Differential response to frequency-dependent interactions: an experimental test using genotypes of an invasive grass

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Differential response to frequency-dependent interactions: an experimental test using genotypes of an invasive grass

Alexandra Collins · E. M. Hart · J. Molofsky

Abstract Positive feedbacks have been suggested as a means for non-indigenous species to successfully invade novel environments. Frequency-dependent feedbacks refer to a species performance being dependent on its local abundance in the population; however, frequency dependence is often described as a monolithic trait of a species rather than examining the variation in response for individual genotypes and fitness traits. Here, we investigate frequency-dependent outcomes for individual genotypes and fitness-related traits for the invasive grass Phalaris arundinacea. We tested for competition-mediated frequency dependence by establishing hexagonal arrays with the center target plant surrounded by either same, different or no genotype neighbors to determine how changing the small-scale frequency neighborhood-influenced invasion success. We used a Bayesian ANOVA approach which allowed us to easily accommodate our non-normal dataset and found that same neighbor plots had greater biomass production than different neighbor plots. Target plants also had greater stem height and aboveground biomass when surrounded by same genotype neighbors. A greenhouse experiment did not support the hypothesis that increased mycorrhizal associations were the cause of positive frequency dependence. We devised a frequency-dependent metric to quantify the extent of fitness-related differences for individual genotypes and found that individual genotypes showed a range of both positive and negative responses to different frequency treatments; however, only positive responses were statistically significant. The small-scale genotypic neighborhood had no effect for the fitness-related traits of leaf number, belowground biomass and total biomass. We demonstrate that individual invasive genotypes respond differently to changing frequency neighborhoods and that growth responses do not respond with the same direction and magnitude. A range of frequency-dependent responses may allow genotypes to invade a wide range of environments.

Keywords Frequency dependence · Intraspecific feedbacks · Mycorrhizae · Phalaris arundinacea · Genotype · Invasive

Introduction

Understanding how non-indigenous species are able to invade and take over native communities has become a central question in ecology. Feedbacks, or the sequence of interactions that determine either a positive or negative effect on a system, have been recognized as an important factor contributing to successful plant invasions due to plant–soil feedbacks (Callaway et al. 2004; Farrer and Goldberg 2009; Klironomos 2002; Laungani and Knops 2009; Nijjer et al. 2007; Reinhart and Callaway 2004; Reinhart et al. 2003; and see reviews by Kulmatiski et al. 2008; Reinhart and Callaway 2006; Wolfe and Klironomos...
2005), fire-suppression feedbacks (Stevens and Beckage 2009), feedbacks with pollinators (Agren 1996) and/or competition-mediated feedbacks among species (Harpole and Suding 2007). Frequency-dependent feedbacks refer to a species performance being dependent on its local abundance in the population and, in general, negative frequency-dependent feedbacks are predicted to maintain species diversity (Bever 2003; Molofsky et al. 2002) while positive frequency-dependent feedbacks are predicted to maintain species uniformity (Bever 2003; Silvertown and Charlesworth 2001; but see Molofsky et al. 2001 for an exception). Thus, invasion potential may be increased if plants exhibit positive frequency-dependent feedbacks (Klironomos 2002; Bever 2003). Yet, most studies have focused only at the species level and our understanding of how intraspecific frequency-dependent interactions influence invasion success remains limited. A fundamental level of variation occurs at the genotype level (Aarssen and Turkington 1983; Antonovics 1976; Harper 1977), and our understanding of how intraspecific frequency-dependent feedback mechanisms increase invasion success may further our ability to predict invasion risk.

Competition-mediated intraspecific frequency-dependent feedbacks driven by neighbors have been recognized for plants (Kelley and Clay 1987; Turkington 1979) and have been shown to influence plant fitness at small spatial scales (Aarssen and Turkington 1985; Antonovics and Ellstrand 1984; Crutsinger et al. 2008). Yet, frequency dependence is generally described as a monolithic trait of all genotypes and all traits (Antonovics and Ellstrand 1984), when in reality genotypes may have considerable variability in their frequency-dependent response. For example, Ronsheim (1996) found that different genotypes of Allium vineale responded differently to neighbor identity. Two out of five genotypes showed evidence for positive frequency dependence while others showed no effect of neighbor identity. Similarly, models investigating frequency dependence often assume that the magnitude and direction of frequency dependence is identical for all individuals (Molofsky and Bever 2002; Molofsky et al. 2001, 2002; but see Eppstein et al. 2006 for an exception). In a two-species model, both the strength and sign of frequency dependence (i.e., whether positive or negative) influenced the patterns of coexistence (Eppstein et al. 2006). Furthermore, the small-scale genotypic diversity of plant populations has been shown to have extended consequences for the diversity of associated herbivores (Crutsinger et al. 2006; Johnson and Agrawal 2005, 2007; Johnson et al. 2006) and for the ability of invasive species to establish (Crutsinger et al. 2008), yet to our knowledge very few studies have examined how individual genotype interactions influence invasive plant performance (but see Vellend et al. 2010).

In addition to understanding the variability in invasive genotype response, determining the mechanisms driving these changes will also better inform our ability to predict genotype trajectories and invasive spread. For plants, negative frequency dependence has been shown to occur through competition and niche partitioning (Antonovics and Kareiva 1988), parasitism (May and Anderson 1983) or mutualism through an asymmetry in benefits to mutualists (Bever 1999), while positive frequency dependence has been shown to occur by preferential predation on rare individuals (Futuyma and Wasserman 1980), host specificity amongst mutualists (Bever 1999) or increased mycorrhizal hyphal connections amongst like genotypes (Ronsheim and Anderson 2001).

Here, we examine the variability of individual genotype response to neighbor identity for multiple growth measures using genotypes of the invasive wetland grass Phalaris arundinacea. Phalaris arundinacea has been shown to have maintained high genotypic diversity in the invasive range (Lavergne and Molofsky 2007) and genotypic diversity is predicted to have its largest ecological effect when the community is dominated by one or a few primary species (Hughes et al. 2008; Whitbam et al. 2006). Our objectives were threefold: (1) to determine the extent of positive or negative frequency-dependent interactions by measuring multiple traits for plant growth, (2) to investigate mycorrhizal hyphal association as the mechanism for positive frequency dependence, and (3) to quantify the extent of growth differences under different small-scale frequency neighborhoods for each individual genotype. We use a flexible statistical approach, Bayesian ANOVA (Qian and Shen 2007), to accommodate both normal and non-normal data. Our work allows for the further understanding of intraspecific frequency-dependent processes and how this may influence invasion success by examining plant performance under different frequency scenarios and the extent to which individual genotypes and growth responses vary in their frequency dependence.

Materials and methods

Study species

Phalaris arundinacea (Poaceae), reed canary grass, is a cool season perennial C₃ grass that is native to temperate zones of the northern hemisphere and is widely distributed throughout Eurasia (Lavergne and Molofsky 2007). Plants grow between 1 and 2 m tall with dense panicles giving rise to both fertile and sterile florets (Gleason and Cronquist 1991). P. arundinacea has high annual seed set as well as a high rate of outcrossing due to self-sterility (Ostrem 1988), and can also reproduce asexually by producing clonal tillers.
**P. arundinacea** is naturally mycorrhizal with colonization varying between 3 and 90% depending on the sampling location (Bauer et al. 2003; Beck-Nielsen and Madsen 2001; Cooke and Lefor 1998; Rickert et al. 1992). An extensive belowground rhizome network makes **P. arundinacea** a competitively dominant species, and it is currently considered an invasive species in many northeastern states of the USA (Kilbride and Pavegli 1999; Lavergne and Molofsky 2004).

**P. arundinacea** is a good study species for experiments on genotypic differences because individuals can be easily genotyped through allozyme screening and rapidly cloned through repeated vegetative tillering. Previous collections of genotypes from the invasive range of **P. arundinacea** have found that genotypic differences amongst plants translate into differences in physiological and morphological characteristics (Broderson et al. 2008; Morrison and Molofsky 1999). Furthermore, differences in these characteristics result in differences in competitive ability and survival (Morrison and Molofsky 1998).

**Frequency dependence experiment**

We experimentally manipulated the intraspecific frequency of genotypes using hexagonal arrays with one “target” plant in the center of the array surrounded by six “neighbor” plants. We used a 3-cm planting density because that corresponds to plant distances observed under field conditions (Collins, personal observation). The experiment consisted of three treatments: (1) target plant with no neighbors, (2) target plant surrounded by six same genotype neighbors, and (3) target plant surrounded by six different genotype neighbors, with each of the six neighbor genotypes representing a different genotype. We established the no neighbor treatment as a control to assess the amount of competition target plants experienced when grown with neighbors. Using the neighbor treatments, we tested for competition-mediated frequency dependence where increased performance of the same neighbor treatment provides support for positive frequency dependence (or reduced intragenotypic competition) and increased performance of the different neighbor treatment provides support for negative frequency dependence (or increased intragenotypic competition). We quantified performance by measuring one fitness response (tiller number) and six growth responses (stem height, leaf number, tiller number and aboveground, belowground and total biomass). Tiller number served as a good proxy for asexual fitness and potential for spread as **P. arundinacea** produced no seeds during the course of our 2-year experiment.

We used a total of seven genotypes and each of the seven genotypes served as the target plant for the three treatments and was replicated three times for a total of 63 plots. All plants were pruned at the time of planting to have two green leaves, 10 cm of stem, 5 cm of roots, 2 cm of rhizome, and one rhizome growing tip. Any genotypes that died after 2 weeks were determined to have died from transplant shock and were replaced. No other plants were replaced after 2 weeks.

The experiment was conducted at a wetland site that had pre-existing stands of **P. arundinacea** (Biological Research Complex (BRC) Burlington, Vermont, (44°27'N, 73°11’W)).

The seven genotypes were collected from one of three established populations of **P. arundinacea** in Vermont (Shelburne Bay (44°24'N, 73°14’W), Gavin Hill (44°35’N, 73°08’W) and Ethan Allen Homestead (44°30’N, 73°14’W)). We used allozymes to identify seven unique genotypes (see Lavergne and Molofsky 2007) and vegetatively propagated all genotypes in the greenhouse prior to being planted at the field site. Clonal replicates of each genotype were generated by cutting stems from existing pots and adding rooting hormone to each node along the length of the stem. Stems were then laid in soil-filled flats and covered with soil. New plants were produced from each node within 6–8 weeks under saturated soil conditions and simulated spring temperatures (22–26°C day/16–20°C night with 12 h days).

All plots were monitored on a bi-weekly basis for both the 2006 and 2007 field seasons (July–September 2006 and May–September 2007) for survivorship, tiller number and the growth response traits of stem height and leaf number. On 4 September 2007, the above- and belowground biomass for all target and neighbor plants was harvested from each plot. Belowground biomass was washed and above- and belowground biomass was dried at 60°C for 48 h before being weighed.

**Mycorrhizae experiment**

If mycorrhizal associations among like genotypes is the mechanism responsible for positive frequency dependence, than genotypes must have genotype-specific responses to particular mycorrhizal strains (Ronsheim 1996). Here, we test the hypothesis that hyphal networks amongst like genotypes will generate a positive feedback. **P. arundinacea** can be highly mycorrhizal (up to 90% colonization), and therefore the presence of mycorrhizal fungi is a plausible mechanism affecting frequency dependence. To test for genotype specific mycorrhizal associations, we conducted a greenhouse experiment where we planted target genotypes with and without genotype neighbors in both mycorrhizal and non-mycorrhizal (fungicide-treated) soil. Target genotypes were planted in similar arrays to our frequency dependence field experiment, with either three same genotype neighbors, three different neighbor genotypes or no neighbors. Genotypes were collected from the
same three populations in Vermont as our frequency-dependent experiment and were grown under greenhouse conditions for multiple generations to remove any maternal effects prior to the establishment of the experiment. The experiment consisted of three neighbor treatments, two soil treatments, and four genotypes all replicated six times for a total of 144 pots. Soil for the experiment was collected from the field site where the frequency dependence experiment was planted in August 2008 and mixed 1:1 with perlite to improve soil aeration. Prior to soil removal, existing *P. arundinacea* plants were removed and roots were cleared and stained to confirm the presence of mycorrhizal fungi at our experimental site. We also collected three composite soil samples that were analyzed by the University of Vermont soil testing laboratory for available soil phosphate.

Plant arrays were established in 1-gallon (c. 4.5-L) pots, and humidity and other abiotic factors were regulated and maintained constant throughout the entire experiment. The temperature of the greenhouse was maintained at 23.8°C day/15.5°C night. We eliminated mycorrhizae by applying the fungicide Topsin-M in solution (70% a.i.; Cerexagri, Philadelphia, PA, USA) at a rate of 50 mg (active ingredient) kg⁻¹ soil (dry mass). Topsin-M was added as 200-mL aliquots to each pot at the beginning of the experiment and re-applied every 3 weeks. Topsin M: thiofanate-methyl: dimethyl [1,2-(phenylene)-bis(iminocarbonothioyl)] bis (carbamate) is a fungicide with a similar mechanism for fungal suppression as the now banned fungicide, benomyl (methyl 1-[butylcarbamoyl]-2-benzimidazole carbamate), and significantly reduces AMF colonization (Wilson and Williamson 2008).

Genotypes were planted in their treatments on 29 July 2008 and we measured both the target and neighbor plants for the traits of stem height, leaf number and tiller number every 3 weeks. We harvested the experiment 10 October 2008 and separated above- and belowground biomass for each individual genotype in each pot. *P. arundinacea* has distinctive rhizomes making individual genotypes easy to distinguish.

**Statistical analysis**

**Bayesian model**

We analyzed our data using Bayesian ANOVAs which are sometimes called Bayesian hierarchical linear random-effects models (Gelman and Hill 2007; Qian and Shen 2007). We chose this approach because it allows us to use a single comprehensive analytical framework for both our normal and non-normal data. These kinds of models can be built with a variety of methods (penalized quasi-likelihood, Laplace approximation) but the Bayesian Markov-Chain Monte-Carlo (MCMC) method is one of the most flexible (Bolker 2008). We used the statistical package R (R Development Core Team) in conjunction with WinBUGS (Gilsle et al. 1994) to fit the following generalized model.

\[
Y_{ijt} \sim N(\mu_{ijt}, \sigma^2)
\]

\[
\mu_{ijt} = \beta_0 + \beta_{ijt} + \beta_{2i} + \beta_{3j} + \beta_{4t}
\]

(1)

In the above generalized formulation, \(\beta_0\) is the grand mean, \(\beta_1\) is the effect of neighbor treatment \(j\), \(\beta_2\) is the effect of study year \(k\), \(\beta_3\) is the effect of genotype \(l\) and \(\beta_4\) is a term for the interaction between genotype \(l\) and neighbor treatment \(j\). Each effect term is then modeled as a random normal variable with a mean of 0 and variance \(\sigma^2\)

\[
\beta_{1-4} \sim N(0, \sigma_{\beta_{1-4}}^2).
\]

(2)

Variance between groups is the estimated variance term in Eq. 2. A second measure of between group variance is known as finite population variance \(s^2\), which is the standard deviation of each \(\beta_{1-4}\). When the number of groups is large, \(s^2\) and \(\sigma^2\) will converge, otherwise \(s^2\) is a more accurate estimate of between group variability (Gelman and Hill 2007). Each of the three response variables were fit to the full model (Eq. 1) with some changes made to accommodate the dataset. Mean stem height was fit to the full model (Eq. 1), while aboveground biomass measurements were fit to a model without study year, since measurements were only made in 2007 when the experiment was harvested.

\[
Y_{ijt} \sim \text{LogNormal}(\mu_{ijt}, \sigma^2)
\]

\[
\mu_{ijt} = \beta_0 + \beta_{ij} + \beta_{2i} + \beta_{3j} + \beta_{4t}
\]

(3)

Tiller number on final harvest was fit to a model that takes into account that tiller number is a Poisson-distributed count variable.

\[
Y_{ijt} \sim \text{Pois}(\mu_{ijt})
\]

\[
\mu_{ijt} = \beta_0 + \beta_{ij} + \beta_{2i} + \beta_{3j} + \beta_{4t}
\]

(4)

Each model was fit in WinBUGS called from R using R2WinBUGS using 3 chains and 50,000 iterations with a 10,000 iteration burn-in period (Sturtz et al. 2005). The significance of an effect was determined by whether a 95% credible interval bounded zero; if an effect bounded zero it was considered to be not significant.

**Frequency dependence for individual genotypes**

We quantified the extent of growth differences for our seven genotypes under both our same and different neighbor treatments. These metrics were calculated using the same three growth measures measured in our Bayesian model: mean stem height, aboveground biomass and final tiller number at harvest. We standardized growth (stem and
tiller) and biomass measures for treatments by subtracting the mean of the target individuals for genotype \( i \) of the same neighbor treatment (\( T_{ia} \)) from the mean of the target individuals for genotype \( i \) from the different neighbor treatment (\( T_{id} \)) and dividing by the mean of the target individual same neighbor treatment (Eq. 5).

\[
E_i = \frac{(T_{ia} - T_{id})}{T_{ia}}
\]

(5)

where, \( E_i \) represents the extent of the growth differences between the two neighbor treatments for genotype \( i \). Here, positive values indicate the extent of positive frequency dependence and negative values indicate the extent of negative frequency dependence. All \( E_i \) estimates were calculated with a permutation test with 10,000 replicates in R to calculate \( P \) values.

**Mycorrhizae experiment**

We analyzed the mycorrhizae data using the Bayesian ANOVA models described above, except instead of a year factor, \( k \) was the fungicide treatment. Stem height, leaf number, tiller number and biomass measures were our response variables and genotype, neighbor identity and soil type were treated as fixed effects. We also modeled interactions between genotype and fungicide treatment and between neighbor treatment and fungicide treatment. All response variables were modeled with the same error distributions as in the preceding models (e.g., leaf with a Poisson distribution) except for tiller number. We normally would model tiller number as a Poisson variable, but because of poor convergence we modeled it as a normal variable after a square root transformation. We again used WinBUGS and R2WinBUGS with 50,000 replicates with a 10,000 replicate burn-in and three chains. Significance was again assessed by whether or not 95% credible intervals bounded 0 for a given treatment.

**Results**

For all growth responses measured, the no neighbor treatment had consistently greater performance than the neighbor treatments indicating that plants in arrays experienced competition when grown with neighbors; however, this result was only statistically significant for tiller number (Fig. 1c). Comparing only the neighbor treatments, we found that the total plot biomass (including both target and neighbor plants) was significantly greater for same neighbor treatments than different neighbor treatments (Fig. 2). Survivorship of target plants was also greater under the same neighbor treatments than the no neighbor treatments; however, this effect was not statistically significant. For target plants, same and different neighbor treatments showed no neighbor treatment or genotype effect for the traits of leaf number, belowground biomass and total biomass. Here, we focus on the fitness response (tiller number)

![Fig. 1](image-url)  
**Fig. 1** Estimates of the effect size of neighbor treatment for *Phalaris arundinacea* on (a) stem height, (b) aboveground biomass and (c) tiller number. Same refers to the same neighbor treatment, Different to the different neighbor treatment and None to the no neighbor treatment. Each estimate is the effect relative to the grand mean of the trait. Thin lines represent 95% credible intervals and the center point is the median of the posterior distribution.
Each median line had different treatment effects and none to the no neighbor treatment. Each estimate is the effect relative to the grand mean of the trait. Thin lines represent 95% credible intervals and the center point is the median of the posterior distribution and growth responses (stem height and aboveground biomass) that showed significant genotype and neighbor treatment effects.

Fitness response

The largest variance component for tiller number (excluding the residual variance) was neighbor treatment (Fig. 3c). The presence of neighbors increased competition such that genotypes performed best with no neighbors and there was no difference in performance between same and different neighbor treatments (Fig. 1c). Tiller number was the only response variable measured that showed a significant effect of genotype (Fig. 3c).

Growth responses

Target plants surrounded by different genotype neighbors had smaller mean stem height, and smaller aboveground biomass than target plants surrounded by same genotype neighbors (Fig. 1a, b). Interestingly, for both stem height and aboveground biomass, there was no significant difference in performance between the same neighbor and no neighbor treatments (Fig. 1a, b) indicating that the presence of same genotype neighbors promoted positive frequency-dependent interactions that may have overcome the negative effects of competition. Individual genotypes differed in performance under neighbor treatments; however, the effect of neighbor treatment was so large that any effect of genotype was not significant, and for stem height, the inclusion of study year explained more variance than genotype (Fig. 3a).

Aboveground biomass had the greatest neighbor treatment × genotype interaction indicating that different

Fig. 2 Estimates of the effect size of neighbor treatment on the total biomass produced by each plot (including target plants and neighbors). Same refers to the same neighbor treatment, Different to the different neighbor treatment and None to the no neighbor treatment. Each estimate is the effect relative to the grand mean of the trait. Thin lines represent 95% credible intervals and the center point is the median of the posterior distribution.

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genotypes responded differently to neighbor frequency. Five genotypes performed better when surrounded by the same neighbor genotypes while two genotypes performed worse (Fig. 4b, inset). Stem height had six genotypes performing better when surrounded with like neighbors (Fig. 4a, inset). As stem height and aboveground biomass were the two traits to show differences in performance between neighbor treatments, and for many genotypes the standard error was large, we examined which genotype effects were significant by quantifying the extent of frequency dependence for individual genotypes.

Frequency-dependent growth for individual genotypes

To exclude the possibility that some genotypes were simply competitively dominant over others, we compared growth responses of genotypes when grown alone and

| Table 1 Frequency-dependent metrics for *P. arundinacea* growth measures for seven unique genotypes using a permutation test with 10,000 replicates |
|-------------|--------|-----------------|--------|
| Stem height | *P* value | Aboveground biomass | *P* value |
| GH10       | 0.05   | -0.32           | 0.63   |
| GH11       | 0.42   | 0.06            | 0.85   |
| GH30       | 0.5    | 0.89            | 0.04   |
| NEA11      | 0.63   | 0.74            | 0.26   |
| NEA24      | 0.61   | 0.86            | 0.02   |
| SB12       | 0.38   | -0.43           | 0.54   |
| SB3        | -0.11  | 0.73            | 0.26   |

Positive effect sizes mean that plants performed better when planted with same genotype neighbors, and negative effect sizes mean that plants performed better when planted with different genotype neighbors. Values in bold are statistically significant (*P* ≤ 0.05) found that there were no significant differences in growth among our seven genotypes. We then examined the interaction effect of the neighbor treatment × genotype interaction by calculating frequency metrics for all genotypes for the growth responses of stem height and aboveground biomass.

In neighbor treatments, genotypes showed a range of both positive and negative frequency-dependent interactions (Table 1); however, all genotypes that exhibited a negative frequency-dependent response were not significant. Two genotypes, NEA 24 and GH 30, had significant positive frequency-dependent interactions for both stem height and aboveground biomass measures. In total, three genotypes showed significant evidence for positive frequency-dependent interactions for stem height (GH30, NEA11 and NEA24) and three genotypes showed evidence for positive frequency-dependent interactions for aboveground biomass (GH11, GH30 and NEA 24) (Table 1; Fig. 4).

Testing the mechanism driving positive frequency dependence

Plants had greater stem height and aboveground biomass when planted in the fungicide-treated soil (Fig. 5). Stem height had a significant interaction with the fungicide treatment because plants in the no neighbor treatment had a greater effect of the fungicide treatment than neighbor treatments and there was no significant difference between the same and different neighbor treatments. Neighbor treatment had a significant effect for the traits of stem height, tiller number and aboveground biomass. All growth measures were significantly greater under the no neighbor treatment and we found no significant difference between the same and different neighbor treatments. The particular genotype used had a significant effect for tiller number; however, we found no significant interactions (Fig. 5).
Analysis of soil samples taken from the site of soil collection found that the mean amount of soil phosphorus was 5.65 ppm which is considered optimum in Vermont.

Discussion

Invasion potential may be increased if plants exhibit positive feedbacks (Klironomos 2002; Bever 2003). Interestingly, here we show that at the genotype level, positive frequency dependence increases invasive plant performance for several, but not all, growth measures. Stem height and aboveground biomass both showed evidence for intraspecific positive frequency dependence. For P. arundinacea, positive frequency-dependent interactions that result in taller stem heights may decrease time to seed set and increase seed production through greater seed recruitment. There appears to be a threshold stem height of approximately 30 cm before plants will set seed (Collins, personal observation). In addition, same neighbor and no neighbor treatments had no significant difference in performance for stem height and aboveground biomass. Thus, the presence of same genotype neighbors did not significantly reduce plant performance and may therefore increase invasive spread, particularly for newly invaded sites. Plants grown with same genotype neighbors may have less belowground competition (e.g., sharing resources underground) and may therefore allocate more energy to aboveground biomass and stem growth.

Positive frequency dependence may also help invasive species overcome potential Allee effects (Allee 1931) that can limit invasive spread (Elam et al. 2007). An Allee effect occurs when an individual’s performance increases with population size or density (Stephens et al. 1999) and can lead to decreased establishment, longer lag times and slower rates of spread (Taylor and Hastings 2005). For P. arundinacea, increased positive frequency-dependent interactions for growth measures such as aboveground biomass may promote the formation of dense monocultures leading to the exclusion of native plants even when propagule pressure is low. We found that genotypes produced greater biomass when planted with same genotype neighbors. Thus, the lack of genotypic variation at new sites of invasion may not limit invasion success. Finding increased productivity in monoculture is in contrast to other studies that have found greater overall productivity in genetically diverse plots (Johnson et al. 2006; Crutsinger et al. 2006). Interestingly, previous studies examining the effect of genotypic diversity have primarily used native species, and we suggest that positive frequency-dependent interactions among invasive genotypes may facilitate invasion into new environments.
We also found considerable variation in how different growth measures respond to changing the frequency neighborhood. Although most genotypes exhibited greater performance when grown with like neighbors, the extent of positive frequency-dependent interactions varied among genotypes and the growth measures examined. Furthermore, three growth measures (leaf number, belowground biomass and total biomass) showed no evidence for any frequency-dependent interactions. Given the relatively small number of genotypes used in our experiment and the large variability in response, this suggests that even at the invasive patch level there is a considerable range of responses to neighbor identity. A range of positive and negative frequency-dependent growth responses within a patch could ultimately impact established patch structure and influence local genotypic diversity (Eppsteiner et al. 2006). Negative frequency-dependent interactions amongst genotypes could promote genotypic diversity within a patch while the presence of genotypes that have positive frequency-dependent interactions could result in the formation of stable genotype clusters (Molofsky et al. 2001). Determining the extent of negative or positive intraspecific frequency-dependent interactions in established invasive populations will require a more rigorous sampling within a single patch as well as the determination of which specific plant traits are under frequency-dependent selection.

To our knowledge, no other study has quantified the extent of performance differences due to frequency-dependent interactions for individual invasive genotypes. What sets our work apart is that, rather than finding no frequency-dependent effect (Bennington and Stratton 1998) or pooling the frequency-dependent effects of all individual genotypes (Antonovics and Ellstrand 1984), we successfully quantify the amount of positive and negative frequency-dependent interactions for different growth measures for each individual genotype used in our study. As an increasing amount of work shows that genotypic diversity greatly influences community structure (Neuhuser et al. 2003; Whitham et al. 2003), quantitative studies such as ours will help to untangle how feedbacks impact genotype coexistence and may in turn clarify patterns we see at higher trophic levels. For example, Booth and Grime (2003) found that increasing genotypic variation of a long-term grassland experiment reduced the rate at which species diversity declined. A follow-up study (Whitlock et al. 2007), examining the role of genetic variation for shaping community composition, found that patterns of abundance in these plant communities could be predicted from knowing the genotypic composition of their component populations. Therefore, knowing how genotypes interact within established invasive patches and how their interaction influences performance could help predict their potential to outcompete native species and for future invasion risk.

We document cases of significant positive frequency-dependent interactions for certain invasive genotypes; however, what remains unclear is what mechanisms are driving these increases in plant performance. Empirical studies have proposed several potential mechanisms leading to positive frequency dependence including predation of rare species (Futuyma and Wasserman 1980), host specificity (Bever 1999) and shared mycorrhizal associations (Ronsheim and Anderson 2001). We predicted that mycorrhizal associations may have affected our positive frequency-dependent result; however, we demonstrated that mycorrhizal associations were likely not the main mechanism affecting positive frequency dependence. Despite the major limitation of our study, where Topsin-M appears to be killing off pathogens in the soil as well as mycorrhizae, we also found that available soil phosphate was high at the site where our field experiment was planted and where soil for our greenhouse experiment was collected. As mycorrhizae can increase the availability of immobile nutrients, especially soil phosphorus (Bolan 1991), and the level of available phosphorus is considered non-limiting at our site, we predict that another mechanism is likely at play. We suggest that future work should explore alternative mechanisms such as whether plant roots are able to recognize self and non-self (Dudley and File 2007; Mahall and Callaway 1996; Murphy and Dudley 2009).

In summary, we show that invasive genotypes differ in their direction and magnitude of frequency-dependent interactions for different growth measures, and for two key traits we find evidence for positive frequency dependence. For those genotypes that show evidence for positive frequency-dependent interactions, we also eliminate mycorrhizal associations as a factor affecting frequency dependence. Furthermore, we demonstrate that growth measures show differential response to neighbor identity, with some growth measures showing no difference in response regardless of the small-scale frequency neighborhood. We suggest that future work should try to understand what characteristics, both genetic and phenotypic, are associated with positive and negative frequency dependence. In this way, understanding scall-scale genotypic patch structure could better inform the potential for invasion risk.

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