

Title: Plant G x Microbial E: Plant genotype interaction with soil bacterial community shapes rhizosphere composition during invasion

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ABSTRACT

It is increasingly recognized that different genetic variants of hosts can uniquely shape their microbiomes. Invasive species often evolve in their introduced ranges, but little is known about the potential for their microbial associations to change during invasion as a result. We asked whether host genotype (G), microbial environment (E), or their interaction (GxE) affected the composition and diversity of host-associated microbiomes in *Centaurea solstitialis* (yellow starthistle), a Eurasian plant that is known to have evolved novel genotypes and phenotypes, and to have altered microbial interactions, in its severe invasion of California, USA. We

conducted an experiment in which native and invading plant genotypes were inoculated with native and invaded range soil microbial communities. We used amplicon sequencing to characterize rhizosphere bacteria in both the experiment and the field soils from which they were derived. We found that native and invading plant genotypes accumulated different microbial associations at the family level in each soil community, often counter to differences in family abundance between soil communities. Root associations with potentially beneficial Streptomycetaceae were particularly interesting, as these were more abundant in the invaded range field soil and accumulated on invading genotypes. We also found that bacterial diversity is higher in invaded soils, but that invading genotypes accumulated a lower diversity of bacteria and unique microbial composition in experimental inoculations, relative to native genotypes. Thus variation in microbial associations of invaders was driven by the interaction of plant G and microbial E, and rhizosphere microbial communities appear to change in composition in response to host evolution during invasion.

INTRODUCTION

Introduced species that proliferate in their new environments are able to do so as a result of a suite of environmental and biological interactions [1, 2]. Biotic interactions are thought to be a particularly important means by which an introduced species could become invasive, because of opportunities for escape from enemies (i.e. the Enemy Release Hypothesis [3–7]) or gain of mutualisms [8–11] in an introduced environment. Altered interactions with enemies or mutualists could have immediate fitness benefits to invaders, but also might allow introduced populations to evolve phenotypes that increase invasiveness, in response to

increased resource availability from mutualists and/or relaxed selection for defense from enemies [12, 13]. While the potential for invaders to adapt in response to differences in species interactions in this way has received considerable attention (e.g. the Evolution of Increased Competitive Ability Hypothesis [14, 15]), it is also the case that invader evolution for any reason, adaptive or otherwise, might itself alter species interactions. The question of whether altered species interactions arise from a novel environment ('E') or from invader evolution (i.e. genotype, 'G'), or their interaction ('GxE') is largely unexplored, though the contributions of these different effects would have important consequences for understanding the mechanisms of invasions and predicting further spread (e.g. see [16, 17]).

For invasive plants, a critical source of both enemy and mutualist interactions is the soil microbial community [5, 18–26]. Some invasive plants perform better with soil microbial communities from invaded ranges, which suggests that initial interactions with soil microbes could facilitate invasion [4, 20, 27–30] but see [31]. Invaders could also benefit from differences in plant-soil feedbacks that accumulate over time [32]. Plants experience negative feedbacks when a host-specific pathogen accumulates in the soil and suppresses their growth [33, 34], or positive feedbacks if the same dynamic occurs with a mutualist [35], and invading species have been observed to benefit from enhanced positive or reduced negative feedbacks in soils from their introduced ranges [24, 29, 35–38]. Invaders might also benefit from interactions with a simplified microbial interaction network in a novel environment, allowing invaders more specificity in exploiting a smaller number of mutualists or defending against fewer pathogens [12, 39].

Regardless of whether microbial pathogens or mutualists differ between native and invaded ranges, we know very little about whether invaders that undergo evolutionary change over the course of their invasion interact with soil microbes differently than native genotypes. It is well documented in the fields of plant disease and agriculture that the genotype of a plant can be tightly correlated with the presence or relative abundance of specific microbes in and around their tissues [40–42]. Plant genotypes can control microbial communities through differences in their resulting phenotype, via traits like root and leaf morphology [43], exudate production [44, 45], and interaction between plant phenotype and environment (e.g. root morphology) [46]. Plant genotypes vary in the amount or type of compounds they produce in defense against microbes [47], or as incidental secondary metabolites that nonetheless impact microbial communities associating with their tissues [48]. These types of genetically-based phenotypic differences have been shown to control plant association with individual microbial taxa, as well as the plant-associated microbial community makeup [44, 45, 49]. The strong existing evidence that plant genotype affects microbial associations suggests that evolution of invader genotype could lead to changes in microbial associations, which could then play a role in invader success and proliferation.

Indeed, invaders are well known to experience rapid evolution for a variety of reasons. Introduced populations can adapt to their new environments and the process of colonization [50–52], but they may also experience non-adaptive genetic changes through several mechanisms, including initial and serial founder events, multiple introductions and admixture of previously isolated subpopulations, hybridization with other species, stimulation of transposable element activity, and/or the revealing of cryptic genetic variation in a novel

environment [39, 53–57]. Given that microbial associations respond to host genotype (as discussed above), such broad opportunities for genetic change in invaders are likely to create opportunities for altered species interactions, particularly with microbial communities that are sensitive to variation in plant growth and chemistry.

We test whether invader evolution alters microbial interactions in the invasive annual forb yellow starthistle (*Centaurea solstitialis*, L.; Asteraceae). This species is native to Eurasia, introduced to the Americas, Australia, and South Africa, and highly invasive in grasslands of North and South America [58]. The *C. solstitialis* invasion of California, USA is particularly well characterized for its evolution of novel genotypes and phenotypes. Previous work has used genome-wide markers to reconstruct its invasion history and detail genetic divergence among populations [59–61]. This work finds that plant populations invading California are unique genotypes, genetically divergent from populations elsewhere, but derived from a single subpopulation in Spain and southern France [59]. Levels of genetic diversity in California populations are similar to those in native range populations, indicating the lack of any strong genetic bottlenecks during the divergence of this invasion [59]. The California genotypes are also distinct from their native progenitors in terms of genetically-based phenotypes, including having evolved larger seeds, larger plant size (leaf number, leaf area, height, and biomass), higher reproduction (reproductive duration and number of capitula), and greater competitive ability against grasses than native genotypes [59, 61–65]. Most recently, these phenotypes are being genetically mapped to the regions of the genome responsible for these evolved differences in the invaders [66].

Invading *C. solstitialis* plants in California also appear to accumulate more biomass from

more favorable (reduced deleterious) interactions with the soil microbial community in the invaded range [30, 67, 68], raising the possibility that the larger and more fecund invading genotypes have evolved to take advantage of release from microbial enemies or gain of mutualists [69]. These experiments tested the performance of *C. solstitialis* plants in soil inoculated with soil microbial communities collected from different parts of the global range but did not test for evolved differences among host plants. Field experiments treating *C. solstitialis* plots with fungicide found that fungi in the invaded range are less favorable for plant performance, which has generated interest in the potential for bacterial communities to play an important role in creating more favorable growth conditions in the invasion [70]. A previous survey of bacteria associated with *C. solstitialis* in the field by Lu-Irving and colleagues [69] found that bacterial diversity, including of known pathogenic groups, was lower on invading plants. The authors hypothesized that the lower diversity of plant-associated bacteria could reflect escape from enemies in the invaded range [69]. Their results suggest that invaders might benefit from altered interactions with the microbial community, but it is not clear to what extent observed microbiome differences are a cause or effect of the evolution of plant invader genotypes.

Here, we identify the roles of both microbial community environment, plant genotype, and their interaction in shaping plant-microbial associations in this system. Based on previous observations of plant microbiome differences between ranges, we predict that soil microbial communities will be the primary source of variation in microbial associations, and we predict that plants (native or invading genotypes) grown with invaded range soil inoculum will accumulate a lower diversity of rhizosphere bacteria. Using the same populations sampled in

previous studies of both plant evolution and microbiome variation in this system [59, 62, 69], we sample field soil from native and invaded ranges to identify the bacterial community available for plant interactions. We conduct a factorial greenhouse experiment, in which both native genotypes and invading genotypes of *C. solstitialis* are grown in native and invaded range soil microbial communities, to identify which members of the soil community recruit to roots of the genotype from each range. We identify bacteria using 16S amplicon sequencing, and test for associations with genotype and soil source. In contrast to our predictions, we find that the interaction of both plant genotype and microbial environment influence specific microbial associations, and we find that previously observed patterns of lower diversity of bacterial communities on invading plants are not inherent features of the microbial environment as formerly hypothesized, but are instead the product of novel plant genotype interactions in the invasion.

METHODS

Study system biology

Centaurea solstitialis was introduced accidentally as a contaminant of alfalfa into South America in the 1600s and then North America in the 1800s [58, 71]. It is an obligately outcrossing diploid annual plant that has four distinct native genetic subpopulations across Eurasia [60, 72–74]. In both the California invasion and its source subpopulation in western Europe, *C. solstitialis* is an annual plant that grows as a rosette with a taproot through a mild and wet winter and spring, then bolts and flowers throughout the dry summer [62, 63, 75].

Soil and seed sampling

Field soils used in this study were collected when seeds were being produced on *C. solstitialis* plants, during dry summer conditions in August 2018. Soils came from four invaded sites in California, USA and four native sites in southern Spain and France, all of which were locations previously included in other studies of this system, including in the comparisons of plant-associated microbial diversity by Lu-Irving and colleagues [69] plant genetic diversity by Barker and colleagues [59], and plant trait variation by Dlugosch and colleagues [76](Supporting Information Table S1). At each site, a 30 m linear sampling transect was established through a *C. solstitialis* patch, and a second 30 m sampling transect was established outside of the patch, parallel to the first and separated by ~5 m, to capture site variation outside of *C. solstitialis* patches. The distance between transects was chosen to be adjacent but outside of active growth by *C. solstitialis*, noting that microbial communities have been observed to vary more by soil characteristics than geographic distance even at small scales [77, 78]. Samples from both transects were ultimately grouped for analysis by site (see below). A soil sampling point was established every two meters along each transect. At each sampling point, a 18 mm diameter soil core was used to sample the topmost 5-10 cm of soil from three adjacent cores (<10 cm apart), which were combined into a single sample for that sampling point, in a sealed plastic bag. This resulted in 15 separate samples per transect (one per sampling point). Larger bulk soil collections (to be used for experimental inoculum) were made near each transect, both inside and outside of *C. solstitialis* patches. For these bulk collections, 1 L plastic bags were filled with soil collected from the top 5-10 cm of soil from a single location. Gloves and tools were sterilized between soil samples by wiping with 70% isopropanol. Soils were stored dry (as

collected) at room temperature until their use in the greenhouse experiment. Seeds were collected from the plant closest to each meter along the transect through the *C. solstitialis* plants, for a total of 30 seed collections per site.

Soil samples were cleaned and homogenized for DNA sampling and use as inoculum. Large rocks and organic particles were removed from each sample manually with sterilized forceps in a biosafety cabinet, and the resulting sample sifted through a sterilized 2 mm sieve. Soil aggregates that did not pass through the sieve were ground gently with a mortar and pestle to break up the aggregates until all soil particles could pass through the 2mm sieve. The sample was stirred to homogenize, and 250 mg was weighed and collected in a 2 ml microcentrifuge tube for DNA extraction.

Experimental inoculations

We used a fully factorial greenhouse experiment to test for the effects of seed source (native vs invaded range) and soil microbial community source (native vs invaded range) combinations on rhizosphere bacterial composition and diversity, using seeds and soils from all sites described above. Soil microbial communities were derived from the bulk collections made from outside of *C. solstitialis* patches at each site (Supporting Information Table S1). All seeds were germinated on the surface of sterile soil (50:50 mix of sand and high porosity soil; HP Pro-Mix™, Quebec, CA) in petri dishes, in a greenhouse set to a maximum of 21°C day and 10°C night at the University of Arizona (Tucson, AZ, USA) College of Agriculture and Life Sciences greenhouses in February 2019. Germination date of each seed was noted to account for differences in germination timing (see Analyses). One week after germinating, seedlings were

potted in individual Deepots™ (410mL; D25L from Stuewe & Sons, Oregon USA) in sterile soil and inoculated with a surface application of 15 mL of unsterilized (live) soil slurry from soils collected in the same locations, or sterilized soil as a control. Slurries were prepared by adding 150 mL of sterile water to 10 mL of soil, mixing, and filtering through sterile cotton gauze.

Deepots were arranged in five spatial blocks in the greenhouse. Each block included all four combinations of plants from the native or invaded range planted into soil inoculum from the native or invaded range. Within each of these four combinations were up to 16 combinations of plants and soil inoculum from the four source populations within each range. Multiple source populations were included to capture variation within ranges, and were grouped as replicates by range for analyses described below. Replication for each plant range x soil inoculum range combination, including all five blocks, ranged from 43 to 72 plants whose rhizosphere communities were successfully sequenced.

Plants were maintained in the greenhouse for two months, when they were harvested by block within a single week. Sterile water was provided through daily misting by a reverse osmosis irrigation system (Evolution-RO, Hydro-Logic Inc., Port Washington, NY, USA). After one month, plants were fertilized every two weeks using autoclaved Hoagland's solution ($\frac{1}{8}$ strength Hoagland Complete Medium, BioWorld, Dublin, OH, USA). At harvest, plants were removed from their pots for rhizosphere collection. The upper 2 to 5 cm of the taproot was collected, together with attached lateral roots and these root tissues were processed and analyzed together. Excess soil was brushed or shaken off, and root samples were placed in individual 50 ml tubes containing 25 ml of sterile wash solution (45.9 mM NaH_2PO_4 , 61.6 mM Na_2HPO_4 , 0.1% Tween 20). Tubes were shaken by hand for 1 minute. Root samples were then removed and

stored on ice in tubes containing 10 ml of wash solution until further processing, stored on ice during transport and then refrigerated at 4°C. Ectorrhizosphere washes were centrifuged at 2,200 g at 4°C for 15 min, supernatants were discarded, and pellets were air-dried and stored at 20°C until DNA extraction. Plant growth and reproduction traits were not collected in this study (but have been extensively studied elsewhere, as described in the Introduction), due to the early age of plants at destructive harvest.

DNA extraction and amplicon sequencing

Microbial DNA was extracted from soil samples using the DNeasy PowerSoil Pro kit (Qiagen, Hilden, Germany). A blank sample of nanopure water was also included in extractions to record any contamination. Resulting double-stranded DNA concentration was quantified using a Qubit fluorometer (Broad Range kit, Invitrogen, Waltham MA, USA).

The 16s rRNA region was amplified using a 2-step polymerase chain reaction (PCR) as in [69] as follows. Target-specific PCR (PCR 1) was conducted by creating a 25.1 µL reaction mixture using 1 µL of microbial DNA, 1.3 µL of 515-F primer (5'-GTGCCAGCMGCCGCGGTAA-3') and 1.3 µL of 806-R primer (5'-GGACTACHVGGGTCTTAAT-3'), 12.5 µL of Phusion Flash master mix (Thermo Scientific, Waltham MA, USA), and 9 µL PCR grade water. Reaction mixtures were placed in an Eppendorf Mastercycler thermal cycler starting with 98°C for 10 seconds, then 25 cycles of this sequence: 98°C for 1 second, 78°C for 5 seconds, 57°C for 5 seconds, 72°C for 15 seconds, 72°C for 1 minute [79]. PCR 1 products were visualized on an agarose gel to determine whether PCR was successful before a second PCR step (PCR 2) was used to incorporate sequencing adapters onto the ends of the amplified PCR products. PCR 2 reaction mixtures were

created using 1 µL of PCR 1 product, 12.5 µL Phusion Flash master mix, and 0.75 µL of a unique barcoded primer combination provided by the University of Idaho's iBEST Genomic Resources Core. Our PCR program ran at 98°C for 10 seconds, then 10 cycles of: 98°C for 1 second, 78°C for 5 seconds, 51°C for 5 seconds, 72°C for 15 seconds, 72°C for 1 minute. Barcoded amplicons were quantified by Qubit fluorometry, pooled in equimolar amounts, cleaned using a MinElute kit (Qiagen, Hilden, Germany), and submitted to the University of Idaho's iBEST Genomic Resources Core for quality control and sequencing with Illumina MiSeq 350 bp pair-end sequencing.

Analyses

Microbial metabarcoding data was processed and analyzed in Qiime 2 version 2019.10 [80], and additional analyses were carried out in R [81], as detailed below. Scripts for processing sequences and replicating all analyses are available on GitHub (<https://github.com/mBerlow/PlantGxMicrobialE.git>). Sequences were denoised in Qiime 2 using the Divisive Amplicon Denoising Algorithm (DADA2) to remove sequence errors and trim primers [82]. Next, sequences were aligned, and a phylogeny was generated using FastTree, rooted at the midpoint [83]. Sequences were grouped at the level of amplicon sequence variants (ASVs, 100% similarity) and taxonomy was assigned using the Silva database, version 138 [84]. ASVs assigned to non-bacterial kingdoms were filtered out for the purposes of our analyses.

Bacterial alpha diversity was measured as richness calculated by the R package 'vegetarian' [85] and Shannon's Diversity Index calculated by Qiime2 after rarefying to 1000

sequences per sample [79]. Rarefaction curves indicated that our sequencing coverage was sufficiently exhaustive to observe plateaus in the accumulation of diversity within nearly all of our samples, with diminishing accumulation of diversity at 1000 sequences (Figure S1)[80]. To test for the effects of soil source (native vs. invaded range) on alpha diversity metrics, we used two-sided t-tests in the stats package in R [81]. To test for effects of experimental treatments on alpha diversity metrics, we used linear models that included effects of seed genotype (native vs. invader), soil source (native vs. invaded range), germination date of the plant (i.e. its age at harvest), experimental block in the greenhouse, and the interaction between seed genotype and soil source (see figure S4 & S5 for normality test results). Non-significant interaction terms ($P > 0.1$) were not included in the final models. Linear models and Tukey's post hoc tests were conducted using the stats package in R [81]. To determine which bacterial taxa were differentially abundant between sample types and treatment groups we used Linear Discriminant Analysis Effect Size (LEfSe) [86]. LEfSe identifies taxa whose abundances differ significantly between treatments using a Kruskal Wallance sum-rank test, investigates biological significance with a Wilcoxon rank-sum test, and finally calculates effect sizes of each differentially abundant taxa using linear discriminant analysis (LDA).

Bacterial community beta diversity was measured as unweighted and weighted UniFrac distances, a dissimilarity measure that accounts for phylogenetic relatedness [87]. Unweighted UniFrac distance accounts for information about presence/absence of ASVs and can be thought of as community membership, while weighted UniFrac distance also accounts for relative abundances and can be thought of as community structure. We used PERMANOVA conducted in the R package vegan [88] with 9999 permutations to test whether beta diversity distances

were predicted by fixed effects of range (native vs. invaded), and sites nested within range (including samples both inside and outside of *C. solstitialis* patches). We calculated partial omega squared effect size using the MicEco package in R.

Given that soil microbial communities can change over very small spatial scales [CITATIONS for this again], we assessed whether the bulk soil collections (used as inocula for the greenhouse experiment) were representative of the soil bacterial diversity sampled across the transects at the same site. To do this, we compared beta diversity distances between each bulk soil sample and the soil transect samples from the same site with the beta diversity distances between that same bulk soil sample and all the soil samples from other sites. Comparisons were made using paired t-tests in R.

RESULTS

We sequenced a total of 351 samples, yielding 16,654,396 paired reads (mean = 44,059, SD = 35,066, min = 12, max = 173,097, see Table S2 for sequence and ASV counts for each sample and Figure S1 for rarefaction curves). Sequences are available on the NCBI Sequence Repository (SUB13812121).

Field environment: Soils

Across our native and invaded range field soil samples we identified 29 phyla, including 294 families of bacteria. Eleven phyla made up 99% of sequences present (Figure 1, Table S3). The most abundant phyla across all samples were Actinobacteriota (an average of 33% of invaded soil bacterial communities, SD = 9%; 41% of native, SD = 14%), Proteobacteria (22% of

invaded, SD = 6%; 20% of native, SD = 14%), and Acidobacteriota (14% of invaded, SD = 5%; 18% of native, SD = 8%).

Bacterial taxa differed between soils from the native and invaded range at both the phylum and family level. Ten phyla were differentially abundant: Actinobacteriota, Acidobacteriota, and Myxococcota were more abundant in native range soils, and Proteobacteria, Armatimonadota, Gemmatimonadota, Planctomycetota, Chloroflexi, Bacteriodota, and Firmicutes were more abundant in invaded range soils (LDA effect size >2, LEfSe; Figure S2). Nineteen bacterial families were more abundant in native range soils, and 55 families were more abundant in invaded range soils (LDA effect size >2, LEfSe; Table S4).

The distribution of soil bacterial diversity also differed between the ranges. Invaded soils had higher alpha diversity than native soils by both metrics (richness $P = 0.035$, Shannon $P = 0.002$, t -test; Figure 2a-b). In terms of beta diversity, both range (native or invaded) and site had a significant effect on community membership and community structure (all $P < 0.0001$, PERMANOVA; Table 1; Figure 3).

We found that the distances between the bulk soils used as inocula and the soil samples from the transects at same site were significantly smaller than the distances between the bulk soils and soil samples from external sites (unweighted unifrac $P = 0.005$, weighted unifrac $P = 0.0001$, paired t -test, Figure S3). In other words, bulk soils used as inocula in the greenhouse experiment were more similar to the soil of the site they were from than to any of our other sites, and were thus representative of those sites for use in and interpretation of our greenhouse experiments.

Experimental inoculation: Rhizosphere

Across our rhizosphere samples we identified 23 phyla and 352 families of bacteria. Eight phyla made up 99% of sequences present (Figure 1, Table S5). The most abundant phyla were Proteobacteria (an average of 70% across samples, SD = 12%), Bacteroidota (13%, SD = 9%), and Actinobacteriota (10%, SD = 6%). There were no phyla that differed between native and invading *C. solstitialis* genotypes within either native or invaded soil treatments, or between native and invader plant genotypes overall (all LDA effect sizes <2, LEfSe).

At the bacterial family level, the interaction of the source of soil microbial communities and plant genotype shaped rhizosphere communities. For invaded range soil inocula, seven bacterial families were differentially abundant, including 6 families that were more abundant on native range plants and one that was more abundant on invaders (LDA effect size > 2, LEfSe; Figure 4). For native range soil inocula, five families differed, with two families more abundant on native range plants and three that were more abundant on invaders (LDA effect size > 2, LEfSe; Figure 4). Only the family Micropepsaceae was differentially abundant between plant genotypes in both native and invaded soil treatments, in which it was more abundant on native genotypes in both treatments.

Contrary to patterns in field soils, alpha diversity (richness and Shannon's H) was higher for bacterial communities on native *C. solstitialis* genotypes than for communities on invader plant genotypes (Figure 2c-d, Table S6). There was not a significant difference in alpha diversity between communities derived from native vs. invaded range soils, nor an interaction between soil range and plant genotype (Table S6).

DISCUSSION

Our goal in this study was to test for the effects of plant evolution and its interaction with the soil microbial environment on plant-microbial associations during invasion. In contrast to previous hypotheses that the invaded environment (i.e. Enemy Release) is responsible for shaping microbiome associations for *C. solstitialis* genotypes invading California, we found that both plant genotype and microbial environment interacted to shape microbial family associations in this invasion. We also found that previously observed patterns of plant-associated microbial diversity in this invasion [69] were not a product of microbial communities available in the environment as hypothesized. Instead, plant genotype drove the pattern of its microbiome diversity, such that invading plant genotypes accumulated a lower diversity of rhizosphere bacteria than native genotypes, despite invaded range soils being overall higher in microbial diversity.

Amplicon sequencing of field soils indicated that bacterial community environments differed significantly between ranges, and among sites within ranges. Differences between native and invaded microbial environments were evident at both the phylum and family levels. For both taxonomic levels most differentially abundant taxa were higher in the invaded range soils than the native range soils (e.g. 55 families were more abundant in the invaded range, versus 19 that were elevated in the native range). Consistent with these differences, alpha diversity was higher in the invaded range soils for Shannon diversity, and marginally higher for species richness. Soil microbial diversity and composition are known to vary geographically for a variety of reasons, including soil physical characteristics (particularly pH), climatic factors, land use and disturbance, and plant species composition [77, 89, 90]. Our collection sites were

typically roadsides adjacent to agricultural fields, with cover of *C. solstitialis* and European grasses, and climatically similar environments [73], in both ranges. Nevertheless, significant side effects within each range indicate that microbial communities were sensitive to subtleties of site characteristics, and the distribution of some microbial taxa (see below) suggest that native range sites might experience more drought stress.

Using these same soils as inocula in our experiment, we found that invading genotypes associated with a lower diversity of bacteria than did native genotypes, regardless of the source of the soil inoculum. This indicates that plant genotypes in this system shape the microbial communities on their roots, and that invaders are not only experiencing novel interactions, their evolution is shaping these interactions. In their observational study of *C. solstitialis*-associated microbial communities in the field, Lu-Irving and colleagues (2019) noted that rhizosphere microbial alpha diversity was positively correlated with the genetic diversity (genetic variation) among plants with a site, within each range, consistent with an important effect of different genotypes accumulating a diversity of different microbial taxa within field site. This relationship could not explain differences in microbial diversity between ranges, however, because plant genetic diversity within sites did not differ substantially between the native and invaded range in this system [69, 73]. Our experiment reveals that it is divergence between native and invading plant genotypes, rather than differences in their population genetic diversity within sites, that explains lower rhizosphere diversity on plants in the invasion.

We found no effect of soil community environment on alpha diversity of root-associated microbes in our experiment, but we did find effects of both soil environment and plant genotype on the abundance of specific taxa in the rhizosphere. Below we discuss known

functions of several of these taxa, but we emphasize that their relationships to *C. solstitialis* are unknown in all cases and must be investigated further. While it is difficult to generalize the functions of many bacterial taxa (especially at higher taxonomic ranks), and some are entirely unknown, current understanding of these groups suggests that additional research on their functional roles in this system would be informative.

In possibly our most important finding regarding differentially abundant taxa, we found that Streptomycetaceae were more abundant in invaded range field soils, and were more abundant on invader genotypes when grown in the native range soils (in which this group was less abundant, suggesting preferential accumulation). This association between Streptomycetaceae and the invaded range, and the apparent accumulation of these bacteria on invading genotypes, is potentially important because the Streptomycetaceae are widely associated with the promotion of plant growth [91]. Their benefits appear to be both direct effects on growth (as ‘biofertilizers’) and indirect effects through their suppression of plant pathogens (the Streptomycetaceae are also responsible for the majority of human antibiotics) [92]. If members of this family have a similar positive relationship with invading *C. solstitialis*, this could explain several observed advantages to invaders in this system, including increased invader growth [62], reduced negative plant-soil interactions in the invaded range [30], and potentially the decrease in diversity of microbes on invader roots, given the antimicrobial properties of Streptomycetaceae [69].

Native genotypes also had a set of interactions that mirrored the invader relationships with Streptomycetaceae. These were taxa that were more abundant in native soils, and also more abundant on native genotypes when grown in the invaded soils in which these same

groups were more depauperate (again suggesting preferential accumulation). These groups have a variety of potential functions of interest, and include an uncultured Gaiellales family (a group that has been associated with plant colonizers, and may regulate organic and fatty acids in soils [93, 94]); a family in Subgroup 1 of the Acidobacteriaceae (a diverse group of oligotrophs abundant in soils, some of which have adapted to specific dry soil conditions; [95, 96]), a family in the CPla-3 termite group (a more highly specialized group in the soil, often in acidic environments; [97]), and the family Polyangiaceae (a group of predatory social bacteria, among the most common predators of bacteria in the soil, with the potential to structure the diversity of the entire microbial community [98–100]).

The most notable association for native plant genotypes, however, appeared to be with the family Micropepsaceae, which was more abundant in invaded range soils, but was overrepresented on native genotypes grown in either invaded or native range soils. This was the only family to be overrepresented on a genotype across both soil environments. The Micropepsaceae has been found to be an indicator taxon associated with specific plant species or site conditions in other studies [101, 102]. For studies that compared different environmental manipulations, the family was associated with early phases of restoration (grasslands [103]), early responses to agricultural planting (tobacco [104]), and response to cold treatments of plants (lettuce [105]), all conditions in which a system had been perturbed recently. In one study, the presence of Micropepsaceae was also associated with reduced suppression of pathogens [106]. If native genotypes are exposed to more pathogens when associating with this family (perhaps during resource acquisition after disturbance), this might favor diversion of resources from growth into defenses, which has been hypothesized to explain the smaller size

of native plant genotypes within *C. solstitialis* [69].

The association patterns of two families suggest that the native environment might be more stressful for plants, particularly in terms of drought. While native and invaded range climatic niches are generally similar in these areas [73], the families Bacillaceae and Terrimicrobiaceae were more abundant in native range soils. Bacillaceae is characterized by the ability to form endospores that are resistant to environmental extremes, including drought [107]. The Terrimicrobiaceae have been observed to be enriched for G3P (Glycerol-3-phosphate) transport-related genes, which are associated with plant-microbe interactions that promote drought tolerance of plants [108]. Both families were also more abundant on invader genotypes in native soils, which is notable because previous research has found that invader genotypes are less drought tolerant, and have suggested that this might be adaptive because invaders occupy an environment with lower competition for limiting resources, primarily summer water, and therefore experience lower water stress in the invasion [62].

Finally, two groups that are more common in invaded range soils could be of interest for their differential associations with each genotype growing in that soil inoculum. Taxa in the Armatimonadales (a group of putatively generalist consumers of organic compounds [97, 109]) In a comparison of patches invaded by *Amaranthus palmeri* to patches of native plants in China, Armatimonadales were one of a small number of families strongly associated with the invader and predictive of soil functional characteristics [110]. Taxa in the Chthonomonadales (a group associated with decomposition of glycine substrates; [111]) were more abundant on invader genotypes in our study. Glycine rich proteins can be a major component of plant cell walls [112] and associated decomposers may reflect greater nutrient cycling in the presence of higher

biomass input [111].

Further work is needed to identify how *C. solstitialis* interactions with these specific bacterial families that we have identified, as well as other components of their microbial communities (i.e. fungi and viruses), affect host plant fitness across ranges. Microbial interactions could be responsible for natural selection that has driven evolution of invaders in this system, given that root microbiomes can have large effects on host plant performance [113, 114]. A study of 15 annual plant species in California grasslands (not including *C. solstitialis*), found that soil microbial communities generated large fitness differences among species, suggesting that microbially-driven selection could be strong [115].

Invaders will also evolve to adapt to other aspects of their new environments, and are expected to evolve increases in traits associated with invasiveness itself [50–52, 62, 116–119]. Such trait changes could be synergistic with more favorable microbial interactions, for example the evolution of increased root investments for resource acquisition could also help plants take advantage of beneficial microbes [120]. The influence of host evolution on microbial interactions during invasion does not appear to have been investigated previously, however. There is some evidence that plant-microbial interactions have changed over time during invasion, which could result from plant evolutionary change. For example, older invading populations of *Solidago canadensis* were found to have increased positive microbial interactions and competitive ability in a common garden experiment, which suggests that these changes were genetically-based [121]. In contrast, the evolution of aboveground herbivore interactions in plant invasions has attracted longstanding interest [12], and recent reviews find abundant evidence for the evolution of these interactions, though not consistently to the advantage or

disadvantage of the invader [122, 123]. Our work demonstrates that belowground microbial interactions should also be expected to evolve during invasion, as a product of both the resident microbial community, and the genetic composition of the host. How these interactions alter fitness and invasiveness over time, and how they interact with available genetic variation in introduced populations, will be important avenues of further research.

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Table 1 - PERMANOVA results for two measures of beta diversity (weighted and unweighted UniFrac).

predictor	df	Pseudo-F	ω^2	P
<i>unweighted UniFrac:</i>				
Range	1	5.48	0.06	0.0001
Site with/in range	5	5.49	0.19	0.0001
<i>weighted UniFrac:</i>				
Range	1	10.15	0.14	0.0001
Site with/in range	6	12.51	0.37	0.0001

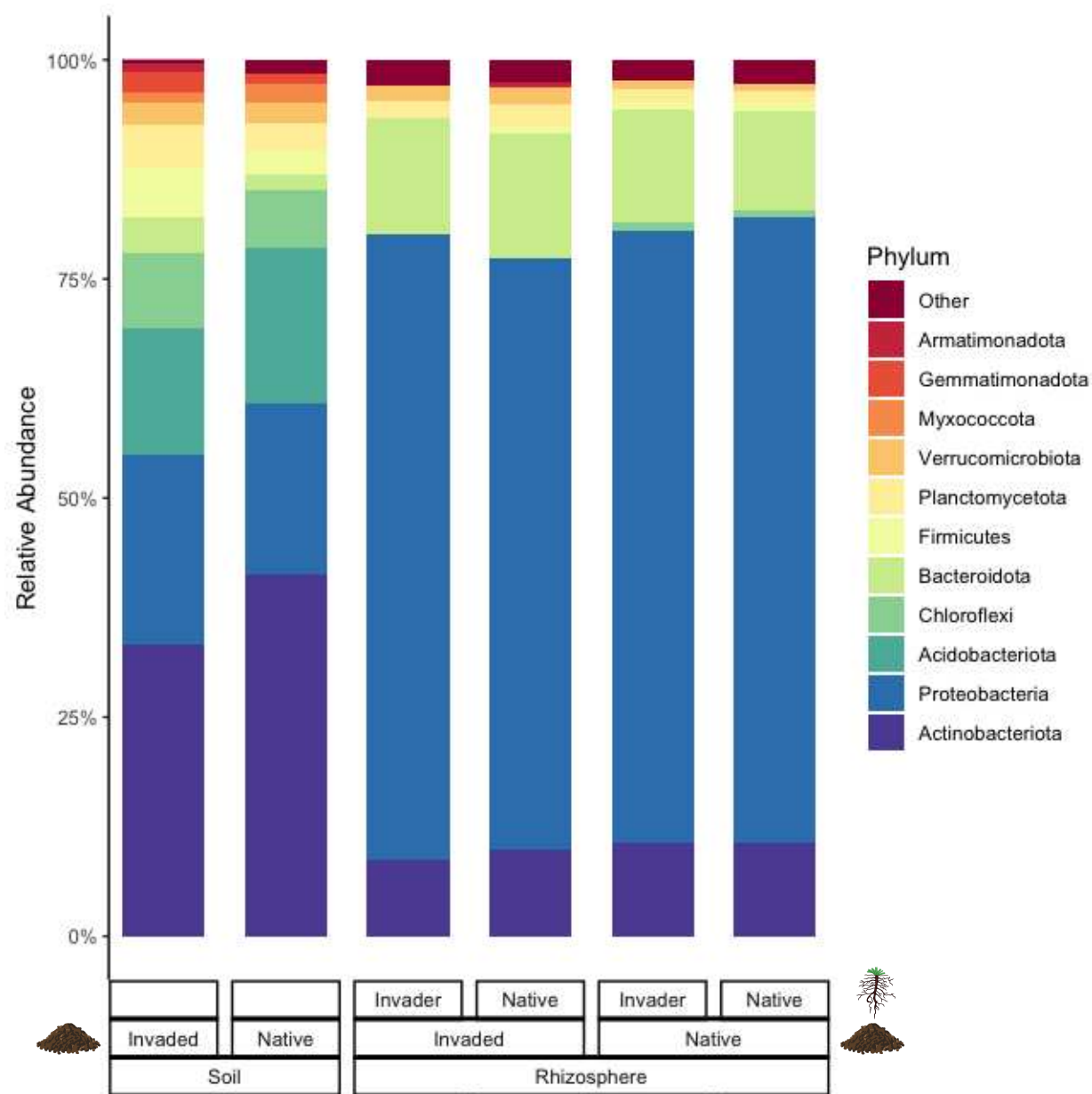


Figure 1 - Relative abundance of bacterial phyla in field soil samples, grouped by range (invaded or native) and rhizosphere samples (root wash) grouped by plant genotype (invader or native) within range. All phyla shown constitute at least 1% of at least one sample for soil, and at least 0.6% of at least one sample for rhizosphere. Relative abundance of each phylum is average across replicates for each group.

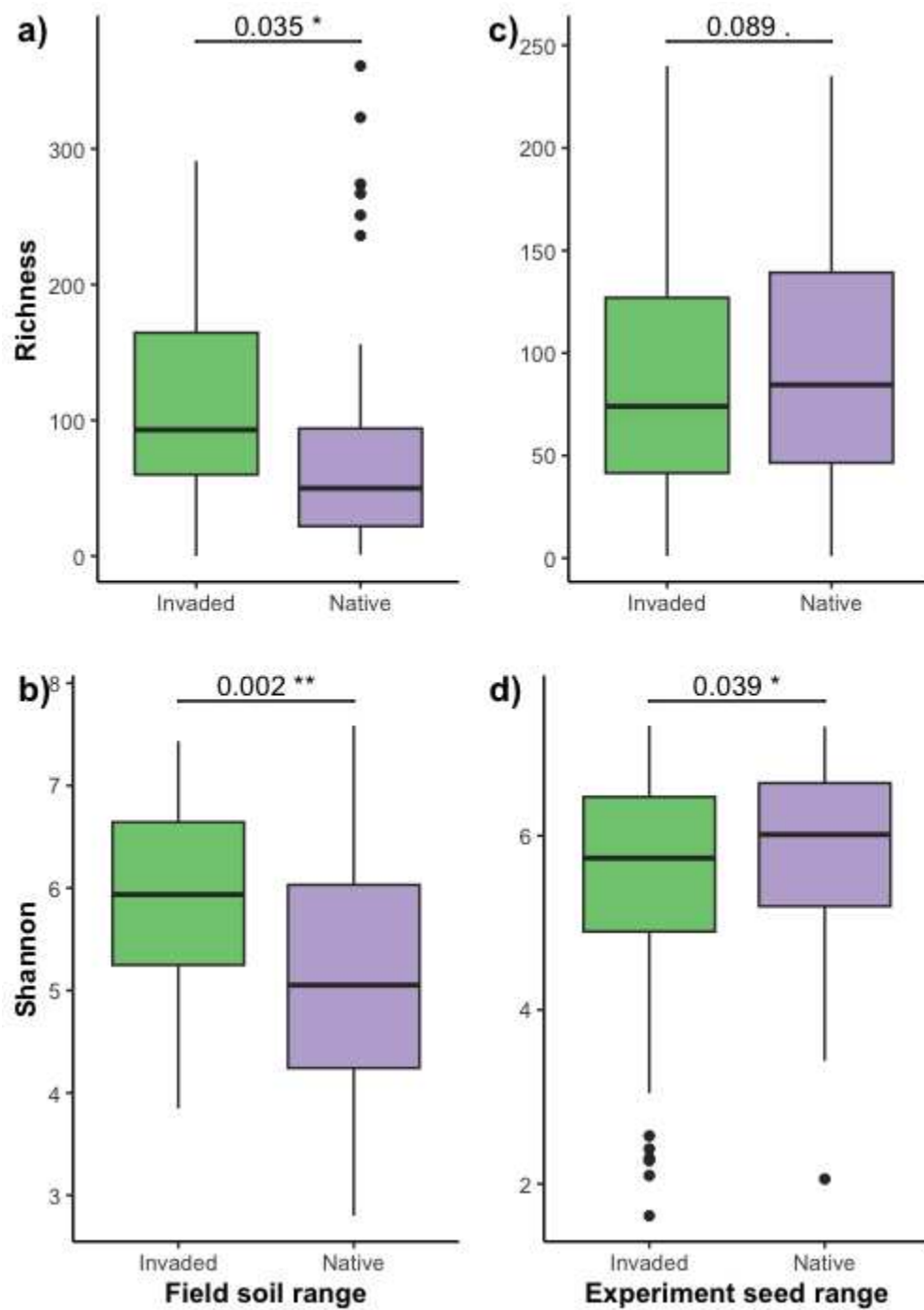


Figure 2 - Box plots showing two measures of alpha diversity (a & c richness, b & d Shannon's

diversity index) for field soil samples (a & b) in invaded range soils (California, USA) and native range soils (Spain and France) and for native and invader yellow starthistle (*C. solstitialis*) genotypes grown (c & d).

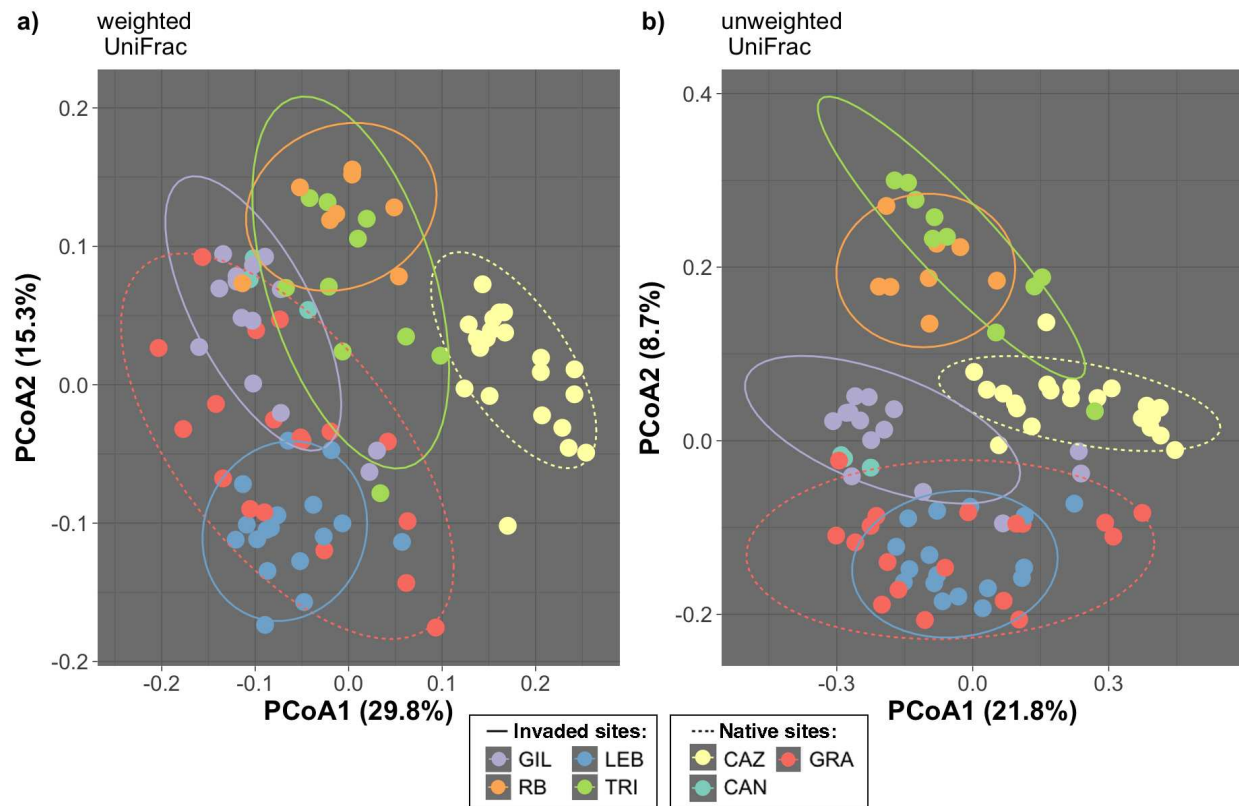


Figure 3 - Principal coordinates ordination of **a)** weighted UniFrac distances (community structure) and **b)** unweighted uniFrac distances (community membership) of field soil samples. Variation explained by each axis is shown in parentheses. Each data point represents one soil sample. Ellipses represent 90% confidence intervals. Solid line ellipses represent invaded sites, and dashed line ellipses represent native sites.

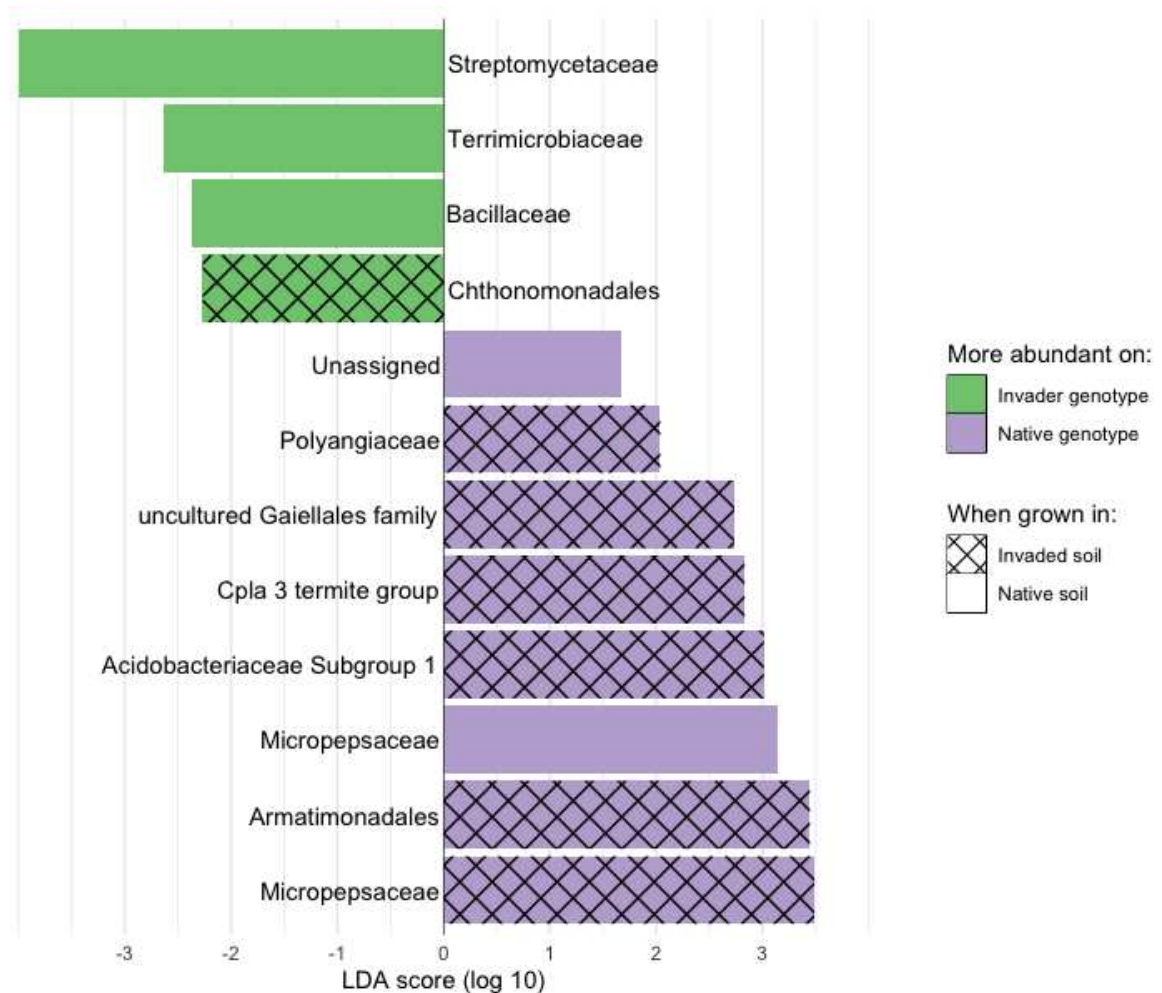


Figure 4 - Bacterial families that are differentially abundant in rhizosphere bacterial communities between invader and native *C. solstitialis* genotypes when grown with invaded and native microbial communities. Shown are Linear Discriminant Analysis (LDA) scores from LEfSe analyses comparing rhizosphere microbial communities between native and invading plant genotypes. Solid color bars are from LEfSe comparing microbial communities of each genotype when grown in native soil, hatched bars for when grown in invaded soil. Note: Micropepsaceae was the only family found to be differentially abundant between genotypes in both invaded and native soil.

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