Effects of native bryophytes on exotic grass invasion across an environmental gradient

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Abstract. Understanding the role of native biodiversity in controlling exotic species invasion is a critical goal in ecology. In terrestrial plant communities, most research has focused on the effects of native vascular plants on invasion by exotic vascular plants. However, in many ecosystems, native bryophytes and other non-vascular plants are common and can affect the establishment, survival, and growth of vascular plants. A more complete picture of how native biodiversity affects exotic plant invasion demands that more studies measure the effects of native bryophytes on exotic vascular plants. Moreover, there is growing realization that the effects of native species on invaders can range from negative to positive and that a complete picture of interactions between native and exotic plants requires measuring interactions in multiple environments. We used both observational and experimental studies to quantify the effects of native bryophytes on vascular plants along a 200-m environmental gradient in a coastal dune in northern California. We found a positive association between vascular plants and bryophytes across the environmental gradient. Our experiments with two exotic annual grass species showed the effects of bryophytes to be species-specific and to vary with environmental context. Bryophytes facilitated the survival of one exotic grass species, Vulpia bromoides, at both ends of the environmental gradient. Bryophytes reduced the survival and inflorescence production of the other exotic grass, Bromus diandrus, at one end of the environmental gradient and had no effect at the other end. Our findings provide a test of the effects of native bryophytes on exotic vascular plant invasion and show that these effects can vary dramatically even across local environmental gradients.

Key words: biological soil crusts; bryophytes; coastal dune; exotic species; facilitation; stress-gradient hypothesis.

INTRODUCTION

Biological communities throughout the world are increasingly invaded by a wide range of non-native species, and it is critical to understand the factors that mediate the success of these taxa in their novel environments (Kennedy et al. 2002). The environmental filtering model of invasion contrasts physical conditions, such as climate and soil type that might restrict exotic invasion, with the effects of biotic interactions such as competition, predation, and mutualism (Kennedy et al. 2002, Kraft et al. 2014). Exotic species are able to invade and thrive when they can pass through abiotic filters and cope with biotic interactions, which are usually assumed to be negative (Rejmánek 1996). Despite considerable evidence that competition can limit exotic plant invasion (Levine 2000, Kennedy et al. 2002), there are also a growing number of examples showing that exotic plants can be facilitated by native species (Badano et al. 2007, Griffith 2010, Cushman et al. 2011, Kleinhesselink et al. 2014). The stress-gradient hypothesis (SGH)
provides a unifying conceptual framework that may help explain how environmental context determines whether native species compete with or facilitate exotic invasion (Bruno et al. 2003). The SGH predicts that the effects of native species on invading exotic species will change across environmental gradients; specifically that effects will be less negative and more positive in more stressful environments (Bertness and Callaway 1994, Bruno et al. 2003, Badano et al. 2007).

Studies examining the effects of competition and facilitation on invasive exotic plants have tended to focus on interactions between vascular plants (Kennedy et al. 2002, Badano et al. 2007). However, exotic plants commonly interact with a much more diverse array of species in their novel environments. For example, deserts, coastal dunes, forest understories, and arctic environments are often colonized by an abundance of non-vascular plants such as bryophytes and lichens (Belnap et al. 2001). In these environments, invading exotic vascular plants is likely to have significant interactions with native non-vascular plants (Deines et al. 2007, Langhans et al. 2009). Despite their small stature, non-vascular plants can affect germination conditions for seeds and the availability of nutrients and water resources in the soil (Serpe et al. 2006, Langhans et al. 2009). In some systems, bryophytes and biological soil crusts more generally (fungi, lichens, cyanobacteria, bryophytes, and algae) have been found to play a role in resisting exotic species invasion, especially invasion by exotic annual grasses (Morgan 2006, Serpe et al. 2006, Deines et al. 2007, Hernandez and Sandquist 2011). Studying interactions between native bryophytes and exotic vascular plants along environmental gradients would provide a novel test of the idea that these interactions should change depending on environmental context.

While some studies have found that bryophytes can facilitate vascular plant germination, survival, and growth (Rayburn et al. 2012), it is unknown how much the effects of native bryophytes on exotic plants can vary across local environmental gradients. In this study, we investigate the effects of native bryophytes on the local distribution of vascular plants across a gradient in environmental conditions and test whether bryophytes affect the germination, survival, growth, and reproductive output of two exotic annual grasses. We test these predictions: (1) The association between vascular plants and moss patches will change systematically across an environmental gradient; (2) the effect of moss on exotic annual grass performance will change depending on position along an environmental gradient.

**Study System**

We conducted our study in a coastal dune system at Bodega Head in Sonoma County, California (38°19′N, 123°3′W). The area has a Mediterranean climate, receiving most of its precipitation between fall and early spring (Barbour et al. 1973). The soil is composed almost entirely of fine to coarse sands low in nutrients and organic matter (McNeil and Cushman 2005, Lortie and Cushman 2007, Cushman et al. 2010, Kleinhesselink et al. 2014). The current study was conducted between two large dune ridges that run in the direction of the prevailing winds (Appendix S1: Fig. S1). This site spans a 220-m gradient in soil conditions and wind speed that creates a large gradient in plant size and species composition, which we refer to as an environmental gradient. The details of this gradient have been previously described by Lortie and Cushman (2007) and Kleinhesselink et al. (2014). As one moves from the southeast to the northwest along the gradient, soil nitrate pools decrease by over 90%, soil water field capacity decreases by 40%, sand particle size doubles, and average surface wind speeds increase by roughly 500% (Lortie and Cushman 2007). These abiotic changes are accompanied by a decrease in average plant height and size: The height of the dominant shrub in this system decreases toward the NW along the gradient, and the average aboveground biomass of individual plants declines by a factor of 10 from the SE to the NW end of the gradient. *Bromus diandrus* (Poaceae; hereafter referred to as a *Bromus*) and two species of *Vulpia* (*V. bromoides* and *V. myuros*) are common exotic annual grass species at this site and throughout coastal California. We lumped the two *Vulpia* species into one group in our study because it was difficult to identify them to species in the field. A previous study showed that *Bromus* cover decreased across the environmental gradient, whereas *Vulpia* cover increased (Lortie and Cushman 2007).
Bryophytes and lichens constitute an important part of the ground cover at this site (Danin et al. 1998). The most abundant bryophyte along the environmental gradient was *Syntrichia ruralis* (Pottiaceae), which is a large drought-tolerant moss common in seasonally dry environments across California (Malcolm et al. 2009). Other bryophytes common at this site were *Homalothecium arenarium* (Brachythecaceae) and *Didymodon vinealis* (Pottiaceae).

**Methods**

**Vascular plant association with bryophytes across the gradient**

We established 23 20-m transects spaced approximately 10 m apart along the 220-m gradient documented by Lortie and Cushman (2007) and Kleinhesselink et al. (2014). Each transect ran perpendicular to the dune gradient (Appendix S1: Fig. S1). The most southeasterly transect occurred at the sheltered end with low wind speed finer textured sand, and the most northwesterly transect occurred at the more exposed end of the gradient with higher wind speeds and coarser textured sand. For the remainder of the paper, we refer to position along the gradient in meters away from the more sheltered southeast end of the gradient and refer to these sides of the gradient as the SE and NW ends of the gradient.

In order to determine cover of native mosses across this study site, and their association with the focal species of annual grass and other vascular plants, we recorded the cover of shrub, bare sand, or moss patches on each of the transects described above. To estimate the cover of mosses, we used point-intercept sampling at 25 randomly chosen points along each transect, recording whether points fell on a moss patch, bare sand, or within a shrub. All moss species were lumped together and recorded as moss. We then recorded the species identity of any vascular plant rooted within 1 cm of the point.

**Effects of bryophytes on Bromus and Vulpia**

We established a field experiment in the winter of 2009/2010 to assess the effect of moss on the germination, survival, growth, and reproduction of both exotic grass species. For each species, we established 18 separate blocks centered on large mats (>15 cm in minimum width) of the native moss *Syntrichia ruralis*. Each block was positioned away from shrubs and other large perennial plants. Nine blocks were located on the sheltered SE end of the gradient, and nine were located at the more exposed NW end of the gradient. Each block consisted of three separate 5 x 10 cm patches (Appendix S1: Fig. S2). Two patches were positioned on naturally occurring mats of moss. We left one as an un-manipulated patch (moss covered), and we removed the moss on the other by pulling the moss off the soil surface (moss removed). The third patch in each block was established on naturally moss-free bare sand as a control. Within each patch, we planted five seeds of the either *Vulpia* or *Bromus*. We planted only five seeds per patch (roughly spaced 2 cm apart) in order to keep seedling density low and prevent any self-thinning from occurring among the planted individuals. To prevent the seeds from blowing out of the patches and to help re-locate the seeds, we glued each seed by its own to a small wire and inserted the wire into the sand. We then pressed each seed into the sand or the moss surface of the patches. Seeds of both species were collected from the field site in the fall of 2009. The *Bromus* seeds were planted in late December 2009, and the *Vulpia* seeds were planted a few weeks later in January of 2010 in separate blocks.

Starting in January 2010, we visited the experimental patches roughly once every two weeks. Because seeds were glued to wires in each patch, we were able to easily distinguish between plants germinating from the seeds we planted from plants emerging from the natural seed bank. We weeded out all other seedlings within the experimental patches to eliminate the effect of competition between our focal plants and other vascular plants. In May 2010, the surviving plants produced inflorescences and began to senesce. At this time, we counted the total number of surviving plants in each patch, harvested their aboveground biomass, and counted the total number of inflorescences. Total aboveground biomass from all focal plants surviving in each patch was dried at 60°C for 48 h and weighed to the nearest milligram.

**Statistical analyses**

We used logistic regression to determine whether vascular plants were associated with
moss patches and whether the association between vascular plants and moss changed across the environmental gradient. For this analysis, we only included points falling outside of shrubs. We modeled the probability of a vascular plant occurring at each sampling point as a function of gradient position (meters away from the SE end of the gradient), moss presence (either moss patch or bare sand), and the interaction between these two factors. When residual deviance was greater than residual degrees of freedom, we used a quasi-binomial model as recommended by Crawley (2007). We fit separate models for all vascular plant species together, for exotic species, for native species, and for the target exotic annual grasses, Vulpia and Bromus, together. We tested significance of the gradient effect, the micro-habitat effect (moss covered or bare sand), and their interaction by comparing model deviance with F-tests in R (R Core Team 2015).

We used logistic regression to model how position on the environmental gradient (SE vs. NW end of the gradient) and treatment (moss removed, moss present, or bare sand) and their interaction affected the final number of surviving Bromus and Vulpia plants in each experimental patch. Because we did not track individual germination and survival, our analysis focuses on the expected probability of one of the five seeds planted in each patch transitioning to an adult plant at the end of the growing season. The model included treatment, gradient position, and their interaction as fixed effects and the 18 experimental blocks as random effects. We fit the model with a generalized linear mixed effects model with a logit link and binomial errors in the lme4 package in R (Bates et al. 2015). We analyzed the log average aboveground biomass of each species using the same model structure but within a linear model with normal errors. Similarly, we analyzed the number of inflorescences produced in each experimental patch using a generalized linear model with a log-link and quasi-Poisson errors. We used the number of surviving plants in each patch as an offset term to control for varying number of surviving plants in each patch. We fit the inflorescence data without the random block effects because models fit with the random effect failed to converge.

We compared model deviance with chi-square or F-tests to evaluate the significance of each of the fixed effects in the models. When we found a significant treatment or treatment × environmental gradient effect, we tested for significant pairwise differences between the treatment means. We adjusted for the multiple comparisons using the Šidák correction implemented with the emmeans package in R (Lenth and Hervé 2015).

RESULTS

Vascular plant associations with moss patches across the gradient

Moss cover was low in the more sheltered SE end of the gradient, peaked toward the middle of the gradient, and declined across the last 50 m of the gradient (Fig. 1). This pattern of moss cover was well described as a quadratic function of distance on the environmental gradient ($F_{2,20} = 26.7$, $R^2 = 0.70$, $P < 0.01$). We found 31 different species of vascular plants rooted at 189 of the 398 sampling points outside of shrubs. Chorizanthe cuspidata (Polygonaceae), a small native annual, was the most frequently encountered species and occurred at 60 sampling points. Lumping all vascular plant species together, we found they were rooted at 104 of 277 points without moss and 85 of 121 moss covered points. The frequency of vascular plants increased toward the NW end of the gradient ($F_{1,396} = 54.3$, $P < 0.01$; Fig. 2) and was greater within moss patches than uncolonized bare sand patches ($F_{1,395} = 17.8$, $P < 0.01$). However, we did not find a gradient position × micro-habitat interaction ($F_{1,394} = 0.70$, $P = 0.40$). Exotic and native species both increased in frequency toward the NW end of the gradient (exotics: $F_{1,396} = 5.0$, $P = 0.03$; natives: $F_{1,396} = 37.7$, $P < 0.01$; Appendix S1: Figs S3 and S4) and were both more frequent within moss patches than uncolonized bare sand patches ($F_{1,395} = 17.8$, $P < 0.01$). However, we did not find a significant treatment × environmental gradient effect, we tested for significant pairwise differences between the treatment means. We adjusted for the multiple comparisons using the Šidák correction implemented with the emmeans package in R (Lenth and Hervé 2015).
interaction ($F_{1, 394} = 1.5$, $P = 0.22$; Appendix S1: Fig. S5).

Effects of moss on Bromus

There was a significant treatment x gradient position interaction on Bromus survival ($\chi^2 = 24.8$, df = 2, $P < 0.01$); Bromus survival was greater in bare sand patches than in moss-covered patches at the more sheltered SE end of the gradient, but this difference disappeared in the more exposed NW end of the gradient (Fig. 3a). Moss treatment had a significant effect on Bromus final biomass ($\chi^2 = 7.44$, df = 2, $P = 0.02$; Fig. 3c); however, this effect was due to biomass being higher in bare sand patches than in patches with moss removed (Fig. 3c). There was no significant treatment x gradient interaction effect on final Bromus biomass ($\chi^2 = 4.27$, df = 2, $P = 0.12$) nor a main effect of gradient position ($\chi^2 = 1.63$, df = 1, $P = 0.19$). Bromus inflorescence production was affected by a treatment x gradient interaction ($F_{2,46} = 5.35$, $P = 0.01$): At the SE end of the gradient, plants in bare sand produced significantly more inflorescences than plants in either moss patches or moss-removed patches, whereas at the NW side of the gradient there were no differences between treatments (Fig. 3e).

Effects of moss on Vulpia

Moss significantly increased Vulpia survival ($\chi^2 = 32.97$, df = 2, $P < 0.01$; Fig. 3b), but this effect did not vary across the gradient ($\chi^2 = 0.58$, df = 2, $P = 0.75$). There was a trend toward higher Vulpia survival at the NW end of the gradient ($\chi^2 = 3.60$, df = 1, $P = 0.06$). Vulpia biomass was not significantly affected by moss treatment ($\chi^2 = 4.97$, df = 2, $P = 0.08$; Fig. 3d), gradient position ($\chi^2 = 0.21$, df = 1, $P = 0.65$), nor was there a treatment x gradient interaction ($\chi^2 = 1.18$, df = 2, $P = 0.55$). Similarly, Vulpia inflorescence production was not affected by moss treatment ($F_{2,40} = 0.43$, $P = 0.65$), gradient position ($F_{1,42} = 0.53$, $P = 0.47$), nor their interaction ($F_{2,38} = 0.18$, $P = 0.83$; Fig. 3f).

DISCUSSION

Our study demonstrates that native bryophytes have important effects on the occurrence of vascular plants in this system as well as the survival of two exotic annual grasses. However,
we found only limited support for our hypothesis that the strength and direction of these effects would change across the environmental gradient. The natural occurrence of vascular plants was significantly greater in moss patches than in bare sand but this positive association did not change significantly across the environmental gradient as we predicted (Fig. 2). This result held for both native and exotic species alike (Appendix S1: Figs. S3 and S4). In our field experiment, we found only a few cases where moss had a facilitative effect on exotic grass performance at one end of the environmental gradient but a competitive effect at the other end. In particular, the effects of moss varied along the environmental gradient for *Bromus* survival (Fig. 3a) and inflorescence production (Fig. 3e). In the more sheltered SE end of the gradient, moss reduced *Bromus* survival compared to both bare sand treatments (Fig. 3a). In contrast, in the more exposed NW end of the environmental gradient *Bromus* survival was greater (but not significantly so) in moss patches than in the moss-removed or the bare sand patches. This indicates that the effects of moss patches on exotic species invasion may change across the gradient depending on local environmental conditions. We also observed a gradient by treatment interaction effect on *Bromus* inflorescence production but this was less supportive of our hypothesis (Fig. 3e). At the SE end of the environmental gradient, *Bromus* produced fewer inflorescences in both the moss-covered and moss-removed patches compared to the bare sand patches. However, the differences between any pair of treatment levels disappeared in the more exposed NW end of the gradient (Fig. 3e). The effects of moss on *Vulpia* did not vary along the environmental gradient (Fig. 3b, d, f). Nevertheless, we did find that moss-covered patches consistently gave *Vulpia* seeds the highest probability of germinating and surviving into adult plants (Fig. 3b). This indicates that moss-covered patches may be key micro-habitats supporting *Vulpia* population growth and invasion in this environment.

Our work shows the potential for native mosses to influence the success of invasive vascular plants in this environment. Mosses and other components of biological soil crusts are often found to have neutral to negative effects on
vascular plant germination (Zamfir 2000, Serpe et al. 2006, Jeschke and Kiehl 2008, Drake et al. 2018), while at the same time they can increase the growth and survival of established plants (Pendelton et al. 2003, Langhans et al. 2009, Ferrenberg et al. 2018). Our experiment showed that moss mats had positive effects on the germination and survival of Vulpia across the environmental gradient. We hypothesize that the beneficial effect of moss on seedling germination and survival may be due to the ability of moss to retain moisture and organic matter in the upper soil layers (Sand-Jensen and Hammer 2012).

Our experiment showed that the effect of moss on germination and survival depended on the annual grass species involved. The germination and survival of Vulpia were facilitated by moss across the gradient, but moss had a negative effect on Bromus at the SW end of the gradient (Fig. 3a). Other studies have also shown that the effects of moss on seedling performance depended greatly on the species of vascular plant (Zamfir 2000, Serpe et al. 2006). Seed size may be an important plant trait that controls how plants respond to bryophyte mats and Bromus seeds are over 10 times larger than Vulpia seeds (Royal Botanic Gardens, Kew 2019). In deep moss mats, small seeds can be at a disadvantage as they fall deep into the moss where it may be too dark to germinate or emerge (Zamfir 2000). On the other hand, large seeds run the risk of being stuck on the surface of a moss mat and not being able to absorb enough water to germinate—Serpe et al. (2006) showed that it was not until seeds were incorporated into the moss layer that they began to absorb moisture. One hypothesis to explain the species-specific difference observed in our study involves an interaction between moss mat density and seed size. In the more sheltered end of the gradient, moss patches may be thicker and denser—although we did not measure this. If this is the case, denser moss could have kept the larger Bromus seeds on the surface of the moss mat and prevented them from absorbing moisture. In contrast, the smaller Vulpia seeds may have fallen more deeply into the moss layer and had better access to moisture. More detailed observations of the thickness of moss mats across the gradient as well as time courses of seed water status would be needed to test this hypothesis.

We expected that the performance of the annual grasses in patches where moss was removed would be similar to the performance in naturally bare sand patches. However, in the more sheltered end of the environmental gradient Bromus survival and inflorescence production were significantly greater in bare sand patches than in patches where moss was removed, while the difference between moss-covered patches and moss-removed patches was not significant (Fig. 3a, e). Likewise, the biomass of Bromus plants in bare sand patches was significantly greater than the biomass of plants in moss-removed patches (Fig. 3c). The difference between Bromus performance in bare sand and moss-removed patches is notable because they are among the strongest effects in the experiment. These results indicate that the environment created by our removal treatment was somehow different from naturally bare sand. We speculate that this effect is either due to depletion of local soil nutrients by the removed moss or due to some residual allelopathic influence of moss in these patches (Michel et al. 2011). Another possibility is the presence of cryptic cyanobacterial or algal crust on what we assumed were naturally bare sand patches. These crusts may have exerted a positive effect on Bromus but would not necessarily be present in the patches where moss was experimentally removed.

We expected that the exotic annual grasses in this system would have their performance limited at the NW end of the environmental gradient where wind speeds are higher and the sand is coarser and lower in nitrogen content. Instead, we found that Bromus and Vulpia often performed as well or better at the NW end of the environmental gradient (Fig. 3). While the decrease in plant size and biomass of other species across the gradient is suggestive of a stress gradient (Lortie and Cushman 2007 and Kleinhesselink et al. 2014), this same gradient may not be stressful for the exotic species in this study. This result runs counter to the hypothesis that exotic species are limited from stressful environments within landscapes because they lack specialized adaptations needed to tolerate the local stresses (Harrison 1999). Instead, our finding supports the idea that stressful environments can sometimes be more easily invaded by exotic plants, perhaps because it offers opportunity to
escape competition from larger native competitors (MacDougall et al. 2006).

CONCLUSION

Native biodiversity plays a critical role in determining which exotic species can invade and where exotic species can invade. We demonstrate that native bryophytes can also influence the success of exotic vascular plants. Moreover, we found the effects of bryophytes on exotic annual grass establishment depended on environmental context and the vital rate being measured. We find evidence that bryophytes can either compete with or facilitate exotic vascular plants and suggest that the role of native bryophytes should more often be considered in conservation and restoration of native vegetation (Bowker 2007, Chiquoine et al. 2016).

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**Data Accessibility**

Data and R scripts to reproduce analyses available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.m27m0k8

**Supporting Information**

Additional Supporting Information may be found online at: http://onlinelibrary.wiley.com/doi/10.1002/ecs2.2769/full