INTRODUCTION

Organisms in natural ecosystems participate in a web of mutualistic and antagonistic species interactions. For plant species, interactions with herbivores, pathogens, mutualists, and competitors occur both aboveground and belowground, comprising two subsystems that feedback with each other to regulate community structure and ecosystem functioning (e.g., Bardgett, 2005; Van der Putten, Vet, Harvey, & Wäckers, 2001; Wardle, 2002). For example, the interaction between plants, their belowground microbial mutualists, and aboveground insect herbivores can affect the fitness and community dynamics of all three partners (reviewed in Gehring & Bennett, 2005; Van der Putten, Vet, Harvey, & Wäckers, 2001; Wardle, 2002).
Arbuscular mycorrhizal fungi (AMF) are belowground plant symbionts that colonize the cortical root cells of vascular plants. The association is common, found in an estimated 80% of vascular plant families, and is characterized by an exchange of soil nutrients and water accessed by AMF in exchange for plant photosynthate. AMF often improve plant establishment, survival, and growth and can also confer tolerance to high concentrations of salts and heavy metals (Hildebrandt, Regvar, & Bothe, 2007; Porcel, Aroca, & Ruiz-Lozano, 2012; Smith & Read, 2008). Plants inoculated with AMF often also show improved growth under water stress (Augé, 2001). Although this symbiosis is thought to be largely mutualistic in natural populations, the outcome of AMF–plant interactions is often context dependent (Hoeksema et al., 2010; Johnson, Graham, & Smith, 1997; Klironomos, 2003). A mutualistic relationship is expected to occur in environments with low soil nutrients, limited water availability, and/or high light levels (Johnson, Rowland, Corkidi, Egerton-Warburton, & Allen, 2003; Johnson et al., 1997). However, under extremely arid conditions the symbiosis may fail because fungal hyphae are unable to grow and persist, and the plant conducts minimal photosynthesis to conserve water via stomatal closure (Dosskey, Boersma, & Linderman, 1991).

Because AMF interactions affect plant physiology in such diverse ways, it is no surprise that they can have complex, indirect influences on a plant’s interactions with other organisms. AMF have been found to affect interactions with a number of plant mutualists, including pollinators (Gange & Smith, 2005; Wolfe, Husband, & Klironomos, 2005) and rhizobia (Larimer, Clay, & Bever, 2014), as well as plant enemies, such as pathogens (Borowicz, 2001) and herbivores (Gehring & Whitham, 2002). Given the potentially large effects of both herbivory (Maron & Crone 2006) and mycorrhizal fungi (Koide & Dickie, 2002) on plant populations, the tripartite interaction of plants, AMF, and herbivores has received considerable attention. Previous experiments on plant–microbe–herbivore interactions have produced highly variable results (Gehring & Bennett, 2009), with the net outcome depending on factors such as herbivore feeding specialization (Koricheva, Gange, & Jones, 2009) and various abiotic stresses (Pineda, Dicke, Pieterse, & Pozo, 2013). Herbivory may increase with AMF because plants acquire more nutrients and grow more quickly resulting in more abundant and high-quality food for herbivores (Smith & Read, 2008; Vannette & Hunter, 2011). Alternatively, herbivory may decrease with AMF because of changes in plant defense, either via immune system “priming” due to fungal colonization of plant tissue (Jung, Martínez-Medina, Lopez-Raez, & Pozo, 2012), or increases in constitutive or inducible defense compounds (Gange and West 1994, Bennett & Bever, 2009). The magnitude of positive or negative microbial effects on plant resistance to herbivores may be amplified under abiotic stress because of cross-talk between plant signaling pathways induced by each stressor (Pineda et al., 2013).

Plant–AMF–herbivore interactions likely depend on light availability, soil nutrient content, and other environmental factors that influence the rates of resource exchange between plants and AMF and therefore plant tissue quality. Populations of plants may adapt to their local abiotic and/or biotic soil environments in part through their response to AMF colonization (Kaepler et al. 2000, Cavender & Knee, 2006; Johnson et al., 2010), which may in turn cause population variability in AMF-driven plant–herbivore interactions. Two studies have examined how plant genotypes differentially respond to AMF, and how this variable response translates into different effects on herbivores, both finding that plant genotype can significantly alter the outcome of this tripartite interaction (Bennett, Millar, Gedroics, & Karley, 2016; Garrido, Bennett, Fornoni, & Strauss, 2010). However, only one study (Rasmussen et al., 2017) has incorporated geographic variation in studies of plant–AMF–herbivore interactions; this study found variation among three populations of Plantago lanceolata in both response to AMF and subsequently, effects on herbivores in feeding trials. While some work has been done looking at plant–microbe–herbivore interactions for species in the process of range expansion (Engelkes et al., 2008), no studies that we know of have sampled plant populations from the center to edge of a species’ current geographic distribution where a strong environmental gradient might contribute to variation in the outcome of species interactions.

In this study, we examined multispecies interactions among plants, AMF, and herbivores using six populations of the annual plant Clarkia xantiana ssp. xantiana (hereafter, xantiana). Xantiana is colonized by AMF in the field and experiences a strong geographic gradient in the strength of its interactions with insect herbivores, especially grasshoppers. Damage by grasshoppers occurs principally on developing fruits and seed loss to herbivores can exceed 40% in some populations (D. Moeller, unpublished data). The severity of insect herbivory declines considerably from center to margin of xantiana’s geographic distribution, and resistance to herbivory within populations reflects this geographic trend, with higher herbivore resistance observed in central populations (D. Moeller, unpublished data). This variation in herbivory occurs across a marked gradient in abiotic factors, where precipitation becomes lower and less predictable toward the range edge (Eckhart et al., 2011). With the complex gradient in abiotic and biotic factors across xantiana’s range, the outcome of interactions between plants, microbial symbionts, and insect herbivores may also change across geographic space and among plant populations.

Here, we examine how AMF and plant source population interact to affect plant performance and the severity of herbivory under water stress using a greenhouse experiment. Specifically, we ask the following questions:

1. Does AMF inoculation affect xantiana growth and phenology under low water availability?
2. Does AMF inoculation alter herbivore preference for xantiana fruits?
3. Does this tripartite interaction vary among plant populations spanning the range of the subspecies?

2 | MATERIALS AND METHODS

2.1 | Natural history of Clarkia xantiana

Clarkia xantiana ssp. xantiana is a primarily outcrossing winter annual forb endemic to southern California. It occurs primarily on steep, sandy slopes in the southern Sierra Nevada at elevations of 500–1500 m (Eckhart & Geber, 1999; Lewis & Lewis, 1955). Seeds germinate in November–December, and plants mature in May–June. The populations used in this experiment are distributed along a strong environmental gradient from the center of xantiana’s distribution to its eastern range edge, with mean annual precipitation and temperature decreasing, and variability in precipitation increasing, as populations approach the eastern range limit (Eckhart et al. 2010, Eckhart et al., 2011). Soils in the center of xantiana’s range are mostly derived from igneous rock, with metasedimentary soils becoming increasingly frequent at the eastern range edge (Eckhart et al. 2010). One population used in this study, S22, occurs on soils derived from metasedimentary rock; all others occur on igneous soils.

Xantiana plants are attacked by multiple herbivores. Fatal mammalian herbivory becomes increasingly frequent toward the eastern range edge (J. Benning, unpublished data), while invertebrate herbivory is highest in central populations (D. Moeller, unpublished data). Defoliation by larvae of Hyles lineata (White-lined Sphinx Moth) occurs occasionally (personal observation), but the most common invertebrate herbivory is by grasshoppers (Melanoplus sp.). These generalist herbivores feed on immature fruits of xantiana by eating the immature, green pericarp and/or immature seeds; this fruit herbivory can have significant fitness consequences for both individuals and populations of xantiana, with some populations losing more than 40% of their fruits on average to grasshopper herbivory (D. Moeller, unpublished data).

Surveys in natural populations show that the subspecies is colonized by AMF in the field (J. Benning, pers. observation).

2.2 | Experimental design

We conducted a greenhouse experiment in St. Paul, Minnesota, USA manipulating AMF inoculation and source population of xantiana in a 2 × 6 full factorial design (n = 50 plants per treatment combination, 600 plants total). We applied each of two AMF treatments—AMF-inoculated (M+) and AMF-free (M−)—to half of the individuals from each population. In order to capture variation in xantiana’s AMF responses and herbivore resistance, we chose source populations that span from the center to edge of its range (Figure 1); the source populations used were (from center to edge): CF (109x, 35°00′94″E), Del (107x, 35°31′27″E), Key (57x, 36°28′64″E), S22 (22x, 36°90′76″E), C3 (21x, 36°91′07″E), and Golf (3x, 37°05′82″E). (Population labels with “x” are included here to correspond to earlier published work with xantiana, but we use the preceding labels throughout the manuscript for simplicity.) We used field-collected seeds from 10 maternal families within each population. All seeds were collected from the field during the summer of 2015.

Xantiana experiences low nitrogen availability in the field (field NO₃ mean ± SD: 4.26 ± 4.94 ppm, averaged across five of the six populations used in this experiment), and we prepared our soil mix (2:1 sand (Fischer Mining Co., MN, USA): growing mix (SunGro LC8; SunGro, MA, USA); this growing mix typically contains 50–140 ppm NO₃) to reflect the low nutrient charge of natural soils. The soil mixture was steam pasteurized for 2 hr at 80°C and then allowed to cool for at least 2 days before planting. We prepared 600 1 L pots (Stuewe & Sons D60 Deepots) for planting by submerging in 0.9% sodium hypochlorite (bleach) solution, and lining the pot bottoms with newspaper.

Arbuscular mycorrhizal fungi treatments were applied to the soil mixture prior to planting. Plants receiving the M+ treatment were planted into pots with a 25-ml layer of Micronized Endomycorrhizal Inoculant (BioOrganics Inc, New Hope, PA, USA) spread ca. 4 inches below the soil surface This commercial AMF inoculum contains a blend of nine endomycorrhizal species (Glomus aggregatum, G. etunicatum, G. clarum, G. deserticola, G. intraradices, G. mosseae, Gigaspora margarita, and Paraglomus brasiliannum) and has been used successfully to induce AMF colonization in a wide range of plant species (Babikova et al., 2014; Berg, Eaton, & Ayres, 2001; Cavender & Knee, 2006; Pischl & Barber, 2016; Wiseman & Wells, 2009). The non-mycorrhizal control treatment (M−) pots only contained 2:1 sand: growing mix. Two seeds were then sown on top of the soil.

Pots were placed in a cold room with 24-hr light (2–150 W bulbs) at 8°C for the first 6 days, then 12°C for six more days. While in the cold room, plants were misted daily. After 12 days in the cold room, and 3 days after the first germinants were observed, pots were moved to the greenhouse. Fourteen pots had no germination; for
those pots, pre-germinated seeds from the assigned population and maternal family were sown on top of the soil.

In the greenhouse, plants were maintained at ~25 to 27°C for the first month on a 14-hr/10-hr L/D cycle and then ~23 to 25°C for the remainder of the experiment on the same L/D cycle. For the first 6 weeks in the greenhouse, plants were watered regularly to ensure early survival. They then received four waterings 2 weeks apart, and then, watering was stopped altogether for the final 7 weeks of the experiment. This watering regime was designed to mimic the seasonal conditions experienced by *xantiana* in the field, as rainfall tapers off and eventually stops altogether toward the end of their growing season. Survival was very high, with 95%–99% of individuals (of 100) surviving for each population. Sample sizes for Key were lower (59) because field collections mistakenly included some individuals of *xantiana*’s sister subspecies, *C. xantiana* ssp. *parviflora*, as these are difficult to distinguish in the field after fruit set. Those *parviflora* individuals were excluded from analysis (this did not skew sample sizes between AMF treatments for Key: M+ *n* = 29; M− *n* = 30).

Once plants began flowering, we generated fruits by crossing random pairs of plants from the same population × AMF treatment, but from different maternal families. Two to six flowers per plant were crossed.

We measured two phenological traits—days to first flower and time spent flowering—and four growth allocation metrics—number of flowers produced, as a proxy for reproductive effort; number of seeds per fruit; average seed weight; and shoot dry biomass. Days to first flower were recorded as the number of days between planting and the day the first flower opened, and time spent flowering was the number of days between the day the first flower opened and the day the last flower senesced. All hand-pollinated fruits not used in herbivore trials were collected at maturity; for each of these fruits, the number of seeds per fruit was counted, and the mass of all seeds from a fruit was quantified. Individual seed weight was calculated by dividing total seed mass from a fruit by the number of seeds in that fruit. After plants had senesced, their stems were clipped at the soil level and desiccated in a drying oven at 55°C prior to measuring shoot dry biomass. All stems were collected over the course of 7 days.

### 2.3 | Herbivore choice trials

We conducted herbivore choice trials on a subset of fruits to assess whether AMF may confer herbivore resistance to host plants, and how those effects may vary among source populations. Our trials occurred only using fruits, which avoids the potentially confounding effects of fruit number, plant architecture, or plant size when using whole plants in herbivory trials. We used young adults of the generalist grasshopper, *Melanoplus sanguinipes* (USDA ARS Northern Plains Agricultural Research Laboratory), as the herbivore in our feeding trials. *Melanoplus devastator*, a close relative, is a common herbivore of *xantiana* in the field. It is unclear whether *M. devastator* and *M. sanguinipes* are different taxa because the main differentiating feature is geographic location, whereas other attributes do not appear to differ (Dingle & Mousseau, 1994; Orr, Porter, Mousseau, & Dingle, 1993). Grasshoppers were maintained in ventilated plastic tubes kept in growth chambers at 30°C, 16:8 L/D. They were fed a diet of organic romaine lettuce and Cheerios (General Mills, Minneapolis, MN) and were thus naïve to variation in *xantiana* tissue quality.

Trial arenas were constructed in plastic Ziploc containers (15.5 × 15.5 × 8.5 cm) with a square piece of medium-density fiberboard placed in the bottom. We included one fruit from each population × AMF treatment in every trial, for a total of 12 fruits per trial.

![Figure 2](image_url)
We drilled 12 holes equidistantly around the perimeter of a circle in the MDF board, and each fruit was pinned into a randomly assigned hole. The MDF board was then covered with sand to mimic field conditions, with the fruits sitting just above the level of the sand, and the tops of the Ziploc containers were replaced with fine mesh to allow air movement (Figure 2). Prior to trials, all fruits were sprayed with water to delay desiccation.

On consecutive days, we conducted two rounds of choice trials which included 20 and 12 trial arenas, respectively. One trial in round two was discarded due to the death of the grasshopper during the trial, resulting in a total of 31 choice trials. Two of the seven grasshopper tubes were randomly selected for each round, before which grasshoppers in those tubes were starved for 27.5 hr. Grasshoppers were then haphazardly chosen from the tubes for trial. One grasshopper was placed into each arena and allowed to feed on xantiana fruits for 17.5 hr. In addition, round one had two control trials, which received no grasshopper, and round two had one control trial. Trials were conducted at 32°C, 16:8 L/D.

We assessed the severity of herbivory using a categorical scoring system. Grasshoppers produce two distinct types of herbivory on xantiana fruits, which we called “scraping” and “chomping.” “Scraping” herbivory appears as rasping on the side of a fruit that may or may not breach the ovary wall; “chomping” is a shortening of the fruit from the tip toward the peduncle, and usually results in a greater fitness loss (from devouring of immature seeds) than “scraping”. Fruits used in herbivory trials were scored on a scale of 0–3 based on the type of herbivory they suffered. A fruit was scored "0" for no/trace herbivory; "1" for 10%–50% scraping; "2" for 50%–100% scraping; or "3" for chomping. A binary measure of herbivory was also used in analysis, for which herbivory scores 1–3 were condensed into the single factor "herbivory present," and 0 scores were "herbivory absent." Herbivory scoring was "blind"; that is, the herbivory scorer was unaware of which treatment combination each fruit came from while scoring.

### 2.4 | Statistical analyses

We used linear mixed models to test for the effects of plant population and AMF inoculation on plant fitness proxies (biomass and flower number), phenology (time to begin flowering and time spent flowering), and seed production (seeds per fruit and seed weight). For fruits that were hand pollinated, seeds per fruit was calculated for each plant as the mean number of seeds per fruit, with the number of analyzed fruits on a plant ranging from one to six. Some fruits that were hand pollinated did not fill any seeds, which most likely reflected unsuccessful hand pollination, rather than abortion due to a lack of resources. Consequently, we removed seedless fruits from seed analyses. Seed weight was calculated for each plant as the mean weight of a single seed. We included all plants with at least one fruit that produced two or more seeds (n = 423).

Treatment effects were examined using a type-II ANOVA using the Afex package in R (R Core Team 2017). Effect sizes were compared between the Kenward–Rogers and Satterthwaite approximations for denominator degrees of freedom; minimal differences were found between these methods, so Kenward–Rogers values are reported. Days to first flower, time spent flowering, dry biomass, number of flowers produced, number of seeds per fruit, seed weight, and seed weight per fruit were included separately as response variables. Population, AMF treatment, and population x AMF interaction were included as fixed effects, and maternal family (nested within population) was included as a random effect. Normality and homoscedasticity of residuals were verified visually. For models that showed variation in response among populations, Tukey’s post hoc test was employed to detect pairwise differences between populations using the Holm correction for multiple comparisons.

Although the inclusion of multiple maternal families was primarily to capture a representative sample of population genetic variation, we also explored the potential for differences in the direction of response to mycorrhizal fungi (positive versus negative) among maternal families. To do this, we performed t tests comparing biomass and reproductive effort of M+ and M− plants for each family. If the 95% CI of the difference between M+ and M− groups did not span zero, we interpreted the family as having a significantly positive or negative response.

We tested for an effect of population and AMF treatment on herbivory levels using both a cumulative link mixed model (CLMM) and a generalized linear mixed model (GLMM). In the CLMM, herbivory score (0–3) was treated as an ordinal categorical response variable; in the GLMM, presence/absence of any herbivory (including both scraping and chomping) was included as a binomial response variable. Population, AMF treatment, and population x AMF interaction were included as fixed effects; trial and maternal family were included as random effects. For CLMM and GLMM models, significance of fixed factors was determined through likelihood ratio tests; nonsignificant factors were dropped from both models. Subsequent pairwise comparisons of factor levels were tested using Tukey’s contrasts at α = 0.05 in the lsmeans package in R. All analyses were conducted using the R statistical platform (R Core Team 2015).

### 3 | RESULTS

#### 3.1 | Plant responses

Mycorrhizae significantly reduced plant biomass ($F_{1,488.2} = 237.5, \ p < 0.001$), flower production ($F_{1,486.1} = 231.4, \ P < 0.001$), seed mass ($F_{1,371.4} = 34.5, \ p < 0.001$), and seed weight per fruit ($F_{1,376.6} = 22.3, \ p < 0.01$) by 30%, 32%, 10%, and 15%, respectively (Figure 3; Supporting information Table S1). They also produced 6% fewer seeds per fruit although the difference was marginally nonsignificant ($F_{1,384.3} = 3.0, \ p = 0.08$). Total seed weight per fruit was significantly lower in CF plants than Del, C3, or Golf (Figure 3b). Seeds from Key and S22 were significantly lighter than all other populations except C3 (Figure 3d). Populations did not differ in their growth and seed production responses to AMF inoculation (Population x AMF term for responses in the order they were mentioned: $F_{5,488.8} = 0.29, \ p = 0.92; \ F_{5,485.8} = 1.12, \ p = 0.35; \ F_{5,371.2} = 1.14, \ p = 0.34; \ F_{5,376.2} = 0.64,$
Individual plants initiated flowering over a 10-day period. Overall, AMF inoculation significantly delayed flowering by 1 day ($F_{1,480.5} = 6.6$, $p < 0.05$; Figure 4a). The largest delay in flowering was observed in the earliest flowering population, Del, where AMF inoculation delayed flowering by 3 days (Tukey’s adjusted $p = 0.02$). Plant populations showed significant variation in time to flowering, although there was no clear geographic pattern (Figure 4a). The only significant population difference in flowering duration was between Del (31.4 ± 2.9 days) and Key (24.2 ± 3.5 days; Tukey’s adjusted $p = 0.03$; Figure 4b). There was no effect of AMF on flowering duration.

3.2 | Herbivore responses

AMF inoculation did not affect herbivore preference (CLMM: LR = 4.1, df = 6, $p = 0.66$; GLMM: LR = 3.1, df = 6, $p = 0.8$), suggesting that AMF infection did not confer greater plant resistance or susceptibility to herbivores. The overall fraction of fruits damaged by herbivores in trials was 56%. Of the damaged fruits, 27% received chomping herbivory and 73% received scraping herbivory.

Since likelihood ratio tests indicated no effect of AMF on herbivory rates, we continued analyses with the reduced model, testing the effect of plant population on herbivory, with trial and maternal family included as random factors. Populations varied significantly in herbivory, with the CLMM (accounting for the severity of herbivory) and GLMM (with herbivory as a binary response variable) models resulting in similar estimates of population differences. Overall, Key and S22
suffered the highest rates of herbivory, and Del and Golf suffered the lowest rates of herbivory in herbivore choice trials. Pairwise comparisons of populations for both GLMM and CLMM indicated significantly more herbivory for Key and S22 than for Del, and CLMM additionally identified Key as experiencing significantly more intense herbivory than Golf (Figure 5). Least-squares means from GLMM indicated that fruits from Del and Golf plants had 43% and 48% probability of herbivory, respectively, while fruits from Key and S22 plants each had 72% probability of herbivory. CLMM results indicated similar trends, with a tendency toward more severe herbivory in Key (proportional odds $= 2.4, p = 0.02$) and S22 (proportional odds $= 1.8, p = 0.1$).

4 | DISCUSSION

4.1 | Herbivore responses to AMF and plant population

AMF interactions with xantiana consistently and negatively affected plant performance, yet had no measurable effects on patterns of herbivory by the most destructive insect herbivore in natural populations (grasshoppers; Melanoplus spp.). The net effect of AMF on herbivory is likely conditional upon how plants respond to AMF in terms of tissue nutrient concentration, defensive chemical concentration, and growth (Gehring & Whitham, 2002; Koricheva et al., 2009). Therefore, the outcome of these aboveground–belowground interactions is complex and context dependent. For example, a review by Gehring and Bennett (2009) reported that 45% of studies investigating the effects of AMF on insect herbivory (from 20 publications) showed a net increase, 35% showed a net decrease, and 21% showed no significant effect. Decreases in herbivory due to AMF may be due to increased accumulation of constitutive defenses in plant parts or because AMF colonization of root cortical cells "primes" the plant immune system. In the latter case, AMF colonization provokes jasmonic acid-based plant immune responses, such that plants can upregulate the production of defense compounds more rapidly upon herbivore attack (Jung et al., 2012). Our study involved the harvest of fruits from plants that experienced no damage prior to herbivory trials and therefore is best suited for evaluating whether AMF influence herbivory via constitutive defense production.

Previous studies have shown that AMF cause greater constitutive defense production in some plant species, which in turn causes reduced herbivory; however, the results are mixed. For example, Dingle and Mousseau (1994) found increased levels of aucubin and catalpol, carbon-based herbivore deterrent compounds, in leaves of AMF-colonized Plantago laureola plants compared to fungicided controls. They reasoned that the change in abundance of these compounds was due to altered carbon/nutrient balance in mycorrhizal plants, but herbivory trials with two different insects showed contrasting results—chewing insects (caterpillars) favored the non-mycorrhizal plants but sucking insects (aphids) showed no preference (see also Roger et al. 2013). Another recent study found that AMF inoculated plants did not suffer significantly different levels of herbivory than control plants (Barber, Kiers, Hazzard, & Adler, 2013), despite differences in leaf nutrient concentrations between treatments. Our results, showing no influence of AMF inoculation on herbivore preference, reflect the conclusions of a recent review (Gehring & Bennett, 2009) that found effects of mycorrhizal interactions on plant–herbivore dynamics vary widely and are highly context dependent. In a meta-analysis of plant–mycorrhizal–herbivore studies (including both ecto- and endomycorrhizal experiments), Koricheva et al. (2009) came to similar conclusion of high variability and context dependency in these aboveground–belowground interactions, but did find a trend of decreased herbivore performance on AMF plants when the herbivore was a polyphagous chewing insect (such as the grasshoppers in this study), but not when the herbivore had a more restricted diet or was a sucking insect. Our results do not fit this pattern, although we did not measure herbivore performance, but rather herbivore preference. Herbivores did respond significantly differently to xantiana populations where two populations (Key, S22) experienced greater damage than the remaining populations. Those populations (Key, S22) occur near or at the range margin; however, the other marginal populations do not exhibit comparably high levels of herbivory (C3, Golf). Therefore, we did not observe a convincing geographic pattern to herbivore preference as we might have predicted based upon field studies that documented greater herbivore resistance in central populations.

4.2 | Plant responses to AMF

We found that all aspects of growth and reproductive effort were reduced in AMF inoculated plants, suggesting that the net effect of AMF inoculation on xantiana was parasitic in our experiment. This relationship did not vary among xantiana populations, as evidenced by a nonsignificant interaction between AMF and population treatments. Symbioses occur on a parasitism mutualism continuum, with the outcome of the symbiosis depending on environmental factors.

**FIGURE 5** Variation in herbivory damage among source populations in herbivore choice trials. Populations are ordered across the x-axis from range center to edge. Severity of herbivory increases with herbivory score (see Materials and Methods). Different letters above populations indicate significant differences according to Tukey’s pairwise comparison of least square means with a significance level of $\alpha = 0.05$
such as nutrient and light availability, and the genotypic identities of the plants and the AMF (Johnson et al., 1997; Klironomos, 2003). A meta-analysis by Hoeksema et al. (2010) found overall positive growth effects in response to mycorrhizal inoculation (both AMF and ectomycorrhizal fungi with non-N-fixing forbs), especially when multiple fungal species were used as inoculum. However, adverse growth effects on forbs in response to AMF are not uncommon. For example, in multiple field and greenhouse studies of agricultural tobacco, *Glomus macrocarpum* consistently and significantly reduced tobacco root length, aboveground biomass, and reproductive effort (Hendrix, Jones, & Nesmith, 1992; Jones & Hendrix, 1987; Modjo & Hendrix, 1986; Modjo, Hendrix, & Nesmith, 1987). Likewise, in a study testing outcomes of many pairwise combinations of plant and AMF species, Klironomos (2003) found large variation in plant growth responses to AMF inoculation, with AMF having both strongly negative and strongly positive effects on plant biomass. In that study, plant growth response ranged from −49% to +46% (% difference in plant growth between AMF and non-AMF plants). In light of these previous results, the AMF effects in the current study (−30% difference) fall within the range previously observed and toward the negative end of the spectrum. Further investigation may reveal whether the adverse growth effects witnessed in the current study were due to environmental aspects of the greenhouse treatments, the combination of *xantiana* and AMF genotypes, or both.

This experiment showed consistent, negative effects of AMF inoculation on not only plant growth, but other performance metrics more directly tied to individual lifetime fitness. Biomass is used as a fitness proxy in most experiments testing effects of AMF (Hoeksema et al., 2010), but in this study we also documented significant reductions in flower number and seed weight, and a marginally significant reduction in seed number, for AMF-inoculated plants. This highlights the fact that reductions in growth due to parasitic AMF interactions likely will be realized as significant reductions in lifetime fitness, even though direct measure of individual plants’ seed output is rarely reported in the AMF literature.

Low soil nitrogen availability has been shown to shift the mycorrhizal symbiosis toward parasitism in other systems (Johnson, Wilson, Wilson, Miller, & Bowker, 2015), and the low nutrient charge of the soil mix in this experiment, chosen to reflect field soil conditions (Methods), could have contributed to the parasitic effects of the AMF inoculum. Additionally, if there were microbial species in the inoculum other than AMF, they could have contributed to the plant response observed. The media of AMF inoculum can also change abiotic soil properties within inoculated pots, such as by increasing soil nutrient availability (Rowe, Brown, & Claassen, 2007), but any positive effects of nutrient addition via inoculum media were outweighed by negative effects of AMF inoculation in this experiment.

Although we saw this community of nine AMF species have adverse effects on *xantiana* in the greenhouse, it remains to be seen what effect AMF have on *xantiana* in a field setting, and how locally sourced AMF species affect growth. Most studies using commercially produced inocula reported increased plant growth in inoculated plants compared to uninoculated plants (Baum, El-Tohamy, & Gruda, 2015), including some that found commercial inoculum to provide benefits similar to or greater than native inoculum (Barber et al., 2013; but see Rowe et al., 2007; White, Tallaksen, & Charvat, 2008). Nevertheless, there is some evidence of genotype × genotype interactions between plants and root endophytes (Johnson et al., 2010), which suggests that local adaptation to microbial communities may influence the outcome of interactions. Greenhouse conditions have also been implicated in studies reporting AMF parasitism, since low levels of light during winter months may sufficiently depress photosynthetic rates that the nutrient benefit provided by AMF is not enough to overcome the carbon cost. Finally, field plants experience a complex biotic community that is absent from the greenhouse, and it is possible that positive effects of AMF are only realized in this more complex biotic context, where plants are interacting with multiple other microbial and invertebrate species (Hoeksema et al., 2010).

In addition to affecting growth and reproductive effort, microbial communities have also been shown to affect plant fitness by altering plant phenology. In arid regions like those inhabited by *xantiana*, plants can avoid periods of low water availability late in the growing season by shifting flowering and flowering earlier (Aronson, Kigel, Shmida, & Klein, 1992; Volls, 2007). For example, Lau and Lennon (2012) showed that the microbial community of the annual plant, *Brassica rapa*, was responsible for a phenological shift to earlier flowering under drought conditions. We found that, overall, AMF inoculation delayed flowering of *xantiana* by 1 day. This AMF-induced phenological shift was stronger in one population, Del, where flowering was delayed by 3 days with AMF inoculation (all plants initiated flowering within a 10-day time period). This delay may be due to direct influence of AMF colonization on biochemical pathways influencing the switch from growth to reproduction in *xantiana*, or an indirect effect where plant resource allocation to AMF hinders development and delays flowering. Earlier flowering time in *xantiana* is favored by selection in some environments (Geber and Eckhart 2005), in part because flowering time strongly influences a plant’s probability of fatal mammal herbivory in some parts of *xantiana*’s range. For example, at the edge of *xantiana*’s range, each day delay in flowering increases a plant’s odds of fatal herbivory by 5%, and this fatal mammal herbivory can have large effects on population growth rates (Benning, Eckhart, Geber, & Moeller, 2018). Our results suggest that microbial communities may play an important role in modulating the expression of this ecologically important trait.

5 | CONCLUSIONS

Plant–AMF interactions have been shown to indirectly affect plant–herbivore interactions by influencing the quantity and quality of plant tissues as well as the production of defensive compounds (Smith and Read 1997, Bennett, Alers-Garcia, & Bever, 2006). In this study, we found consistent negative effects of a community of nine AMF species on plant growth and reproduction but no indirect effects on herbivory to fruits by insect herbivores. The negative effect of AMF on plant growth does not appear to similarly influence plant tissue quality from the perspective of grasshoppers. Populations
sampled from across a geographic gradient in the abiotic and biotic environment did not differ in plant responses to the AMF community. There was some variation in herbivore preference for fruits from different plant populations, but not a clear geographic pattern. Overall, our results are consistent with previous observations that plant–AMF interactions vary considerably from mutualistic to parasitic, and that plant–herbivore interactions may not be modified by AMF even when AMF effects on plants are strong.

ACKNOWLEDGMENTS

We thank the National Science Foundation for supporting our research on *Clarkia xantiana* (DEB-1255141 to Moeller including an REU supplement for LGB and DEB-1701072 to Moeller and Benning), and Stefan Jaronski of the USDA ARS Northern Plains Agricultural Research Laboratory for generously providing a grasshopper colony for our experiment. The manuscript was much improved by discussions with A. Pringle, M. Hart, and M. Keller-Pearson, and from thoughtful comments by J. Lau, the Lau lab, and the Moeller lab.

AUTHOR CONTRIBUTIONS

LB, JB, and DM conceived the ideas and methodology. LB wrote the manuscript; LB and JB collected and analyzed the experimental data. All authors contributed critically to the drafts and gave final approval for publication.

DATA ACCESSIBILITY STATEMENT

All experimental data are available at datadryad.org: https://doi.org/10.5061/dryad.176gv58.

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REFERENCES

Aronson, J., Kigel, J., Shmida, A., & Klein, J. (1992). Adaptive phenology of desert and Mediterranean populations of annual plants grown with and without water stress. *Oecologia*, 89, 17–26. https://doi.org/10.1007/BF00319010

Augé, R. M. (2001). Water relations, drought and vesicular-arbuscular mycorrhizal symbiosis. *Mycorrhiza*, 11, 3–42.

Babikova, Z., Gilbert, L., Bruce, T., Dewhirst, S. Y., Pickett, J. A., & Johnson, D. (2014). Arbuscular mycorrhizal fungi and aphids interact by changing host plant quality and volatile emission. *Functional Ecology*, 28, 375–385. https://doi.org/10.1111/1365-2435.12181

Barber, N. A., Kiers, E. T., Hazzard, R. V., & Adler, L. S. (2013). Context-dependency of arbuscular mycorrhizal fungi on plant-insect interactions in an agroecosystem. *Frontiers in Plant Science*, 4, 338.

Bardgett, R. (2005). The *Biology of Soil*: A Community and Ecosystem Approach. Oxford, England, UK: OUP Oxford. https://doi.org/10.1093/acprof:oso/9780198525035.001.0001

Baum, C., El-Tohamy, W., & Gruda, N. (2015). Increasing the productivity and product quality of vegetable crops using arbuscular mycorrhizal fungi: A review. *Science Horticuluture*, 187, 131–141. https://doi.org/10.1016/j.scienta.2015.03.002

Bennett, A. E., Alers-Garcia, J., & Bever, J. D. (2006). Three-way interactions among mutualistic mycorrhizal fungi, plants, and plant enemies: Hypotheses and synthesis. *American Naturalist*, 167, 141–152.

Bennett, A. E., & Bever, J. D. (2009). Trade-offs between arbuscular mycorrhizal fungal competitive ability and host growth promotion in *Plantago lanceolata*. *Oecologia*, 160, 807–816. https://doi.org/10.1007/s00442-009-1345-6

Bennett, A. E., Millar, N. S., Gedroics, E., & Karley, A. J. (2016). Plant and insect microbial symbionts alter the outcome of plant-herbivore-parasitoid interactions: Implications for invaded, agricultural and natural systems. *Journal of Ecology*, 104, 1734–1744. https://doi.org/10.1111/1365-2745.12620

Benning, J., Eckhart, V. M., Geber, M. A., & Moeller, D. A. (2018). Biotic interactions limit the geographic range of an annual plant: herbivory and phenology mediate fitness beyond a range margin. *bioRxiv*, 300590.

Berg, E. S., Eaton, G. K., & Ayres, M. P. (2001). Augmentation of AM fungi fails to ameliorate the adverse effects of temporal resource variation on a lettuce crop. *Plant and Soil*, 236, 251–262. https://doi.org/10.1023/A:1012729112136

Borowicz, V. A. (2001). Do arbuscular mycorrhizal fungi alter plant-pathogen relations? *Ecology*, 82, 3057–3068.

Cavender, N., & Knee, M. (2006). Relationship of seed source and arbuscular mycorrhizal fungi inoculum type to growth and colonization of big bluestem (*Andropogon gerardii*). *Plant and Soil*, 285, 57–65. https://doi.org/10.1007/s11104-005-5228-2

Dingle, H., & Mousseau, T. A. (1994). Geographic variation in embryonic development time and stage of diapause in a grasshopper. *Oecologia*, 97, 179–185. https://doi.org/10.1007/BF00323147

Dosskey, M. G., Boersma, L., & Linderman, R. G. (1991). Role for the photosynthetic demand of ectomycorrhizas in the response of Douglas fir seedlings to drying soil. *New Phytologist*, 117, 327–334. https://doi.org/10.1111/j.1640-1238.1991.tb04914.x

Eckhart, V. M., & Geber, M. A. (1999). Character variation and geographic distribution of *Clarkia xantiana* A. Gray (*Onagraceae*): Flowers and phenology distinguish two subspecies. *Madroño*, 46, 117–125.

Eckhart, V. M., Geber, M. A., Morris, W. F., Fabio, E. S., Tiffin, P., & Moeller, D. A. (2011). The geography of demography: Long-term demographic studies and species distribution models reveal a species border limited by adaptation. *American Naturalist*, 178, S26–S43. https://doi.org/10.1086/661782

Eckhart, V. M., Singh, I., Louthan, A. M., Keledjian, A. J., Chu, A., Moeller, D. A., & Geber, M. A. (2010). Plant-soil water relations and species border of *Clarkia xantiana*ssp. *xantiana* (*Onagraceae*). *International Journal of Plant Sciences*, 171, 749–60. https://www.jstor.org/stable/10.1086/654845

Engelkes, T., Morriën, E. M., Verhoeven, K. J. F., Bezemer, T. M., Biere, A., Harvey, J. A.,... van der Putten, W. H. (2008). Successful range-expanding plants experience less above-ground and below-ground enemy impact. *Nature*, 456, 946–948. https://doi.org/10.1038/nature07474

Gange, A. C., & Smith, A. K. (2005). Arbuscular mycorrhizal fungi influence visitation rates of pollinating insects. *Ecological Entomology*, 30, 600–606. https://doi.org/10.1111/j.0307-6946.2005.00732.x

Gange, A. C., & West, H. M. (1994). Interactions between arbuscular mycorrhizal fungi and foliar-feeding insects in *Plantago lanceolata* L. *The New Phytologist*, 128, 79–87. https://doi.org/10.1111/j.1469-8137.1994.tb03989.x

Garrido, E., Bennett, A. E., Fornoni, J., & Strauss, S. Y. (2010). Variation in arbuscular mycorrhizal fungi colonization modifies the expression of tolerance to above-ground defoliation. *Journal of Ecology*, 98, 43–49. https://doi.org/10.1111/j.1365-2745.2009.01586.x
The effect of mycorrhizal associations along the mutualism–parasitism continuum. New Phytologist, 135, 575–585. https://doi.org/10.1046/j.1469-8137.1997.00729.x

Johnson, N. C., Rowland, D. L., Corkidi, L., Egerton-Warburton, L. M., & Allen, E. B. (2003). Nitrogen enrichment alters mycorrhizal allocation at five mesic to semiarid grasslands. Ecology, 84, 1895–1908.

Johnson, N. C., Wilson, G. W. T., Bowker, M. A., Wilson, J. A., Miller, R. M., & Tilman, D. (2010). Resource limitation is a driver of local adaptation in mycorrhizal symbioses. Proceedings of the National Academy of Sciences of the United States of America, 107, 2093–2098. https://doi.org/10.1073/pnas.0906710107

Johnson, N. C., Wilson, G. W., Wilson, J. A., Miller, R. M., & Bowker, M. A. (2015). Mycorrhizal phenotypes and the Law of the Minimum. New Phytologist, 205, 1473–1484. https://doi.org/10.1111/nph.13172

Jones, K., & Hendrix, J. W. (1987). Inhibition of root extension in tobacco by the mycorrhizal fungus Glomus macrocarpum and its prevention by benomyl. Soil Biology & Biochemistry, 19, 297–299. https://doi.org/10.1016/0038-0717(87)90012-5

Jung, S. C., Martinez-Medina, A., Lopez-Raez, J. A., & Pozo, M. J. (2012). Mycorrhiza-induced resistance and priming of plant defenses. Journal of Chemical Ecology, 38, 651–664. https://doi.org/10.1007/s10886-012-0134-6

Kaeppeler, S. M., Parke, J. L., Mueller, S. M., Senior, L., Stuber, C., & Tracy, W. F. (2000). Variation among maize inbred lines and detection of quantitative trait loci for growth at low phosphorus and responsiveness to arbuscular mycorrhizal fungi. Crop Science, 40, 358–64.

Klironomos, J. N. (2003). Variation in plant response to native and exotic arbuscular mycorrhizal fungi. Ecology, 84, 2292–2301. https://doi.org/10.1890/02-0413

Koide, R. T., & Dickie, I. A. (2002). Effects of mycorrhizal fungi on plant populations. Plant and Soil, 244, 307–317. https://doi.org/10.1023/A:1020204004844

Koricheva, J., Gange, A. C., & Jones, T. (2009). Effects of mycorrhizal fungi on insect herbivores: A meta-analysis. Ecology, 90, 2088–2097. https://doi.org/10.1890/08-1555.1

Larimer, A. L., Clay, K., & Bever, J. D. (2014). Synergism and context dependency of interactions between arbuscular mycorrhizal fungi and Rhizobia with a Prairie legume. Ecology, 95, 1045–1054. https://doi.org/10.1890/13-0025.1

Lau, J. A., & Lennon, J. T. (2012). Rapid responses of soil microorganisms improve plant fitness in novel environments. Proceedings of the National Academy of Sciences of the United States of America, 109, 14058–14062. https://doi.org/10.1073/pnas.1202319109

Lewis, H., & Lewis, M. E. (1955). The genus Clarkia. Berkeley, Calif: University of California Press.

Maron, J. L., & Crone, E. (2006). Herbivory: Effects on plant abundance, distribution and population growth. Proceedings of the Royal Society B, 1601, 2575–84. https://doi.org/10.1098/rspb.2006.3587

Modjo, H. S., & Hendrix, J. W. (1986). The mycorrhizal fungus Glomus macrocarpum as a cause of tobacco stunt disease. Phytopathology, 76, 688–691. https://doi.org/10.1094/Phyto-76-688

Modjo, H. S., Hendrix, J. W., & Nesmith, W. C. (1987). Mycorrhizal fungi in relation to control of tobacco stunt disease with soil fumigants. Soil Biology & Biochemistry, 19, 289–295. https://doi.org/10.1016/0038-0717(87)90011-3

Orr, M. R., Porter, A. H., Mousseau, T. A., & Dingle, H. (1993). Molecular and morphological evidence for hybridization between two ecologically distinct grasshoppers (Melanoplus sanguinipes and M. devastator) in California. Heredity, 72, 42–54.

Pineda, A., Dicke, M., Pieterse, C. M. J., & Pozo, M. J. (2013). Beneficial microbes in a changing environment: Are they always helping plants to deal with insects? Functional Ecology, 27, 574–586. https://doi.org/10.1111/1365-2435.12050

Pischl, P. H., & Barber, N. A. (2016). Plant responses to arbuscular mycorrhizae under elevated temperature and drought. Journal of Plant Ecology, 10, 692–701.

Porcel, R., Aroca, R., & Ruiz-Lozano, J. M. (2012). Salinity stress alleviation using arbuscular mycorrhizal fungi. A review. Agronomy for Sustainable Development, 32, 181–200. https://doi.org/10.1007/s13593-011-0029-x

Rasmussen, P. U., Amin, T., Bennett, A. E., Green, K. K., Timonen, S., Van Nouhuys, S., & Tack, A. J. M. (2017). Plant and insect genetic variation mediate the impact of arbuscular mycorrhizal fungi on a natural plant–herbivore interaction. Ecological Entomology, 42, 793–802. https://doi.org/10.1111/een.12453

R Core Team (2017). R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. URL https://www.R-project.org/

Roger, A., Getaz, M., Rasmann, S., & Sanders, I. R. (2013). Identity and combinations of arbuscular mycorrhizal fungal isolates influence plant resistance and insect preference. Ecological Entomology, 38, 330–338. https://doi.org/10.1111/een.12022.

Rowe, H. I., Brown, C. S., & Claassen, V. P. (2007). Comparisons of mycorrhizal responsiveness with field soil and commercial inoculum for six native montane species and Broums tectorum. Restoration Ecology, 15, 44–52. https://doi.org/10.1111/j.1526-100X.2006.00188.x

Smith, S. E., & Read, D. J. (1997). Mycorrhizal Symbiosis. Academic Press.

Smith, S. E., & Read, D. J. (2008). Mycorrhizal symbiosis, 3rd ed.. New York: Elsevier.

Van der Putten, W. H., Vet, L. E. M., Harvey, J. A., & Wäckers, F. L. (2001). Linking above- and belowground multitrophic interactions of plants, herbivores, pathogens, and their antagonists. Trends in Ecology & Evolution, 16, 547–554.

Vannette, R. L., & Hunter, M. D. (2011). Plant defence theory re-examined: Nonlinear expectations based on the costs and benefits of resource mutualisms. Journal of Ecology, 99, 66–76. https://doi.org/10.1111/j.1365-2745.2010.01755.x

Volis, S. (2007). Correlated patterns of variation in phenology and seed production in populations of two annual grasses along an aridity gradient. Evolutionary Ecology, 21, 381–393. https://doi.org/10.1007/s10682-006-9108-x

Wardle, D. A. (2002). Communities and ecosystems: Linking the aboveground and belowground components. Princeton, NJ: Princeton University Press.
White, J. A., Tallaksen, J., & Charvat, I. (2008). The effects of arbuscular mycorrhizal fungal inoculation at a roadside prairie restoration site. Mycologia, 100, 6–11. https://doi.org/10.1080/15572536.2008.11832494

Wiseman, P. E., & Wells, C. E. (2009). Arbuscular mycorrhizal inoculation affects root development of Acer and Magnolia species. Journal of Environmental Horticulture, 27, 70–79.

Wolfe, B. E., Husband, B. C., & Klironomos, J. N. (2005). Effects of a belowground mutualism on an aboveground mutualism. Ecology Letters, 8, 218–223. https://doi.org/10.1111/j.1461-0248.2004.00716.x

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**How to cite this article:** Bolin LG, Benning JW, Moeller DA. Mycorrhizal interactions do not influence plant–herbivore interactions in populations of *Clarkia xantiana* ssp. *xantiana* spanning from center to margin of the geographic range. *Ecol Evol.* 2018;8:10743–10753. [https://doi.org/10.1002/ee3.4523](https://doi.org/10.1002/ee3.4523)