Comparisons in the native and introduced ranges reveal little evidence of climatic adaptation in germination traits

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1 Abstract

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Plant invasions are increasing due to globalization and environmental change, including through anthropogenic climate change. Yet we lack an understanding of how some species become widespread invaders while others do not. Two competing mechanisms have been posited:

1) post-introduction rapid evolution to the novel environments of the introduced range and 2) broad environmental tolerance in the native population that makes invaders tolerant of diverse introduced environments. Each mechanism has implications for how invaders respond to climate change: either needing to evolve to future climates, or already being tolerant of diverse current/future climates. Disentangling these mechanisms requires investigating how evolution versus tolerance drive essential invasion traits (germination success and timing; growth rate). Here, we tested for evidence of rapid evolution in these traits by using growth chambers to provide common climates for seven herbaceous plant species sampled from multiple populations in their native (European) and introduced (North American) ranges. Chambers provided two levels of stratification—to simulate different winter lengths—and four temperature levels post-stratification—to simulate different spring conditions. We used Bayesian multilevel models to

examine responses, while controlling for population and seed family. Across all species, trait responses were largely similar between native and introduced populations, except in response to particular climates representing cold winters and warm springs (where introduced populations germinated later and grew faster). Our results suggest that broad environmental tolerance, not rapid evolution, likely underlies invasion success for these invaders—and may sustain their spread with continued warming—but suggests that species may evolve in response to specific combinations of winter and spring climatic regimes.

Keywords: Climate change ecology, Invasion ecology, Rapid evolution, Broad environmental tolerance, Phenology, Plant-climate interactions, Growth chamber experiment, Germination, Bayesian multilevel models, Invasive plants.

26 1 Introduction

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- Exotic plant invasions can transform biodiversity and ecosystems (Bellard et al., 2016; Mack et al., 2000; Pejchar & Mooney, 2009). These invasions are increasing: globalization is facilitating extra-
- range plant dispersal (Helmus et al., 2014), and human alteration of ecosystems may provide new
- niche space (Blois et al., 2013; Harte et al., 2015; Inouye, 2008; Tilman & Lehman, 2001). Upon
- dispersing to a new environment, introduced species can thrive by filling vacant niches (Elton, 1958)
- or outperforming native plants (Daehler, 2003; Davis & Pelsor, 2001).
- 33 Changing environments, especially with anthropogenic climate change, could select for species
- that can take advantage of newly created temporal niches (Godoy & Levine, 2014; Wolkovich &
- 35 Cleland, 2011), and related resources, through shifts in the timing of flowering, fruiting, and other
- 36 life history events (Franks et al., 2007). Recent research shows invaders differ in their sensitivity to
- 37 climate (Reeb et al., 2020) and are able to shift their phenology more than native species in their
- introduced communities (Reeb et al., 2020; Wolkovich & Cleland, 2014; Zettlemoyer et al., 2019).
- 39 But whether these invaders are inherently more phenologically flexible (i.e., similar phenological
- 40 responses across both their native and introduced ranges), or whether their phenology can rapidly
- evolve post-introduction has received little attention.
- Two major biological mechanisms for how plants become widespread in novel environments are par-
- 43 ticularly relevant to understanding how phenology and invasions may intersect: 1) post-introduction

rapid evolution and 2) broad environmental tolerance in the native population. A large body of literature supports this first mechanism of rapid evolution (e.g., Clements & Ditommaso, 2011; Colautti & Lau, 2015; Lee, 2002; Prentis et al., 2008; Reznick & Ghalambor, 2001). Rapid evolution can enable introduced species to adapt to vacant niches and take advantage of variable and highresource environments. This includes by evolving greater competitive ability when released from natural enemies (Blossey & Notzold, 1995; Bossdorf et al., 2005, i.e., enemy release hypothesis; though this hypothesis is likely less explanatory than is often assumed Colautti et al., 2004), or by evolving adaptive plasticity (Richards et al., 2006). For example, a study found that genetic adaptation drove adaptive phenotypic variation in flowering time between high-altitude and desert populations of Capsella bursa-pastoris (Brassicaceae) in California (Linde et al., 2001). Invasion can expose populations to divergent selection regimes; in one case this has even led to reproductive isolation and thus speciation in as few as 13 generations (Hendry et al., 2000). If post-introduction rapid evolution is this central to invader success, it would have important implications for invasive species management: managers should treat invasives not as static, homogeneous species, but as constantly adapting populations (Lee, 2002). It would also suggest invaders will continually genetically evolve with climate change and thus estimates of their responses today may not forecast their future climatic responses.

Despite the support for the importance of post-introduction rapid evolution for widespread invaders, a competing body of literature suggests that invaders need not evolve to become widespread in novel environments. Instead, broad environmental tolerance, plasticity, weediness, and generalist adaptations to human-dominated environments within the native population may give invaders sufficient advantages to become widespread, obviating the necessity of post-introduction rapid evolution (Baker, 1965; Bock et al., 2015; Rejmanek & Richardson, 1996; Richards et al., 2006; Schwartz, 1994). A meta-analysis of 117 studies found that invasion success was associated with general performance-related traits, and concluded that it may be possible to predict future invaders by these traits (van Kleunen et al., 2010). In contrast to the rapid evolution hypothesis outlined above, this model of invasions emphasizes invasion prevention and, for invasions that cannot be prevented, suggests treating them as a homogeneous population across their introduced range. It would also suggest that today's estimates of invaders' responses to climate can be used to forecast

73 their future performance and, potentially, their future distributions with climate change.

While these two hypotheses—post-introduction evolution or broad environmental tolerance—are not exhaustive, they represent two major mechanisms that could explain observed differences in the phenological flexibility of invaders (Reeb et al., 2020; Wolkovich & Cleland, 2014; Zettlemoyer et al., 2019) and could be tested by exposing populations from both the introduced and native 77 ranges to common climates. To date most research on the phenology of invaders has focused on the invaders in their introduced communities, often using observational datasets (e.g., Wolkovich et al., 2013) or experimental warming in the field (e.g., Zettlemover et al., 2019). But neither 80 of these methods or even single-location common gardens (i.e., testing individuals from only one 81 part of the range or in only one site, Conner & Hartl, 2004; Vitasse et al., 2009) are sufficient to discriminate the two mechanisms. Reciprocal common garden experiments—with native and invader populations—can test these theories (e.g., Lamarque et al., 2015; Williams et al., 2008), but they are relatively rare and typically only include one or two species due to the immense effort they require. Growth chamber experiments are easier to control and execute, thereby enabling a larger number of species to be tested and compared simultaneously. Moreover, growth chambers can precisely vary the environments that plants experience and provide high-resolution assessment of small differences in trait responses.

Here, we report on a growth chamber experiment of seven highly invasive herbaceous plant species collected from their native (Europe) and introduced (North America) ranges, many of which appear responsive to climate (Wolkovich & Cleland, 2014). Four of our seven study species (Capsella bursa-pastoris, Chelidonium majus, Plantago lanceolata, and Rumex crispus) were included in a phenology monitoring dataset (the Concord Phenology Dataset, Willis et al., 2008), which showed that these species flower 4.5 days earlier on average than they did in the 1800s (compared to less than a day earlier on average across all 372 species in the dataset). This suggests that these invasive species exhibit flexible phenologies—flexibility that may be key to their success.

While much work in studying invaders' phenology has focused on flowering and leaf-out (e.g., Zohner et al., 2017), we focused on germination and growth traits here as they are some of the most important for granting invasive success (Maillet & Lopez-Garcia, 2000; Sattin & Sartorato, 1997): invasive success requires the capacity to germinate in novel environments and grow rapidly enough

to compete with native flora (see Gioria & Pyšek, 2017, and papers reviewed therein; Chrobock 102 et al., 2011; Grime et al., 1988). Therefore, germination success (whether a seed germinates), ger-103 mination timing (days between exposure to warm temperature and germination), and growth rate 104 (cm/day) may represent key invasion traits. At least some of these traits appear to be sensitive 105 to environmental differences (Leger & Rice, 2007; Paudel & Battaglia, 2013). In particular they 106 should respond strongly to two major germination cues: stratification length and spring temper-107 ature (Finch-Savage & Leubner-Metzger, 2006). In temperate ecosystems, many species require 108 cold stratification, which simulates winter, before their seeds can germinate, a requirement that 109 helps ensure that seeds do not germinate during a mid-winter warm period (Baskin & Baskin. 1998: Popay & Roberts, 1970: Wulff et al., 1994). Not surprisingly then, winter length is a key 111 niche variable (Harte et al., 2015) that may show substantial spatial variation, independent of other 112 climate variables (Bonan, 2003). Given sufficient stratification length, spring temperature dictates 113 the appropriate germination time and growth rate (Egli & Wardlaw, 1980; Guilioni et al., 2003). 114 Based on the importance of winter and spring climates, we designed a full-factorial experiment of 115 two stratification lengths and four spring (post-stratification) temperatures, examining responses 116 of germination success, time to germination, and growth rate of introduced (American) and na-117 tive (European) conspecific populations across the eight climatic regimes. Because these invasive 118 species have flourished and become widespread in their introduced range, we hypothesized that the 110 seeds from the invading populations (North America) will either a) respond differently to spring 120 temperature and stratification treatments than the native populations (Europe)—demonstrating 121 rapid evolution, or b) both introduced and native populations will respond similarly to temperature 122 and stratification treatments, and the most fitness-like trait, germination success, will be high and 123 invariant across the treatments—demonstrating broad environmental tolerance.

125 2 Materials and Methods

126 2.1 Study species

Following Richardson's definition of invasive species (Richardson et al., 2011; Richardson et al., 2000, see Supplement for details), seeds were collected from eight herbaceous species that originated

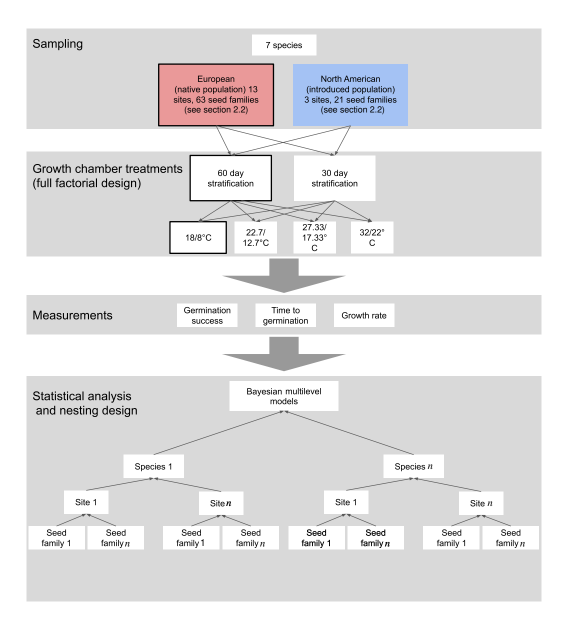


Figure 1: Conceptual diagram showing our methods. We collected seeds from seven species of plants in both their native (pink) and introduced (blue) ranges. These colors are used in figures throughout to refer to these different ranges. We collected seeds from multiple sites and seed families. We then exposed our seeds to eight treatments in a full-factorial design, and measured whether a seed germinated, how long it took to germinate, and how fast it grew. We then modeled how these traits were affected by origin (native vs. introduced) and the eight treatments. Our model accounted for species, site, and seed family variation by using nested random effects. Boxes with bolded outlines represent the reference levels in our models.

in Europe but were introduced to the US, where they have spread and become very widespread (Uva et al., 1997): Alliaria petiolata (Brassicaceae), Capsella bursa-pastoris (Brassicaceae), Chelidonium majus (Papaveraceae), Dactylis glomerata (Poaceae), Plantago lanceolata (Plantaginaceae), 131 P. major, Rumex crispus (Polygonaceae), and Taraxacum officinale (Asteraceae) (see Haines et al. 132 (2011) for authorities). Alliaria petiolata exhibited minimal germination, and so was removed from 133 the analysis. These species represent a mix of perennials, biennials, and annuals. Many were 134 intentionally introduced for medicinal or forage uses (for additional details, including time since 135 colonization, see Supplement). All of these species are weedy, widespread invaders in the US, with 136 many impacting crop production and ecosystems (e.g., Froese & Acker, 2003; Wolfe et al., 2008). 137

138 2.2 Seed collection

We collected mature seeds from European native populations and North American introduced 139 populations from 15 June to 5 September 2015 (see Figure 1 for an overview of our methods). 140 Ideally, our samples would represent the full extent of these species' native and introduced ranges. However, their native ranges span Europe, Asia, and North Africa, while their introduced ranges 142 span North America, with some species flourishing from Florida to Alaska (see Supplement for 143 details). Furthermore, we could visit only a limited number of sites during the time when these species would be producing seeds. Consequently, our sample was not representative of the native 145 and introduced ranges of all species, but instead the result of a more targeted sampling effort in the 146 likely source (Europe) of populations introduced to New England, where invaders have been wellstudied (Willis et al., 2008). European seeds thus came from a total of 63 individuals across 13 sites 148 in nine European countries: Austria, Denmark, France, Germany, Liechtenstein, The Netherlands, 149 Norway, Slovenia, and Switzerland. North American seeds came from a total of 21 individuals 150 across three sites in Massachusetts, USA: Harvard Forest LTER (Petersham) Arnold Arboretum 151 at Harvard University (Boston), and Walden Pond (Concord) (see Figure 2). Multiple seeds were 152 collected from each parent plant (seed family). Elevation ranged from 0-1202 m in Europe and 153 20–300 m in USA. Seeds were collected in paper envelopes and stored at room temperature until early September 2015, when they were cleaned and returned to envelopes. 155

Climate: To examine how climate varied between populations and continents, the mean March,
April, and May temperatures (~1 km² resolution) for 1970-2000 for each population location were
downloaded from WorldClim Version 2 (Fick & Hijmans, 2017) and compared (see Figure 2).
Climates were similar in the native/introduced populations, but showed differences that may be
sufficient to drive populations to adapt after invasion, including overall colder March temperatures
and warmer May temperatures in the introduced range.

162 2.3 Experimental Design

To test phenological responses to climate, seeds were exposed to eight treatments representing vary-163 ing climates. Seeds were first subjected to either a long or short stratification treatment, and then planted in one of four spring temperature treatments. All treatments were carried out in growth 165 chambers. For each treatment, 20 representatives of each species (with seven species this equals 140 166 seeds per treatment) and an additional five representatives from each site of Plantago lanceolata 167 (the most heavily sampled species, with 13 populations) leading to a total of 205 seeds per treatment. Site representatives were drawn from the greatest array of seed families (ranging from 8-48 169 seeds/seed family; this inequality was addressed using multilevel models—see 'Statistical Analysis,' 170 below), and seed family representation was equal across treatments. While when aggregated across all species, the total number of sample sites was greater in the native range (13), the number of 172 sites representing each species were usually similar in both ranges. Thus, native/introduced ranges 173 were evenly represented for most species (except for Plantago lanceolata, which had more samples 174 from the native population). 175

76 2.4 Stratification

We stratified all seeds at 4°C, 70% humidity, 380 ppm of CO_2 (e.g., Meekins & McCarthy, 1999; Popay & Roberts, 1970) on moistened Whatman 1 qualitative filter paper in sterile, vented, lightversion Greiner bio-one 94x16 petri dishes in darkness (Baskin & Baskin, 1998; Popay & Roberts, 1970) in a single Biochambers TPC-19 Reach-In Growth Chamber for either 30 days (reference level) or 60 days. These two stratification treatments represent intermediate stratification lengths for our species: studies show that our species require stratification lengths between 16 days (Popay

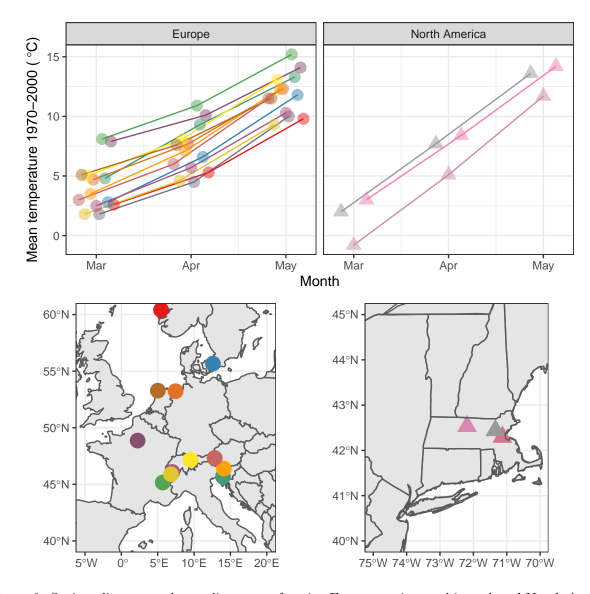


Figure 2: Spring climates and sampling map of native European sites and introduced North American (New England) sites. Climate shows average (1970–2000) March, April, and May temperatures at each site. Note that spring temperature at native (European) sites are similar to spring temperature experienced by introduced (North American) sites, but also show key differences: introduced sites are exposed to greater increases in temperature from March to May. Data from WorldClim Version 2 (Fick & Hijmans, 2017).

& Roberts, 1970) and 120 days (Meekins & McCarthy, 1999). We began the 60-day stratification treatment in late September 2015; other seeds remained in paper envelopes at room temperature until they were stratified in late October 2015. Water was added to petri dishes every 30 days.

186 2.5 Germination

On November 23, 2015, seeds from both stratification treatments were transferred into individual pots with soil (see Experimental Design, above), which were placed into four different growth chambers (three Biochambers TPC-19 and one Biochambers LTCB-19 Reach-In Growth Chamber) and subjected to four different germination treatments. Temperature varied across treatments—all other measured variables were kept constant, and treatments were rotated through growth chambers to control for unmeasured chamber effects. (Seeds that germinated during stratification were not included in the analysis, but this was a small number and unlikely to affect results.)

Germination Temperature: Our four treatments used four temperatures between 18 and 32°C (see next paragraph). Optimal weed germination typically occurs at 20-30°C (Hartmann & Kester, 2010; Popay & Roberts, 1970; Steinbauer & Grigsby, 1957; Wulff et al., 1994). We used this sightly broader spectrum to ensure a sufficient variance in germination response.

Thermoperiocity: Our treatments employed daily fluctuations in temperature (thermoperiocity) of 10°C (see e.g., International Seed Testing Association, 1954; Steinbauer & Grigsby, 1957; Toole, 1963), translating to treatment temperatures of: 18/8°C (reference temperature), 22.67/12.67°C, 27.33/17.33°C, and 32/22°C. For conciseness, in the following figures and text, we refer to temperature treatments by just the high temperature value. All treatments were subjected to 8 hours at the high temperature and the remaining 16 hours at the low temperature (Baskin & Baskin, 1998; Popay & Roberts, 1970; Probert, 2000; Roberts & Totterdell, 1981).

Light type, period, & luminance: We used T5HO fluorescent lights (Toole, 1963), which have a high R:FR ratio as exposure to a high R:FR ratio generally increases germination success (though some studies find germination requires high R:FR ratio or is insensitive, Pons, 2000; Popay & Roberts, 1970; Wulff et al., 1994). We exposed all treatments to eight hours (coinciding with the higher temperature, Baskin & Baskin, 1998) of 75 micromol/m²/second, which yielded a daily

photon dosage of 2.16 mol/m². This amount of light should be sufficient to evoke germination response in all species (Pons, 1991). Because none of our species are known to exhibit highirradiance response and growth chambers provide less light than normal natural conditions, we erred on the side of high light (see Supplement for additional details).

Planting substrate & water: We planted each seed in its own tray cell, on top of Fafard Growing Mix (a mixture of fine peat moss, fine perlite, and vermiculite) soil. This planting arrangement ensures light availability (Tester & Morris, 1987) and provides higher germination success than filter paper (Andrews & Burrows, 1974). Every two days, seeds were watered until all of the soil had become wet (Steinbauer & Grigsby, 1957); but not so much that a film of water covered the seeds (Association of Official Seed Analysts, 1960).

Germination and growth rate monitoring: Collection of germination and growth data was 220 masked to population. Seeds were checked during the light period for germination every two days. 221 Germination was defined as the growth of shoot or radical through the seed coat (Baskin & Baskin. 222 1998; Popay & Roberts, 1970). Germination date for each seed was recorded. Germination was 223 monitored until 29 Jan 2016, for a total observation length of 67 days (this is longer than the typical 224 two-week germination trials according to Baskin & Baskin, 1998; Wulff et al., 1994). Aboveground 225 linear height of each seedling was measured five times: 7 Dec 2015, 15 Dec 2015, 21 Dec 2015, 4 Jan 2016, and 29 Jan 2016. On 1 Jan 2016, the plants were moved from the growth chambers to 227 a greenhouse subject to the following conditions: natural photoperiod (approximately 10 hours of 228 light/day), 20 to 25°C, and 65% humidity.

230 2.6 Statistical analysis

To test for evidence of post-introduction rapid evolution across seven species, while accounting for variation due to population and seed family, we used a Bayesian multilevel modeling framework (Carpenter et al., 2017). These multilevel models are robust and generally provide high power and unbiased estimates, especially for fixed effects (Paccagnella, 2011). This approach yielded estimated (fixed) effects that fully incorporate these multiple levels of variance to produce overall estimates both for each species and generalized across species.

Plant height was roughly linear with time (see Figure S1), so growth rate was defined as β in the 237 linear model: $height = \alpha + \beta * day + error$, where error is normally distributed. This growth rate 238 was calculated for each seed that germinated. The four temperature treatments were recoded as 239 three indicator binary factors, allowing non-linear responses to temperature. For all models (growth 240 rate, germination success, and germination timing), stratification length, continental origin, and temperature were treated as binary fixed effects, with the full suite of 2- and 3-way interactions 242 included. Europe, 18/8°C, and 30 days were reference levels for origin, stratification length, and 243 temperature, respectively (see Figure 1). Seed family was treated as a random effect, nested within 244 sampling population, nested within species (with both random slopes and intercepts). Growth rate 245 was modeled with a normal error distribution: 246

 $y_i \sim N(\mu_i, \sigma)$

$$\mu_{i} = \alpha + \beta_{1} \times origin + \beta_{2} \times strat$$

$$+ \beta_{3} \times 22.7^{\circ}C + \beta_{4} \times 27.3^{\circ}C + \beta_{5} \times 32^{\circ}C$$

$$+ \beta_{6} \times origin \times strat + \beta_{7} \times origin \times 22.7^{\circ}C$$

$$+ \beta_{8} \times origin \times 27.3^{\circ}C + \beta_{9} \times origin \times 32^{\circ}C$$

$$+ \beta_{10} \times strat \times 22.7^{\circ}C + \beta_{11} \times strat \times 27.3^{\circ}C$$

$$(2)$$

 $+ \beta_{12} \times strat \times 32^{\circ}C_{+}\beta_{13} \times origin \times strat \times 22.7^{\circ}C$

 $+\beta_{14} \times origin \times strat \times 27.3^{\circ}C + \beta_{15} \times origin \times strat \times 32^{\circ}C)$

(1)

Where the α (intercept) and β (slope) coefficients were all specified with the same normally-distributed nested random effects (γ): seed family nested within sampling population, nested within species—sp[pop[sfamily[i]]] (not shown above). Thus, for each γ in $[\alpha, \beta_1 : \beta_{15}]$:

$$\gamma_{sp[k]} \sim N(\mu_{\gamma}, \sigma_{\gamma})$$
 (3)

$$\gamma_{sp[pop[j]]} \sim N(\mu_{\gamma_{sp[k]}}, \sigma_{\gamma_{sp[k]}}) \tag{4}$$

$$\gamma_{sp[pop[sfamily[i]]]} \sim N(\mu_{\gamma_{sp[pop[i]]}}, \sigma_{\gamma_{sp[pop[i]]}})$$
 (5)

Where sp = species, indexed with k, pop = sampling population, indexed with j, sfamily = seed family, indexed with i, and strat = stratification. Germination success was modeled similarly to growth rate, but using a binomial error distribution and logit link function, while germination timing was modeled with a Poisson error distribution and log link function.

All models were estimated using four chains, each with 2000 iterations (1000 devoted to warmup), and wide priors. All models were built with Stan (Carpenter et al., 2017) using rstanarm
version 2.17.4 (Goodrich et al., 2018) in R (R Development Core Team, 2015). Chain convergence
was confirmed using the Gelman–Rubin statistic/ \hat{R} close to one (Gelman & Rubin, 1992). Model
implementations were validated using simulated data; model fits were assessed using posterior
predictive checks (Gelman et al., 2004).

Average predictive comparisons: The interactions of treatments (stratification and temper-257 ature) and random effects (species, population and seed family) make this model complex, and 258 can make clear interpretations of parameter estimates difficult. To address this, we calculated average predictive comparisons (Gelman & Pardoe, 2007) for each stratification and temperature 260 level. These estimates average over interaction terms and the full mixed (fixed and random) effects, 261 to provide a single estimate per level that includes all modeled uncertainty. Additionally, unlike model output from Poisson and Binomial models, which are given in transformed units, average 263 predictive comparisons yield estimates that are in the units of the dependent variable (but always 264 positive) (Gelman & Pardoe, 2007) and thus allow comparisons across effects. We note that average 265 predictive comparisons can be complicated to implement in certain designs; because we have a full-266 factorial experiment, our stratification and temperature variables are appropriately balanced (i.e., 267

every temperature occurs at every stratification level) and independent (i.e., every combination of input values is equally likely to co-occur), we calculated average predictive comparisons without any weighting requirement, thus simplifying the computation. See Supplement for equations and details.

272 3 Results

Germination success: Germination success was high: across all species, populations, and seed 273 families, 76% of seeds germinated. Multiple seeds from every species germinated in every treatment 274 combination. Overall, germination success was insensitive to stratification, temperature, or origin— 95% credible intervals (henceforth, 'CrI') for all effects were clustered around zero (Figures 4, S2; 276 Table S2). Regardless of the climatic conditions, seeds germinated at fairly constant, high levels. 277 Seeds from the introduced and native ranges germinated at similar levels and responded similarly to treatments (see 'origin,' 'strat,' '22.7°C,' '27.3°C,' '32°C,' 'origin × strat,' 'origin × 22.7°C,' 279 'origin \times 27.3°C,' 'origin \times 32°C,' 'strat \times 22.7°C,' 'strat \times 27.3°C,' 'strat \times 32°C,' 'origin \times strat 280 \times 22.7°C,' 'origin \times strat \times 27.3°C,' 'origin \times strat \times 32°C' labeled in red in Figure 4 and in Table 281 S2). Seeds from different local populations of *Plantago lanceolata* also germinated at similar levels 282 (see Figure S5). 283

Germination timing: The mean time to germination across all species, populations, and seed 284 families was 12.33 days. Overall, stratification and seed origin had no noticeable effect (see 'origin' 285 and 'strat' in Figure 4 and Table S3). All species showed advanced germination at the higher 286 temperatures (see '27.3°C,' and '32°C' in Figures 4, S3; Table S3). However, Plantago lanceolata did show advanced germination in response to temperature × stratification interaction (see Figure S5). 288 Moreover, all species showed a significant positive interaction effect of origin, stratification and the 280 highest temperature (95% CrI: 0.33–0.85 days; see 'origin × strat × 32°C' in Figure 4 and Table S3). 290 That is, the introduced population showed advanced germination at the short stratification/highest 291 temperature combination (Figure 5). Populations showed fairly homogeneous responses, though 292 temperature × stratification interactions did show some inter-population variability (see Figure 293 S5).

Growth rate: The mean growth rate was 1.2 mm/day. Overall, growth rate was the most 295 sensitive response variable to treatments, though it was still unaffected by population origin and stratification alone (see 'origin' and 'strat' in Figures 4, S4; Table S4). Growth was slower at 297 warmer temperatures for all species, but especially Dactylis glomerata (see '22.7°C', '27.3°C', and 298 '32°C' in Figure 4; negative effects indicate slower growth). This effect was larger for each higher 299 temperature; this is in contrast to germination timing, where the change with temperature was 300 more constant (see comparison in absolute change displayed in Figure 3). However, this decreased 301 growth rate at high temperatures was not uniform across all treatments: in response to one of the 302 higher temperatures (27.3°C) seeds of all species from North America grew 0.56mm faster per day 303 (95% CrI: 0.14–0.98) (see 'origin × 27.3°C' in Figure 4 and Table S4). However, this effect was 304 erased in seeds stratified for 30 days (see Figure 5, and 'origin \times strat \times 27.3°C' in Figure 4 and 305 Table S4).

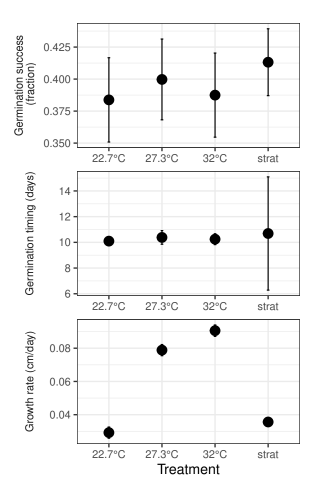
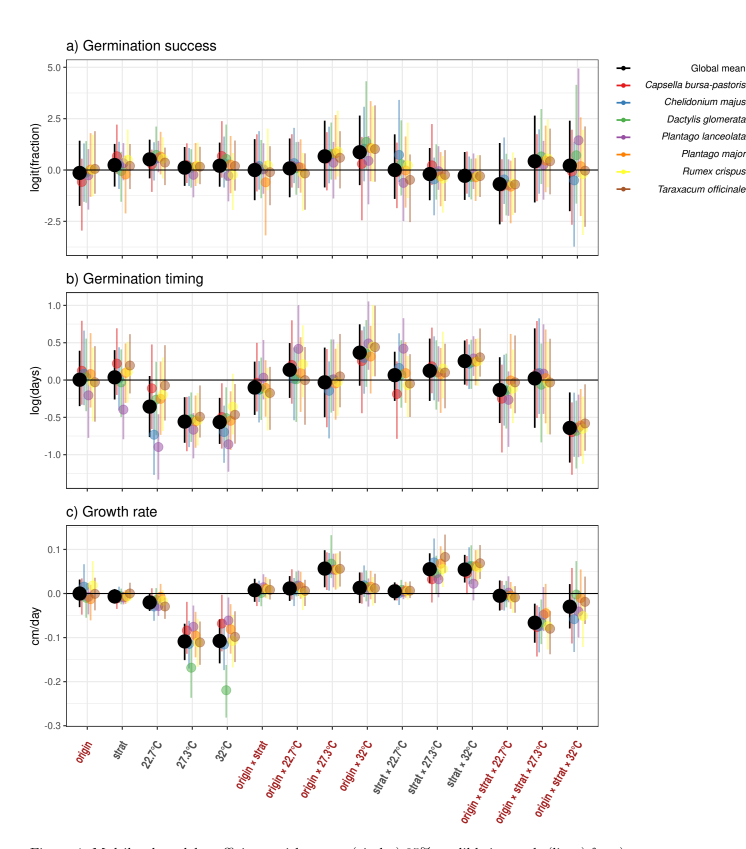


Figure 3: Average predictive comparisons (± standard error) indicating absolute value differences of germination success (top), germination timing (middle), and growth rate (bottom) show how much change in the dependent variable results from a one unit change in the predictor variable while at once integrating over uncertainty from other effects in the model (including population origin). strat refers to the stratification treatment, while 22.7°C, etc., refer to each temperature binary indicator variable. Higher temperatures had indistinguishable effects on germination timing (middle), but sequentially bigger effects on growth rate (bottom). These calculations include all species from both native and introduced populations. See Supplement for further explanation.



Global mean

Dactylis glomerata

Plantago major Rumex crispus

Figure 4: Multilevel model coefficients with means (circles) 95% credible intervals (lines) for a) germination success, b) germination timing and c) growth rate, showing overall effects across species (black circles, 'global mean') and species-specific random effects (colored circles, for intercept coefficients, see Tables S2, S3, S4). The reference level for temperature is a high of 18°C, sixty days is the reference level for stratification and Europe is the reference level for population origin. Parameters testing for effects only of stratification or temperature are in black, while parameter testing for rapid evolution vs. broad environmental tolerance (i.e., those parameters containing 'origin') are in red.

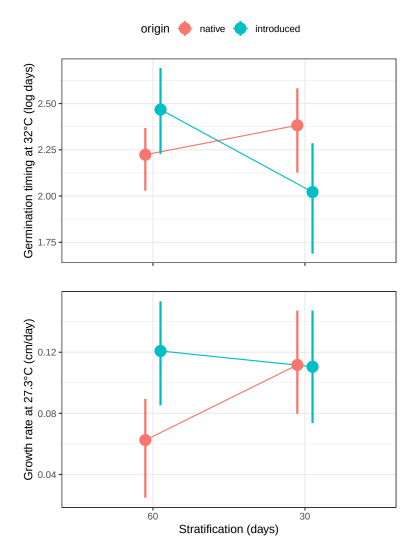


Figure 5: Significant interaction effects of native vs. introduced origin on germination timing (top) and growth rate (bottom). There were no significant effects of origin on germination success. Uncertainty intervals represent posterior predictions across all species (showing 50% credible intervals). For all coefficient estimates, see Figure 4 and Tables S3, S4).

307 4 Discussion

This study leveraged the power of a multi-species growth chamber experiment of native and introduced populations to investigate the importance of post-introduction rapid evolution for widespread
plant invasions across a range of winter-to-spring climatic regimes. All seven widespread, weedy,
highly invasive plant species responded similarly to climate treatments across the subset of populations we studied, suggesting that broad tolerance underlies these invasive species rather than

post-introduction rapid evolution. Across all species, we found only isolated support for post-313 introduction rapid evolution (when winters were long and springs were warm) of key invasion 314 traits—germination success, timing, and growth rate. Instead, our results suggest that these traits 315 do not need to evolve for these species to invade: wide environmental tolerance in the native 316 populations may instead provide sufficient capacity to exploit novel environments (Baker, 1965). 317 Post-introduction rapid evolution may provide a helping hand, but—at least for these traits and 318 for these widespread invaders—rapid evolution does not appear essential for invasion success. This 319 is an encouraging result for forecasts of invader responses to climate change as it suggests we may 320 be able to use current estimates to extrapolate to future responses (up to a point). 321

322 Variation across invasion traits

Our results varied somewhat across traits, highlighting the potential benefits of conditioning biological invasion mechanisms on specific invasion traits (Maillet & Lopez-Garcia, 2000). We found
that post-introduction rapid evolution appeared to play no role in germination success, but may
play a role in germination timing and growth rate—under certain treatment conditions (see Figure

727 5). This suggests that research and theory may be more productive by aiming to identify which
raits are a) broadly tolerant of environmental conditions or b) more likely to rapidly evolve under
results are a) broadly tolerant of environmental conditions or b) more likely to rapidly evolve under

The evidence for broad environmental tolerance in our results was especially pronounced in germina-330 tion success, where all species germinated well, with little regard for climatic conditions. This result 331 suggests that, rather than evolving upon invasion, these widespread invaders drew on the broad 332 environmental tolerance in their native populations. Given the relationship between germination 333 success and fitness (e.g., Domic et al., 2020), this invariant and high germination across climates 334 may be consistent with adaptive phenotypic plasticity (Baker, 1965). Some have suggested that, 335 while initially species may not need to evolve, they may evolve once achieving a foot-hold (Lamar-336 que et al., 2015). However, many of the study species (e.g., Dactylis glomerata) have occupied 337 their introduced range for centuries, yet still show little sign of an evolving, or evolved, germination 338 success in our experiment (i.e., overall, seeds from native and introduced ranges respond similarly 339 to environmental treatments).

Overall, germination timing and growth rate showed few signs of post-introduction evolution. How-

ever, there was some evidence that particular responses have evolved: North American (introduced) populations germinated later and grew faster under longer winter/higher spring temperature combinations (Figure 5). Taking the climate of North American populations into account (Figure 2), 344 this rapid post-introduction evolution of growth rate may be adaptive. North American populations 345 experience climates with longer winter stratification (lower mean March temperatures) and hotter spring temperatures (higher mean May temperature). Thus, the capacity to grow faster after being 347 exposed to a long stratification treatment and high temperatures may provide fitness advantages. 348 Furthermore, germinating later after long winters might help avoid harsh spring conditions (e.g., 349 frosts), which could have costs for later growth and reproduction. Our sampling and modeling framework accounted for variation across seed family, which suggests that our results are not due 351 to maternal effects. Future work designed to estimate maternal effects, as well as consider residual 352 founder effects (Shirk et al., 2014) and genetic drift (Eckert et al., 1996), could provide important insight into the mechanisms of this potential evolution. 354

The convergence between experienced climate and the observed change in growth rate, consistent with adaptive post-introduction rapid evolution, suggests that there is an interdependent
relationship between trait responses and multivariate environments (i.e., seasonal combinations
of winter length and temperature). Considering such interdependencies in the introduced range
may be crucial for predicting how plants evolve post-introduction. Not only can these trait evolution/environment relationships be useful for understanding invasions, they can also help characterize plant capacities to adapt to the multifaceted effects of anthropogenic climate change.

362 Implications for invader responses to climate change

Our evidence for broad environmental tolerance suggests that these widespread invasive species
may continue to perform well with continued climate change, without any evolution in these traits.
This inference may hold for other widespread species, too. Plant invasions have long been used as
a natural experiment for studying plants more generally (e.g., Yoshida et al., 2007). In that light,
these results can be seen as a test of how widespread species may react to rapid climatic change,
where the climate change experienced when a plant colonizes a new environment (i.e., introduced
range) is a proxy for the anthropogenic climate change that plants are experiencing now. Our
results showing that all species germinated earlier at the three higher temperatures, combined with

the invariability of germination success, suggests the prevalence of broad environmental tolerance.

These results indicate that widespread plants may have the capacity to maintain their germination success and germinate/grow rapidly despite the changing climate. While our study focused on widespread invaders, future research should test if temperate plant species with small range sizes share this broad environmental tolerance, or if these localized species may become inferior competitors as the climate changes. If the latter case is true, then climate change may increase the dominance of widespread species.

Our findings that species may adapt their growth rate under certain conditions suggests that invasive species may have the capacity to adapt to the changing winter and spring temperature regimes
that are expected under anthropogenic climate change (IPCC, 2015). If species are adapting to
specific combinations of winter × spring climatic regimes, then forecasters would need to consider
evolutionary responses to multivariate or seasonal environments. Our results also echo the importance of designing experiments that vary both winter length and spring temperature in order to
observe responses to climate change (e.g., Bernareggi et al., 2016).

385 Study limitations & extensions

Our results come from a limited number of individuals and populations collected from the introduced range (see Figure 2; Table S1). The small amount of geographic variation captured in the
introduced range may have introduced bias. Our sampling sites show substantial climate variation
(Figure 2), highlighting potentially important climatic differences, but additional sampling across
the introduced range would provide insights into whether our results generalize to other regions or
if context- (or climate-) dependency is the norm in the introduced range. To this aim, we suggest
future research could build upon our findings by sampling across distinct introduced-range climates
to help understand which traits evolve where, post-introduction.

We harnessed the benefits of growth chambers to provide a common set of precisely controlled multivariate environments for seven species; however, the benefits of this design trade off with a lack of realism. In contrast to reciprocal field common garden experiments, which can integrate important factors (Blois et al., 2013; Germain et al., 2018), our approach lacked most biotic interactions and natural climatic variation. Yet our approach let us tease apart the multivariate nature of climate (stratification × temperature) and examine evidence for post-introduction rapid

evolution across a large range of introduced climates. We believe combining similar growth chamber designs with Bayesian modeling approaches, which integrate across multiple levels of variance (species, population, seed family), provides a tractable approach for other populations, other traits, and other combinations of climate factors (including precipitation). Such future small-scale growth chamber studies could enable robust meta-analyses capable of identifying the traits and climate responses for which post-introduction rapid evolution is, or is not, essential for invasion success, and may guide where best to invest the intensive resources required for reciprocal field common garden experiments.

408 Conclusions

Our results show that post-introduction rapid evolution of germination and growth traits is unlikely to be essential for all plant invasions and that current phenological flexibility seen in invaders was 410 likely present in their native ranges. This suggests that broad environmental tolerance may be 411 important for invasion success in these seven widespread invaders. Post-introduction rapid evolution 412 may still play a role, especially in more extreme or different environments. Linde et al. (2001) found 413 that Capsella bursa-pastoris evolved to colonize high-altitude and desert environments in California. 414 In contrast, our temperate population comparisons showed little sign of rapid evolution, suggesting 415 the tolerance traits contained in temperate native populations may be suitable as long as the 416 introduced environment is not too different (Baker, 1965). Our findings provide support for the 417 speculation by van Kleunen and colleagues (2010) that future invasions can be predicted by species' 418 characteristics (such as broad environmental tolerance), but perhaps only for specific traits (such 419 as germination success). Consequently, managers can perhaps best guard against future invasions 420 by targeting widespread weedy species and preventing them from dispersing beyond their native 421 ranges. Likewise, our results suggest that current estimates of invaders' responses to diverse climates may forecast their future responses under continued climate change. 423

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- tributed critically to the drafts and gave final approval for publication.
- Data, code R code, Stan code, and data will be deposited on the Knowledge Network for Biocomplexity (KNB) repository.

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