

Seed traits and phylogeny explain plant's geographic distribution

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25 **Abstract.** Understanding the mechanisms that shape the geographic distribution of plant species is a central theme of biogeography. Although seed mass, seed dispersal mode and phylogeny have long been suspected to affect species distribution, the link between the sources of variation of these attributes and their effects to the distribution of seed plants are poorly documented. This study aims to quantify the joint effects of key seed traits and phylogeny on species' distribution. We collected seed mass and seed
30 dispersal mode from 1,426 species of seed plants representing 501 genera of 122 families and used 4,138,851 specimens to model species distributional range size. Phylogenetic generalized least squares regression and variation partitioning were performed to estimate the effects of seed mass, seed dispersal mode and phylogeny on species distribution. We found that species distributional range size was significantly constrained by phylogeny. Seed mass and its intraspecific variation were also important in
35 limiting species distribution, but their effects were different among species with different dispersal modes. Variation partitioning revealed that seed mass, seed mass variability, seed dispersal mode and phylogeny together explained 46.82% of the variance in species range size. Although seed traits are not typically used to model the geographic distributions of seed plants, our study provides direct evidence showing seed mass, seed dispersal mode and phylogeny are important in explaining species geographic
40 distribution. This finding underscores the necessity to include seed traits and the phylogenetic history of species in climate-based niche models for predicting the response of plant geographic distribution to climate change.

Keywords. dispersal mode, distributional range size, phylogeny, seed mass, seed mass variability

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1 Introduction

Understanding the ecological and evolutionary processes that govern the geographic range of species
50 can provide insights into their potential adaptive response to global climate change (Gaston and Fuller,
2009; Kubota et al., 2018). It is well known that the geographic ranges of species can span 12 orders of
magnitude, and closely related species may vary enormously in their range (Brown et al., 1996). Many
factors contribute to this variation, although dispersal ability and energy requirements associated with
establishment and persistence in varying habitats have been considered to be the two most important
55 ones (Morin and Chuine, 2006; Zhou et al., 2021). Given that seeds are the predominately mobile stage
of sessile plants, and seed mass generally reflects the amount of energy that a seed contains and its
mobility (Coomes and Grubb, 2003), it seems likely that seed mass could play an important role in
governing the geographic ranges of seed plants.

Seed mass can influence the colonization and competition ability of plant species along different
60 environmental gradients (Chen et al., 2018; Bu et al., 2019). Large-seeded species more often occupy
habitats that have high levels of energy (i.e., tropical or low elevation habitats) and tend to be better
competitors in these environments (Moles and Westoby, 2004), where they typically have higher
germination rates (Akaffou et al., 2021), and greater seedling survivorship (Mukherjee et al., 2019).
Small-seeded species, however, usually occupy low energy habitats. They often produce a large amount
65 of seeds, allowing them to arrive in new (possibly harsher) habitats through wind dispersal (Greene and
Quesada, 2005; Morin and Chuine, 2006; Sonkoly et al., 2017). Furthermore, seed mass has been shown
to decrease along increasing environmental extremes, indicative of the superior colonization ability of
small-seeded species in low energy habitats compared to that of large-seeded species (Procheş et al.,
2012; DeMalach et al., 2019). While some studies (e.g., Morin and Chuine, 2006; Procheş et al., 2012)
70 indicate that species with small and light seeds tend to possess large geographic ranges, there is a need

to further quantify the relationship between seed mass and distributional range size across a broader suite of species and at a wider spatial scale.

Seed traits, including seed mass, could also vary considerably within species, which may be driven by plasticity genes or even molecular signals across the genome (Nicotra et al., 2010). Therefore, 75 intraspecific seed mass variation reflecting a species' high genetic diversity can enable adaptive response to varying environmental conditions and changing climate (Cochrane et al., 2015; Yang et al., 2016), so that to occupy more local habitats (Silvertown, 1989; Sides et al., 2014). Although intraspecific seed mass variation could be an important factor influencing the geographic distribution of plants, few studies have evaluated this source of variation in a regional scale.

80 The seed dispersal mode of a particular species, a key trait responsible for dispersal distance, can also greatly influence species geographic range (Oakwood et al., 1993; Chen et al., 2019b). The seed dispersal ability of a plant species is often a trade-off with other life-history characteristics, such as seed mass, morphologies and persistence in the soil, which in turn can affect seed germination, and the survival and growth of seedlings (Nathan, 2001; Chen and Valone, 2017). However, little is known 85 about the effect of dispersal modes on species distribution. It is also because of the tradeoff between dispersal modes and seed mass variation (Moles et al., 2007; Chen et al., 2019a), discerning the relative importance of seed mass and dispersal on the geographic distribution of seed plants is important but elusive.

Because species from a common ancestor typically experience similar selection pressures in 90 similar habitats, e.g., adaptive niche convergence (Losos, 2008; Grossenbacher et al., 2015), the geographic distribution of species is likely correlated in phylogenetic relationships. Furthermore, phylogenetic relatedness could also influence other ecological processes such as niche partitioning in overlapping habitats or variation in life-history traits, seed traits included, which in turn may influence the distribution range size of species (Moles et al., 2005). Therefore, a species' age or the degree of

95 phylogenetic relatedness could invoke biogeographic limits to expansion (Martin and Husband, 2009) or promote the evolutionary divergence of species and the variation in seed traits (Donoghue et al., 2001; Moles et al., 2005). Although a species' geographic range could well be dependent on its evolutionary history (Felsenstein, 1985), few studies have included phylogeny to discern the effect of seed traits on species distribution.

100 In this study, we attempted to quantify the effects of seed mass, intraspecific seed mass variation, dispersal mode and phylogeny on species geographic range size. We hypothesized that species possessing small seeds with high variability in seed mass, coupled with a strong dispersal capacity, would have larger distributional range sizes than species with contrasting seed traits, and furthermore, species distribution range would be phylogenetically conserved. We collected data on seed mass and
105 seed dispersal mode from 1,426 plant species distributed mainly across China. We specifically aimed to answer two questions: (1) What are the joint effects of seed mass, seed dispersal and phylogeny on species geographic range size? and (2) Are there significant phylogenetic signals associated with species geographic range size?

2 Materials and methods

110 2.1 Seed mass data

Our dataset contains seeds of 1,426 species, representing 501 genera and 122 families of seed plants. All species occur in China, of which about 30% are endemic to China. Seeds from two to 136 populations for each of the species (a total of 17,223 populations) were obtained from the Germplasm Bank of Wild Species (GBWS: <http://www.genobank.org/>). In addition, 549 populations for 454 of the 1,426 species
115 (one to six populations per species) were obtained from the Kew Gardens Seed Information Database (<https://www.kew.org/kew-gardens>). Seeds stored in GBOWS were collected from populations within

the natural distribution range of the species, and dried for 1 to 6 months in a drying room where the relative humidity and temperature were maintained at 15% and 15 °C, respectively. After drying, 50 seeds were randomly sampled from each population for five times (sampling with replacement) and weighed the sampled seeds to the nearest 0.1 mg each time, resulting in five weights for the population. The five weights were averaged and converted to the 1000-seed weight of the population. For each species, the 1000-seed weights across all populations were further averaged and this “grand” average was used as the seed mass for the species. Seed mass variability (i.e., intraspecific variation in seed mass), ranging from zero to one, was calculated for each species as the absolute difference between the maximum 1000-seed weight and the minimum 1000-seed weight across all the populations of the species divided by the maximum value, which is a common measure of plant trait variation (Valladares et al., 2000; Rozendaal et al., 2006). This measure is more suitable than the coefficient of variation (CV), which is sensitive to small changes in mean values when the mean is close to zero; and some plants in this study, such as orchids, have very small seed mass.

2.2 Species distributional range size

In this study, we estimated the distributional range size for each of the 1,426 species using ArcGIS10.2 (ESRI, Redlands, CA, USA) from the global distribution of the species. Thus, the range sizes of the species were the global distribution range. Firstly, the specimen distributional information of each species was obtained from the Global Biodiversity Information Facility (GBIF: <https://doi.org/10.15468/dl.umswqd>, accessed on 04 August 2019), the Chinese Virtual Herbarium (<http://www.cvh.ac.cn/>) and the Biodiversity of the Hengduan Mountains and Adjacent Areas of South-Central China database (BHMAASCC: <http://hengduan.huh.harvard.edu/fieldnotes>). Specimens lacking data on GPS locations, having duplication, containing incorrect coordinates, and those taken from gardens and small oceanic islands were filtered out from our analysis. In addition, species that

140 were cultivated, introduced, invasive, or naturalized were also excluded from our dataset. After
excluding these species records, 4,138,851 specimens of the 1,426 seed plant species were obtained.
Secondly, *shapefile* (containing points) of each species was produced from the coordinates of the
specimens. The *shapefile* was transformed into *raster* using the World Sinusoidal Projection at a spatial
resolution of 100 km using ArcGIS10.2. The distributional range size of each species was calculated by
145 multiplying the number of grids the *raster* contained by 10,000 km² (100 x 100 km). In order to assess
the impact of different spatial resolutions used in calculating species distributional range size, *raster*
with the spatial resolution of 50 km was also used to calculate the range size. Because the distributional
range size calculated at this resolution was highly correlated with the distributional range size calculated
at the resolution of 100 km ($r = 0.993$, $P < 0.001$; Fig. A1), we thus only used the distributional range
150 size calculated at the spatial resolution of 100 km in subsequent analyses.

2.3 Dispersal modes

Based on the published literature and floras, dispersal modes were classified to autochory (self-dispersal,
e.g., by explosive seed release from fruits or gravity, $n = 223$ species), zoochory (dispersal by animals
through ingestion or attachment to an animal body, $n = 468$ species), and anemochory (dispersal by
155 wind, $n = 735$ species) according to the morphological features of their seeds or fruits
(Pérez-Harguindeguy et al., 2013). For example, seeds or fruits with wings, hairs or pappus were
considered wind dispersed (anemochory); seeds or fruits with an aril or flesh offering a succulent
reward for consumers were classified as zoochory; and seeds or fruits lacking modifications pertaining
to the other two categories were classed as autochory (unassisted dispersal) (Qi et al., 2014).

160 2.4 Construction of phylogenetic tree and statistical analyses

For all the species used in our analysis, the scientific names were checked and standardized according to

the Plant List (<http://www.theplantlist.org/>). Different varieties and subspecies of a given species were considered to belong to the same species. The phylogenetic tree was extracted from a previously published supertree using the ‘phylo.maker’ function in R package *V.PhyloMaker* (Jin and Qian, 2019),
165 which was based on the APG classification of flowering plants (Zanne et al., 2014). The ‘multi2di’ function in the *ape* package was used to randomly resolve polytomies in the phylogenetic tree. To test the phylogenetic signal in species distribution, ‘phylosig’ function in the R package *phytools* was used to calculate Pagel’s λ , which is ranged between 0 and 1. $\lambda = 0$ means that the evolution of the trait is phylogenetically independent, and $\lambda = 1$ indicates that trait evolution follows the Brownian motion. Any
170 value of λ significantly higher than zero is regarded to have a phylogenetic signal approaching Brownian motion to a different degree (Arène et al., 2017).

Because closely related species tend to have similar traits, interspecific analyses can be compromised by phylogenetic relatedness (Felsenstein, 1985; Lynch, 1991). In our case, species’ range size is not phylogenetically independent. We thus used a phylogenetic generalized least squares (PGLS)
175 regression to determine the effects of seed mass (SM), intraspecific variation in seed mass (ISM) and dispersal mode (DM) on the distributional range size (RS) of species (Swenson, 2014). The SM×DM and ISM×DM interaction terms were also included in the PGLS model, in order to show effects of SM and ISM on distributional range size among dispersal modes. The regression model was $RS = \beta_0 + \beta_1SM + \beta_2ISM + \beta_3DM + \beta_4SM \times DM + \beta_5ISM \times DM$. The PGLS was implemented using ‘gls’ function
180 in *nlme* package, and the possible phylogenetic dependence in species’ range size was incorporated in a form of a phylogenetic variance-covariance matrix in *gls*.

We further used ‘varpart’ function in *vegan* package to partition the variances in range size explained by seed mass, seed mass variability, dispersal mode, and genus (regarded as phylogeny). Because our phylogenetic tree had some polytomies at the species-level, genera were used as a
185 surrogate in the phylogeny. Variation partitioning is a linear model, which does not require the type of

explanatory variables, and hence is suitable to our data structure (Borcard et al., 2018).

In the analyses of this study, the values of species range size and seed mass were \log_e -transformed to reduce data skewness and downplay extreme values; and the \log_e -transformed seed mass and seed mass variability were standardized to make their coefficients (i.e., effect size) comparable. Seed mass and seed mass variability were each standardized by subtracting the smallest value across all 1,426
190 species and divided by the difference between the largest value and the smallest value. All statistical analyses in this study were conducted using R4.0.2 (R Core Team, 2020).

3 Results

3.1 Effects of phylogeny on species distributional range size

195 We detected a strong phylogenetic signal in species distributional range size for the sampled species ($\lambda = 0.627$, $P < 0.001$), with the signal being stronger in gymnosperms ($\lambda = 0.975$, $P < 0.05$) than in angiosperms ($\lambda = 0.423$, $P < 0.001$). The phylogenetically closely related species had more similar range size than that for distantly related species.

3.2 Effects of seed traits on species distributional range size

200 The results of the phylogenetic generalized least squares regression showed that seed mass had a negatively strong association with species distributional range size (effect size = -13.974, $P < 0.001$; Fig. 1, Table A1), while the effect of seed mass variability on species distributional range size was not significant (effect size = 0.459, $P = 0.109$). Dispersal mode was also significantly associated with species' range size. In the PGLS model, autochorous (explosive/gravity dispersal) species was treated as
205 the baseline dispersal mode. Compared to zoochory (dispersal by animal ingestion or attachment to an animal body) and anemochory (dispersal by wind), autochorous species had significantly larger range

size after the effects of seed mass and seed mass variability were accounted in the interaction terms between seed traits and dispersal modes (Fig. 1, Table A1). The interaction terms between seed mass/seed mass variability and dispersal mode (i.e., seed mass×anemochory, seed mass×zoochory and seed mass variability×zoochory) were significantly positive (effect size = 7.527, $P < 0.001$; effect size = 12.637, $P < 0.001$; effect size = 1.824, $P < 0.001$ respectively), indicating the distributional range sizes of anemochorous and zoochorous species were strongly subject to seed mass and its intraspecific variation (Fig. 1, Table A1).

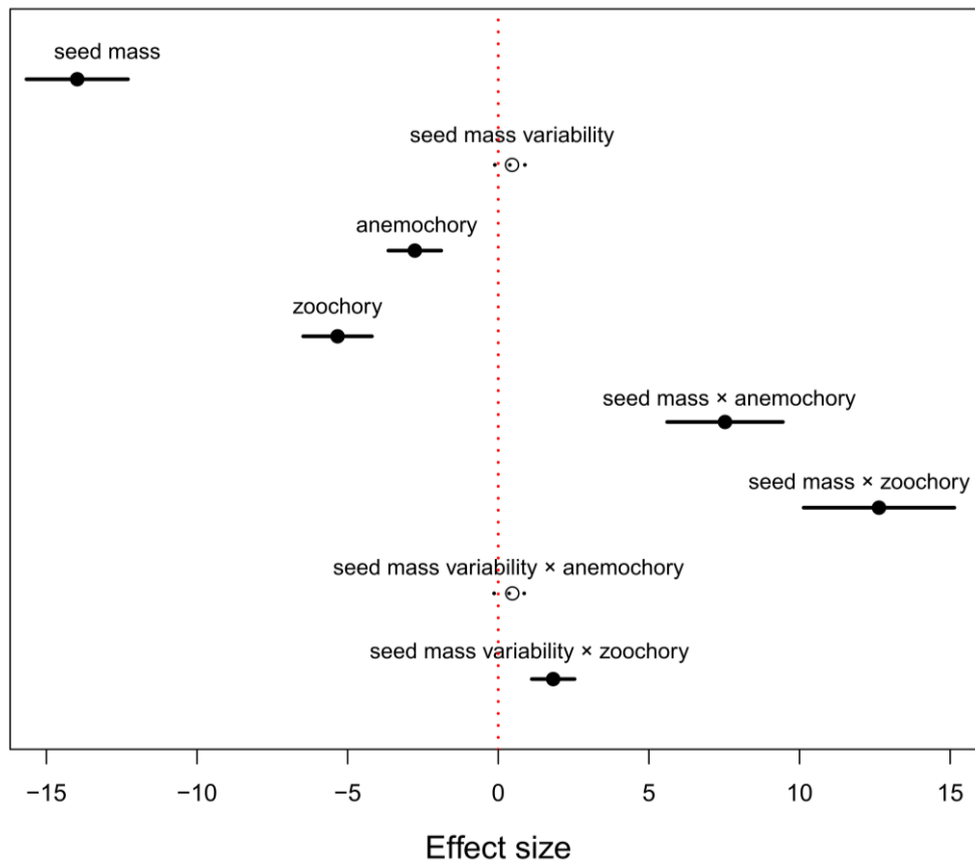
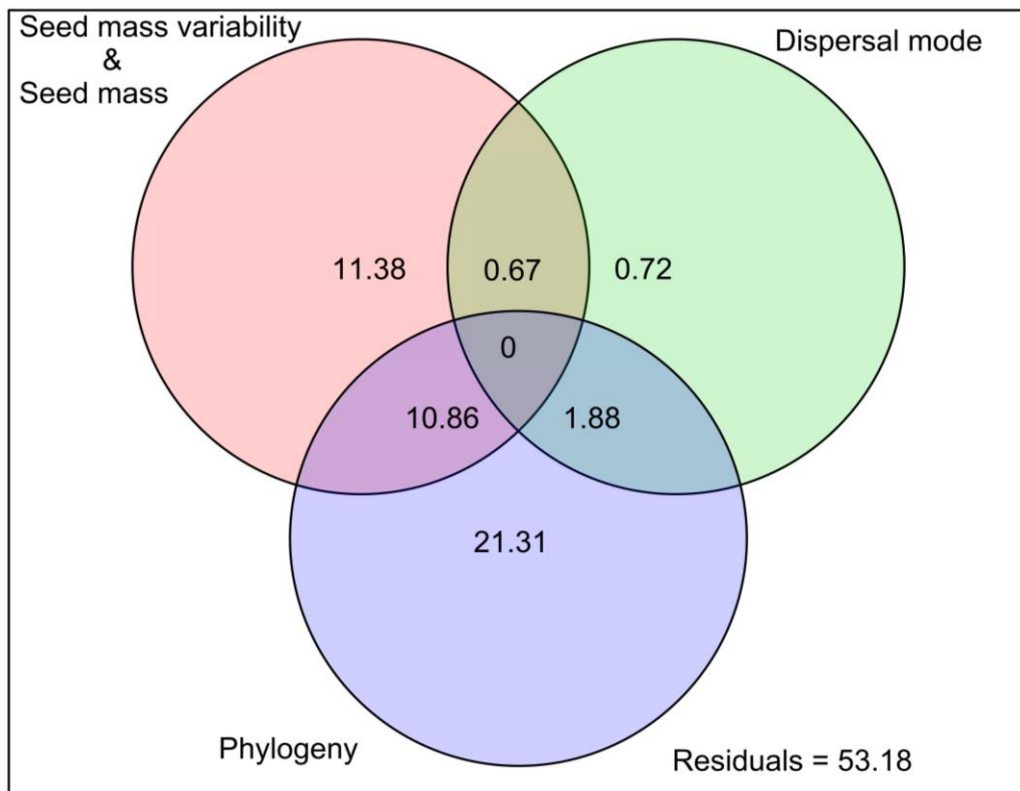


Figure 1. Effects of seed mass and seed mass variability on species distributional range size in autochorous, zoochorous and anemochorous species. In the PGLS model, autochory was treated as a baseline dispersal mode. The black segments represent the effect sizes are statistically significantly different from 0 ($P < 0.05$), while the pointed lines with open circle indicate non-significant effect sizes.

3.3 Joint effects of seed traits and phylogeny on species' range size

220 Variation partitioning showed that the effects of seed mass, seed mass variability, dispersal mode and phylogeny together explained 46.82% of the variance of species' range size (Fig. 2). Of the explained variation, seed mass (including mass variability) contributed a pure 11.38% fraction, phylogeny contributed a pure 21.31%, and a small fraction from the pure dispersal mode (0.72%). We also noted a considerable joint effect of seed traits and phylogeny (13.41%) on species' range size (Fig. 2).



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Figure 2. Variation partitioning of seed mass, seed mass variability, dispersal mode, and phylogeny for species' range size.

4 Discussion

4.1 The relationship between phylogeny and species distributional range size

230 We found a significant phylogenetic signal associated with species distributional range size. This result suggests that closely related species are more similar in distribution range size than distantly related species. It corroborates some studies (e.g., Hunt et al., 2005; Martin and Husband, 2009), but does not support those of Webb and Gaston (2003) which showed the distributional range sizes of closely related species were not more similar to each other than expected by chance. This discrepancy may be due to
235 the different evolutionary history of the studied taxa as well as the heritability of their life-history traits, which can play a critical role in the establishment and persistence of species, and thus influence their distributional range sizes (Angert and Schemske, 2005; Umaña et al., 2018). It is worth noting that Webb and Gaston (2003) studied birds that have much stronger dispersal ability than seed plants, which may explain the difference between the two studies. Seed traits associated with range size can also
240 change over evolutionary time, which in turn could alter the range size of a species' distribution (Blomberg et al., 2003). Furthermore, the geographic distribution range of a species can be influenced by its ecological tolerances associated with life-history traits (Geber and Griffen, 2003; Latimer and Zuckerberg, 2021). Our results imply that the geographic distribution of related plant species may have a similar response to patterns of climate change at a regional scale, due in part, to phylogenetic
245 constraints on the distributional range of species. Here, it seems likely that closely related species have commonly evolved seed traits that result in shared adaptative strategies to climate change, although this causal mechanism requires further empirical study in the field.

4.2 Effects of seed traits on the distribution of species

We found a very strong negative relationship between seed mass and species range size, meaning larger
250 seeds having smaller range size (Fig. 1, Table A1). This result is consistent with previous studies that also found a significant relationship between seed mass and range size (Morin and Chuine, 2006;

Procheş et al., 2012). Different from the effect of seed mass, seed mass variability had no or a weak positive association with distributional range size.

The PGLS model showed that the range sizes of zoochorous (animal-dispersed) and anemochorous (wind-dispersed) species were significantly smaller than that of autochorous (explosive/gravity dispersed) species (Fig. 1). This may appear counterintuitive at the first glance but was resulted after the effects of the interactions between seed mass (and mass variability) and dispersal mode were taken accounted. These strong positive interaction terms (except the interaction between seed mass variability and wind dispersal) shown in Fig. 1 indicate that the range sizes of species with different dispersal modes are strongly subject to seed mass (and also mass variability). For example, zoochorous species with large seed mass and mass variability have significantly larger range size than species that have similar seed traits but dispersed by explosive gravity. This dependence of species distributional range size on the interactions between seed mass and dispersal mode is further confirmed by a simpler PGLS model that excludes all the interactive terms between seed mass (and mass variability) and dispersal mode. The results of this model in Appendix Table A2 show that zoochorous species had significantly larger range size than that of autochorous and anemochorous species ($P < 0.001$), while the latter two groups were not significantly different ($P = 0.257$).

Although intraspecific seed mass variability did not seem to affect distributional range size of autochorous and anemochorous species, the variability was strongly positively associated with range size of zoochorous species. This may be because species with large variation in seed mass could have greater colonization ability in various habitats and seeds of zoochorous species with long dispersal distance have more chances to arrive at heterogeneous habitats than seeds of autochorous and anemochorous species. Given that small- and large-seeded species are shown to adapt to different habitats (Silvertown, 1989), it seems likely that zoochorous species may experience trade-offs between competition ability and dispersal ability through seed mass variation (Chen et al., 2018), resulting in a

similar effect for seed mass on species distributional range size at the geographic scale.

It is interesting to note that Sides et al. (2014) found that species with greater intraspecific variation in specific leaf area (SLA) have wider ecological breadth. Due to its potential role in modulating the response of plant species to environmental changes, greater intraspecific functional variability enables species to adjust to a wider range of competitive and abiotic conditions (Sides et al., 2014; Basnett and Devy, 2021). Plastic responses of seed mass to heterogeneous environments may be related to molecular signals at a single gene or across the entire genome (Nicotra et al., 2010) and thus influence the distributional range size of species (Savolainen et al., 2007). Distributional patterns of plant species may reflect the fact that individuals within a species have different levels of genetic variation in association with seed mass, thus facilitating the species to adapt to a broad spectrum of environments (Völler et al., 2012).

4.3 Effects of seed mass, seed dispersal and phylogeny on species' range size

Our results show that seed traits and phylogeny jointly affect species distributional range size, indicating that species distribution may be limited by ecological and evolutionary processes (Fig. 2). There are two possible reasons for this relationship: (1) the evolution of both seed mass and dispersal mode is phylogenetically conserved (Gallagher and Leishman, 2012; Chen et al., 2018; Kang et al., 2021); and (2) seed mass and seed dispersal mode are not evolutionarily independent but are constrained by evolutionary history, e.g., phylogenetic divergence in dispersal syndrome is related to divergences in seed mass (Moles et al., 2005). However, we also need to recognize that more than 50% of the variance in species distribution in our study remains unexplained. This result suggests that climatic tolerance, competition, colonization ability and other geographic factors could also be important for affecting species distribution (Morin and Chuine, 2006).

5 Conclusions

This study provides evidence that seed mass, intraspecific seed mass variation, seed dispersal mode and phylogeny contribute to explaining species distribution variation on the geographic scale. We found that (1) species distributional range size was significantly constrained by phylogeny, seed mass and its intraspecific variability, and seed dispersal mode; (2) the effects of seed mass and seed mass variability on species distribution varied among dispersal modes; and (3) seed mass, dispersal mode and phylogeny together explained 46.82% of the variance associated with species distributional range size. Despite that more than half of the variation in species distribution is left unexplained, our study clearly shows the importance of including seed life-history traits in modeling and predicting the impact of climate change on species distribution of seed plants.

Data availability. The data are available from the freely accessible databases cited in the manuscript.

Authors contribution. DZL, LMG and FH designed the study; KC and XYY collected data; KC conducted statistical analysis and generated the graphs; KC, KSB and LMG wrote the manuscript; DZL, FH and XYY revised the manuscript. All authors reviewed and approved the final manuscript.

Competing interests. All authors have no conflict of interest.

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References

- 325 Akaffou, S.D., Kouame, A.K., Gore, N.B.B., Abessika, G.Y., Kouassi, H.K., Hamon, P., Sabatier, S., and
Duminil, J.: Effect of the seeds provenance and treatment on the germination rate and plants
growth of four forest trees species of Côte d'Ivoire, *J. For. Res.*, 32, 161-169,
<https://doi.org/10.1007/s11676-019-01064-y>, 2021.
- 330 Angert, A.L. and Schemske, D.W.: The evolution of species' distributions: Reciprocal transplants across
the elevation ranges of *Mimulus cardinalis* and *M. lewisii*, *Evolution*, 59, 1671-1684,
<https://doi.org/10.1111/j.0014-3820.2005.tb01817.x>, 2005.
- Arène, F., Affre, L., Doxa, A., and Saatkamp, A.: Temperature but not moisture response of germination
shows phylogenetic constraints while both interact with seed mass and lifespan, *Seed Sci. Res.*, 27,
110-120, <https://doi.org/10.1017/S0960258517000083>, 2017.
- 335 Basnett, S. and Devy, S.M.: Phenology determines leaf functional traits across *Rhododendron* species in
the Sikkim Himalaya, *Alp. Botany*, 131, 63-72, <https://doi.org/10.1007/s00035-020-00244-5>, 2021.
- Blomberg, S.P., Garland, T., JR., and Ives, A.R.: Testing for phylogenetic signal in comparative data:
Behavioral traits are more labile, *Evolution*, 57, 717-745,
<https://doi.org/10.1111/j.0014-3820.2003.tb00285.x>, 2003.
- 340 Borcard, D., Gillet, F., and Legendre, P. (Eds.): Numerical ecology with R, Springer, New York,
America, 2018.
- Brown, J.H., Stevens, G.C., and Kaufman, D.M.: The geographic range: Size, shape, boundaries, and
internal structure, *Annu. Rev. Ecol. Syst.*, 27, 597-623,

<https://doi.org/10.1146/annurev.ecolsys.27.1.597>, 1996.

- 345 Bu, H.Y., Zhang, Y.M., Zhao, D., Wang, S.Y., Jia, P., Qi, W., Liu, K., Xu, D.H., Ge, W.J., and Wang, X.J.: The evolutionary correlation associated with seed mass and altitude on nutrient allocation of seeds, *Seed Sci. Res.*, 29, 38-43, <https://doi.org/10.1017/S0960258518000387>, 2019.
- Chen, K., Burgess, K.S., Yang, X.Y., Luo, Y.H., Gao, L.M., and Li, D.Z.: Functional trade-offs and the phylogenetic dispersion of seed traits in a biodiversity hotspot of the Mountains of Southwest China, *Ecol. Evol.*, 8, 2218-2230, <https://doi.org/10.1002/ece3.3805>, 2018.
- 350 Chen, S.C., Pahlevani, A.H., Mal'kov á L., Riina, R., Thomson, F.J., and Giladi, I.: Trade-off or coordination? Correlations between ballochorous and myrmecochorous phases of diplochory, *Funct. Ecol.*, 33, 1469-1479, <https://doi.org/10.1111/1365-2435.13353>, 2019a.
- Chen, S.C., Tamme, R., Thomson, F.J., and Moles, A.T.: Seeds tend to disperse further in the tropics, *Ecol. Lett.*, 22, 954-961, <https://doi.org/10.1111/ele.13255>, 2019b.
- 355 Chen, T.C. and Valone, T.J.: Rodent granivory strengthens relationships between seed size and plant abundance in a desert annual community, *J. Veg. Sci.*, 28, 808-814, <https://doi:10.1111/jvs.12529>, 2017.
- Cochrane, A., Yates, C.J., Hoyle, G.L., and Nicotra, A.B.: Will among-population variation in seed traits improve the chance of species persistence under climate change?, *Global Ecol. Biogeogr.*, 24, 12-24, <https://doi.org/10.1111/geb.12234>, 2015.
- 360 Coomes, D.A. and Grubb, P.J.: Colonization, tolerance, competition and seed-size variation within functional groups, *Trends Ecol. Evol.*, 18, 283-291, [https://doi.org/10.1016/S0169-5347\(03\)00072-7](https://doi.org/10.1016/S0169-5347(03)00072-7), 2003.
- DeMalach, N., Ron, R., and Kadmon, R.: Mechanisms of seed mass variation along resource gradients, 365 *Ecol. Lett.*, 22, 181-189, <https://doi.org/10.1111/ele.13179>, 2019.

- Donoghue, M.J., Bell, C.D, and Li, J.H.: Phylogenetic patterns in northern hemisphere plant geography, *Int. J. Plant Sci.*, 162, S41-S52, <https://doi.org/10.1086/323278>, 2001.
- Felsenstein, J.: Phylogenies and the comparative method, *Am. Nat.*, 125, 1-15, <https://doi.org/10.1086/284325>, 1985.
- 370 Gallagher, R.V. and Leishman, M.R.: A global analysis of trait variation and evolution in climbing plants, *J. Biogeogr.*, 39, 1757-1771, <https://doi.org/10.1111/j.1365-2699.2012.02773.x>, 2012.
- Gaston, K.J. and Fuller, R.A.: The sizes of species' geographic ranges, *J. Appl. Ecol.*, 46, 1-9, <https://doi.org/10.1111/j.1365-2664.2008.01596.x>, 2009.
- Geber, M.A. and Griffen, L.R.: Inheritance and natural selection on functional traits, *Int. J. Plant Sci.*,
375 164, S21-S42, <https://doi.org/10.1086/368233>, 2003.
- Greene, D.F. and Quesada, M.: Seed size dispersal and aerodynamic constraints within the *Bombacaceae*, *Am. J. Bot.*, 92, 998-1005, <https://doi.org/10.3732/ajb.92.6.998>, 2005.
- Grossenbacher, D., Briscoe-Runquist, R., Goldberg, E.E., and Brandvain, Y.: Geographic range size is predicted by plant mating system, *Ecol. Lett.*, 18, 706-713, <https://doi.org/10.1111/ele.12449>, 2015.
- 380 Hunt, G., Roy, K., and Jablonski, D.: Species-level heritability reaffirmed: A comment on "on the heritability of geographic range sizes", *Am. Nat.*, 166, 129-135, <https://doi.org/10.1086/430722>, 2005.
- Kang, X., Zhou, J., Abuman, Du, G., and Qi, W.: Multi-factor control of seed mass of species on the eastern part of the Qinghai-Tibetan Plateau: Integration of environmental filters, local adaptation and correlated evolution, *Environ. Exp. Bot.*, 187, 104471, <https://doi.org/10.1016/j.envexpbot.2021.104471>, 2021.
- 385 Kubota, Y., Kusumoto, B., Shiono, T., and Ulrich, W.: Environmental filters shaping angiosperm tree assembly along climatic and geographic gradients, *J. Veg. Sci.*, 29, 607-618, <https://doi.org/10.1111/jvs.12648>, 2018.

390 Latimer, C.E. and Zuckerberg, B.: Habitat loss and thermal tolerances influence the sensitivity of
resident bird populations to winter weather at regional scales, *J. Anim. Ecol.*, 90, 317-329,
<https://doi.org/10.1111/1365-2656.13332>, 2021.

Losos, J.B.: Phylogenetic niche conservatism, phylogenetic signal and the relationship between
phylogenetic relatedness and ecological similarity among species, *Ecol. Lett.*, 11, 995-1003,
395 <https://doi.org/10.1111/j.1461-0248.2008.01229.x>, 2008.

Lynch, M.: Methods for the analysis of comparative data in evolutionary biology, *Evolution*, 45,
1065-1080, <https://doi.org/10.1111/j.1558-5646.1991.tb04375.x>, 1991.

Martin, S.L. and Husband, B.C.: Influence of phylogeny and ploidy on species ranges of North
American angiosperms, *J. Ecol.*, 97, 913-922, <https://doi.org/10.1111/j.1365-2745.2009.01543.x>,
400 2009.

Moles, A.T. and Westoby, M.: Seedling survival and seed size: A synthesis of the literature, *J. Ecol.*, 92,
372-383, <https://doi.org/10.1111/j.0022-0477.2004.00884.x>, 2004.

Moles, A.T., Ackerly, D.D., Webb, C.O., Tweddle, J.C., Dickie, J.B., Pitman, A.J., and Westoby, M.:
Factors that shape seed mass evolution, *PNAS*, 102, 10540-10544,
405 <https://doi.org/10.1073/pnas.0501473102>, 2005.

Moles, A.T., Ackerly, D.D., Tweddle, J.C., Dickie, J.B., Smith, R., Leishman, M.R., Mayfield, M.M.,
Pitman, A., Wood, J.T., and Westoby, M.: Global patterns in seed size, *Global Ecol. Biogeogr.*, 16,
109-116, <https://doi.org/10.1111/j.1466-8238.2006.00259.x>, 2007.

Morin, X. and Chuine, I.: Niche breadth competitive strength and range size of tree species: A trade-off
410 based framework to understand species distribution, *Ecol. Lett.*, 9, 185-195,
<https://doi.org/10.1111/j.1461-0248.2005.00864.x>, 2006.

- Mukherjee, J.R., Jones, T.A., Monaco, T.A., and Adler, P.B.: Relationship between seed mass and young-seedling growth and morphology among nine blue bunch wheatgrass populations, *Rangeland Ecol. Manag.*, 72, 283-291, <https://doi.org/10.1016/j.rama.2018.11.006>, 2019.
- 415 Nathan, R.: The challenges of studying dispersal, *Trends Ecol. Evol.*, 16, 481-483, [https://doi.org/10.1016/S0169-5347\(01\)02272-8](https://doi.org/10.1016/S0169-5347(01)02272-8), 2001.
- Nicotra, A.B., Atkin, O.K., Bonser, S.P., Davidson, A.M., Finnegan, E.J., Mathesius, U., Poot, P., Purugganan, M.D., Richards, C.L., Valladares, F., and van Kleunen, M.: Plant phenotypic plasticity in a changing climate, *Trends Plant Sci.*, 15, 684-692, <https://doi.org/10.1016/j.tplants.2010.09.008>,
420 2010.
- Akwood, M.O., Jurado, E., Leishman, M., and Westoby, M.: Geographic ranges of plant species in relation to dispersal morphology, growth form and diaspore weight, *J. Biogeogr.*, 20, 563-571, <https://doi.org/10.2307/2845727>, 1993.
- Pérez-Harguindeguy, N., Díaz, S., Garnier, E., Lavorel, S., Poorter, H., Jaureguiberry, P., Bret-Harte, M.S., Cornwell, W.K., Craine, J.M., Gurvich, D.E., Urcelay, C., Veneklaas, E.J., Reich, P.B.,
425 Poorter, L., Wright, I.J., Ray, P., Enrico, L., Pausas, J.G., de Vos, A.C., Buchmann, N., Funes, G., Quétier, F., Hodgson, J.G., Thompson, K., Morgan, H.D., ter Steege, H., van der Heijden, M.G.A., Sack, L., Blonder, B., Poschlod, P., Vaieretti, M.V., Conti, G., Staver, A.C., Aquino, S., and Cornelissen, J.H.C.: New handbook for standardised measurement of plant functional traits
430 worldwide, *Aus. J. Bot.*, 61, 167-234, <http://hdl.handle.net/11299/177647>, 2013.
- Procheş, Ş., Wilson, J.R.U., Richardson, D.M., and Rejmánek, M.: Native and naturalized range size in *Pinus*: Relative importance of biogeography introduction effort and species traits, *Global Ecol. Biogeogr.*, 21, 513-523, <https://doi.org/10.1111/j.1466-8238.2011.00703.x>, 2012.
- Qi, W., Guo, S., Chen, X., Cornelissen, J.H.C., Bu, H., Du, G., Cui, X., Li, W., and Liu, K.:
435 Disentangling ecological allometric and evolutionary determinants of the relationship between

seed mass and elevation: Insights from multiple analyses of 1355 angiosperm species on the eastern Tibetan Plateau, *Oikos*, 123, 23-32, <https://doi.org/10.1111/j.1600-0706.2013.00448.x>, 2014.

Jin, Y. and Qian, H.: *VPhyloMaker*: An R package that can generate very large phylogenies for vascular plants, *Ecography*, 42, 1353-1359, <https://doi.org/10.1111/ecog.04434>, 2019.

R Core Team.: R: A language and environment for statistical computing, R Foundation for Statistical Computing, Vienna, Austria. URL <https://www.R-project.org/>, 2020.

Rozendaal, D.M.A., Hurtado, V.H., and Poorter, L.: Plasticity in leaf traits of 38 tropical tree species in response to light; relationships with light demand and adult stature, *Funct. Ecol.*, 20, 207-216, <https://doi.org/10.1111/j.1365-2435.2006.01105.x>, 2006.

Savolainen, O., Pyhäjärvi, T., and Knürr, T.: Gene flow and local adaptation in trees, *Annu. Rev. Ecol. Evol. S.*, 38, 595-619, <https://doi.org/10.1146/annurev.ecolsys.38.091206.095646>, 2007.

Sides, C.B., Enquist, B.J., Ebersole, J.J, Smith, M.N., Henderson, A.N, and Sloat, L.L.: Revisiting Darwin's hypothesis: Does greater intraspecific variability increase species' ecological breadth?, *Am. J. Bot.*, 101, 56-62, <https://doi.org/10.3732/ajb.1300284>, 2014.

Silvertown, J.: The paradox of seed size and adaptation, *Trends Ecol. Evol.*, 4, 24-26, [https://doi.org/10.1016/0169-5347\(89\)90013-X](https://doi.org/10.1016/0169-5347(89)90013-X), 1989.

Sonkoly, J., Deák, B., Valkó, O., Molnár V., A., Tóthmérész, B., and Török, P.: Do large-seeded herbs have a small range size? The seed mass-distribution range trade-off hypothesis, *Ecol. Evol.*, 7, 11204-11212, <https://doi.org/10.1002/ece3.3568>, 2017.

Swenson, N.G. (Eds.): *Functional and phylogenetic ecology in R*, Springer, New York, America, 2014.

Umaña, M.N., Zhang, C., Cao, M., Lin, L., and Swenson, N.G.: Quantifying the role of intra-specific trait variation for allocation and organ-level traits in tropical seedling communities, *J. Veg. Sci.*, 29, 276-284, <https://doi.org/10.1111/jvs.12613>, 2018.

- 460 Valladares, F., Wright, S.J., Lasso, E., Kitajima, K., and Pearcy, R.W.: Plastic phenotypic responses to light of 16 congeneric shrubs from a Panamanian rain forest, *Ecology*, 81, 1925-1936, [https://doi.org/10.1890/0012-9658\(2000\)081\[1925:PPRTLO\]2.0.CO;2](https://doi.org/10.1890/0012-9658(2000)081[1925:PPRTLO]2.0.CO;2), 2000.
- Völler, E., Auge, H., Prati, D., Fischer, M., Hemp, A., and Bossdorf, O.: Geographical and land-use effects on seed-mass variation in common grassland plants, *Basic Appl. Ecol.*, 13, 395-404, 465 <https://doi.org/10.1016/j.baae.2012.06.006>, 2012.
- Webb, T.J. and Gaston, K.J.: On the heritability of geographic range sizes, *Am. Nat.*, 161, 553-566, <https://doi.org/10.1086/368296>, 2003.
- Yang, X., Huang, Z., Venable, D.L., Wang, L., Zhang, K., Baskin, J.M., Baskin, C.C., and Cornelissen, J.H.C.: Linking performance trait stability with species distribution: The case of *Artemisia* and its 470 close relatives in northern China, *J. Veg. Sci.*, 27, 123-132, <https://doi.org/10.1111/jvs.12334>, 2016.
- Zanne, A.E., Tank, D.C., Cornwell, W.K., Eastman, J.M., Smith, S.A., FitzJohn, R.G., McGlinn, D.J., O'Meara, B.C., Moles, A.T., Reich, P.B., Royer, D.L., Soltis, D.E., Stevens, P.F., Westoby, M., Wright, I.J., Aarssen, L., Bertin, R.I., Calaminus, A., Govaerts, R., Hemmings, F., Leishman, M.R., Oleksyn, J., Soltis, P.S., Swenson, N.G., Warman, L., and Beaulieu, J.M.: Three keys to the 475 radiation of angiosperm into freezing environments, *Nature*, 506, 89-92, <https://www.nature.com/articles/nature12872>, 2014.
- Zhou, Q., Wu, J., Cui, X., Li, X., Liu, Z., Musa, A., Ma, Q., Yu, H., Liang, W., Jiang S., and Wang, Y.: Geographical distribution of the dispersal ability of alien plant species in China and its socio-climatic control factors, *Sci. Rep.*, 11, 7187, <https://doi.org/10.1038/s41598-021-85934-8>, 480 2021.

APPENDICES

485 Table A1. The phylogenetic generalized least squares regression for modeling the effects of seed mass, seed mass variability, dispersal mode, seed mass \times dispersal mode and seed mass variability \times dispersal mode interaction terms on species distributional range size. The graphic presentation of the results of this table is given in Figure 1 in the main text.

Variable	Effect size \pmSE	<i>t</i>-value	<i>P</i>-value
Intercept	18.406 \pm 5.612	3.279	0.001
Seed mass	-13.974 \pm 0.842	-16.593	<0.001
Seed mass variability	0.459 \pm 0.286	1.604	0.109
Anemochory	-2.769 \pm 0.438	-6.318	<0.001
Zoochory	-5.333 \pm 0.570	-9.358	<0.001
Seed mass \times anemochory	7.527 \pm 0.960	7.838	<0.001
Seed mass \times zoochory	12.637 \pm 1.250	10.105	<0.001
Seed mass variability \times anemochory	0.468 \pm 0.303	1.545	0.123
Seed mass variability \times zoochory	1.824 \pm 0.355	5.140	<0.001

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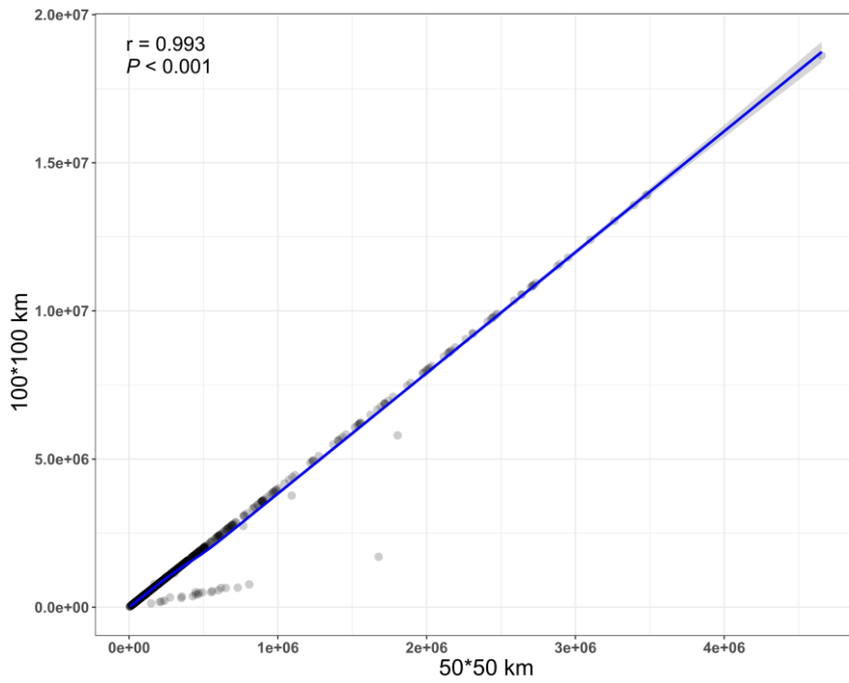
Table A2. The phylogenetic generalized least squares regression for modeling the effects of seed mass, seed mass variability and dispersal mode, without interaction terms, on species distributional range size.

505 In the model, autochory (explosive/gravity dispersal) was treated as the baseline dispersal mode. The results in the table show zoochorous species had significantly larger range size than that of autochorous species ($P < 0.001$), while the range size of anemochorous (wind dispersal) species and that of autochorous species were similar ($P = 0.257$).

Variable	Effect size \pmSE	<i>t</i>-value	<i>P</i>-value
Intercept	16.018 \pm 5.988	2.675	0.008
Seed mass	-7.424 \pm 0.422	-17.611	<0.001
Seed mass variability	1.1 \pm 0.092	11.974	<0.001
Anemochory	0.323 \pm 0.285	1.133	0.257
Zoochory	1.16 \pm 0.295	3.928	<0.001

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520 Figure A1. Relationship between distributional range size calculated at the spatial resolution of 50 km and the range size calculated at the spatial resolution of 100 km.