology* 28:43–46.

Yang QP, Yang GY, Song QN, Shi JM, Ouyang M, Qi HY, Fang XM. 2015. Ecological studies on bamboo expansion: process, consequences and mechanism. *Chinese Journal of Plant Ecology* 39:110–124.

Yoshida T, Goka K, Ishihama F, Ishihara M, Kudo SI. 2007. Biological invasion as a natural experiment of the evolutionary processes: introduction of the special feature. *Ecological Research* 22:849–854.

Yessoufou K, Mearns K, Elansary HO, Stoffberg GH. 2016. Assessing the phylogenetic dimension of Australian *Acacia* species introduced outside their native ranges. *Botany Letters* 163:33–39.

YuMing Y, ChaoMao H. 2010. China’s bamboo-culture/resources/cultivation/utilization. Technical Report 33—International Network for Bamboo and Rattan (INBAR).

Yuming Y, Kanglin W, Shengji P, Jiming H. 2004. Bamboo diversity and traditional uses in Yunnan, China. *Mountain Research and Development* 24:157–165.