Realistic time-lags and litter dynamics alter predictions of plant-soil feedback across generations

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1 Summary

Plant-soil feedback is a critical process in natural plant communities. However, it remains
 unclear whether greenhouse-measured microbial effects manifest in natural systems with
 temporally separated growing seasons as classic experiments often overlook seasonal time
 lags and litter dynamics.

We modified the classic two-phase experiment to study plant-soil feedback for three Cali fornian annual plant species. Our response phase used soil inoculum obtained either im mediately after plant conditioning, after a six-month dry period with the conditioning plant
 removed, or after a dry period with the litter of the conditioning plant. We characterized soil
 bacterial and fungal communities in different treatments and employed recent advancement
 in plant-soil feedback theory to predict plant coexistence.

Temporal delays and the presence of litter caused distinct responses in the fungal and bacterial communities, resulting in divergent microbial compositions at the end of the response phases.
 The delayed response treatments also affected microbially mediated stabilization, fitness differences, and invasion growth rates differently across species pairs, influencing predictions of plant coexistence.

Our study highlights that the interplay between seasonal delays and litter dynamics prevents
 the direct extrapolation of plant-soil feedback measurements across multiple seasons, emphasizing the necessity of considering natural history when predicting microbially mediated
 plant coexistence.

21 Keywords

Annual plants, Invasion growth rate, Litter decomposition, Microbial community, Modern coexis tence theory, Natural history, Seasonality

24 Introduction

The interactions between plants and soil microbes have gained increasing recognition as a pivotal 25 force in shaping plant communities (Bever et al., 2010, van der Putten et al., 2013). The effects of 26 these interactions on plant community dynamics are most commonly studied under the plant-soil 27 feedback (PSF) framework, which captures the effects of bidirectional interactions in which plants 28 simultaneously alter and are affected by the soil microbial community (Bever et al., 1997). To 29 implement this framework in empirical studies, PSF is often quantified through two-phase experi-30 ments that separate the feedback process into distinct "conditioning" and "response" phases (Bever 31 et al., 1997, 2012). Plant performances during the response phase are measured to predict how 32 soil microbes influence plant coexistence (Crawford et al., 2019, Yan et al., 2022). However, despite 33 a vast body of literature showing that soil microbes can exert strong controls over plant species 34 coexistence, connecting the predictions from such two-phase studies to the observed dynamics of 35 plant communities in nature remains challenging (Forero et al., 2019, Beals et al., 2020, Beckman 36 et al., 2022, Png et al., 2023). A promising approach for addressing this challenge is to adopt the 37 classic two-phase design to better reflect the natural conditions under which PSF arises in the field 38 (Gundale & Kardol, 2021). 39

Greenhouse experiments of plant-soil feedback typically simplify the temporal dynamics of 40 feedback by conducting the conditioning and response phases sequentially, without any temporal 41 separation between them. While this design likely captures the effects of microbial feedback among 42 plants growing concurrently, whether these same effects manifest in communities characterized 43 by temporally separated plant growing seasons or in communities where time-lags occur between 44 soil conditioning and its subsequent recolonization is less clear. For example, Esch & Kobe (2021) 45 found that in a temperate hardwood forest, Prunus serotina adults cultivate a soil community that 46 suppresses the growth of conspecific seedlings, but this suppressive effect erodes within months 47 of plant death. Thus, the long-term consequences of soil conditioning are unclear if there are time 48 lags between adult death and subsequent arrival/growth of seedlings in the conditioned soils, 49 which is especially likely in plant communities characterized by dispersal and/or seed limitation 50 (Ehrlén & Eriksson, 2000). Similarly, in systems where plant dynamics are highly seasonal, the 51 conditioning effects that build up during one growing season may not translate directly to affect 52

plants in subsequent growing seasons if the soil community is reshaped during the intervening 53 period (Barnard et al., 2013). Such dynamics are likely to be especially relevant in Mediterranean-54 type annual plant communities frequently used in PSF experiments (e.g., Bonanomi et al., 2012, 55 Siefert et al., 2019, Kandlikar et al., 2021), where winter growing seasons are punctuated by dry 56 summers of plant senescence (Elmendorf & Harrison, 2009). Furthermore, recent theoretical 57 studies have demonstrated that the temporal dynamics of plant-soil feedbacks can substantially 58 alter predictions of microbially mediated plant coexistence (Ke & Levine, 2021, Miller & Allesina, 59 2021). Thus, both empirical and theoretical evidence suggests that incorporating the natural 60 temporal dynamics of plant communities into studies of plant-soil feedback might enable more 61 robust predictions of how soil microbes shape plant coexistence in nature. 62

Another aspect of the soil conditioning process that is largely overlooked in two-phase plant-63 soil feedback experiments is that, in nature, the soil microbial community is shaped not only by the 64 active conditioning effects of plants as they grow but also by the dead tissue (i.e., litter) that plants 65 deposit onto the soil. Specifically, recent literature has shown that plant litter of different species 66 can influence microbial communities by introducing phyllosphere microbes to the soil (Whitaker 67 et al., 2017, Fanin et al., 2021, Minás et al., 2021) and by releasing chemicals and nutrients that 68 affect soil microbial community assembly (Veen et al., 2021). These litter-induced changes in the 69 microbial community can subsequently result in different plant-soil feedback on the responding 70 plants (Veen et al., 2019, Aldorfová et al., 2022). For example, in systems with distinct phenology or 71 seasonality, using soil collected at the end of the growing season rather than after decomposition 72 would fail to capture the full impact of litter dynamics. Despite the role of litter dynamics in 73 shaping soil communities in all systems, this process is largely overlooked in plant-soil feedback 74 experiments, which typically remove all plant material at the conclusion of the conditioning phase. 75 Incorporating the role of litter in plant-soil feedback is thus an important step for bridging the gap 76 between classic experiments and natural conditions. 77

To better predict the long-term consequences of plant–soil feedback in natural systems, we also need theoretically robust metrics to extrapolate greenhouse experimental results. The original theory of plant–soil feedback popularized a pairwise feedback metric that quantifies how soil microbes drive frequency-dependent stabilization (e.g., via host-specific pathogens; Bever *et al.*, 1997,

Eppinga et al., 2018). Recent theoretical advances have integrated plant-soil microbe interactions 82 with modern coexistence theory (Kandlikar et al., 2019, Ke & Wan, 2020), which utilizes invasion 83 growth rates to predict species coexistence (i.e., quantifying whether each plant can establish in its 84 competitor's monoculture equilibrium from low density; Turelli, 1978, Chesson, 2000). Specifically, 85 plant coexistence requires the stabilizing effects of microbes to overcome microbially mediated fit-86 ness differences, with the former capturing how microbes benefit both plants by driving negative 87 frequency dependence while the latter capturing how microbes disproportionately impact one 88 plant species over the other (Kandlikar et al., 2019, 2021, Yan et al., 2022). Evaluating coexistence 89 outcomes on the basis of species' invasion growth rates can also yield important insights for eluci-90 dating the underlying interactions in experimental data (Grainger et al., 2019, Ke & Wan, 2020, 2023). 91 Examining the impact of experimental manipulation through these theoretical metrics enables a 92 more nuanced understanding of the pathways through which plant-soil feedback influences plant 93 coexistence. 94

Here, we conducted an experiment to address two questions about the role of soil microbes 95 in shaping plant coexistence in annual grasslands: (1) How do seasonal time lags and plant lit-96 ter decomposition interact with the conditioning process to alter the soil microbial community? 97 (2) How do these changes to the soil community scale up to impact the predicted consequences 98 of plant-soil feedback? To address these questions, we modified the two-phase greenhouse ex-99 periment and conducted three fully factorial response treatments. These treatments used soil 100 inoculum obtained either immediately after plant conditioning, after a six-month dry period time 101 lag with the removal of the conditioning plant, or after a similar dry period with the litter of the 102 conditioning plant left intact. We quantified the absolute abundance of soil bacterial and fungal 103 communities at the end of the conditioning and response phases, enabling us to evaluate how the 104 soil inocula from each response treatment triggered different microbial communities. We then 105 employed modern coexistence theory to predict the consequences of plant-soil feedback based 106 on microbially mediated stabilization, fitness difference, and invasion growth rates (Kandlikar 107 et al., 2019). Our results demonstrated that both time lag and plant litter altered the outcome of 108 plant-soil feedback, with varying effects across species pairs. This work underscores the need to 109 incorporate natural history when predicting microbially mediated plant coexistence. 110

Methods

112 Study system

We focused on three native Californian winter annual plants: a legume Acmispon wrangelianus 113 (ACWR; Fabaceae), a grass Festuca microstachys (FEMI; Poaceae), and a forb Plantago erecta (PLER; 114 Plantaginaceae). In spring 2019, we collected seeds from the University of California Sedgwick 115 Reserve in Santa Barbara County, California, USA (34°41' N, 120°02' W), where all three species 116 co-occur. In this Mediterranean-type climate, annual plants complete their life cycle and senesce 117 in the hot, dry summer lasting about six months (May-October mean temperature = 18.9°C, 118 mean monthly precipitation = 4.57 mm; data from 2014–2023). The new generation germinates 119 following rain in the cool, wet winters (November-April mean temperature = 12.3°C, mean monthly 120 precipitation = 54 mm). In September 2020, prior to winter rains, we collected field soil from 121 Sedgwick Reserve to serve as microbial inoculum. To ensure that the field microbial community 122 was not pre-conditioned by species in our experiment, we collected soil from four locations where 123 there were no individuals of our focal species in a 1 m radius. The soils were kept at 4° and 124 transported to the lab within 12 hours, where equal amounts of soil from each location were sifted 125 through a 2 mm sieve and homogenized. Prior to the experiment setup, we subsampled the field 126 soil and stored it at -80° for later DNA sequencing of the microbial community. One fraction of the 127 field soil was then used to inoculate the conditioning phase pots, and the remainder was stored at 128 0°C until further use in the response phases as a reference soil treatment. 129

¹³⁰ Greenhouse experiment and soil sampling

¹³¹ We modified the classic two-phase experiment to study how seasonal time lag and plant litter ¹³² affect the soil microbial community and plant competitive outcomes. Specifically, our growth ¹³³ chamber experiment consisted of three fully factorial response treatments, using soil inocula that ¹³⁴ went through different handling treatments to represent these natural history factors (Fig. 1). We ¹³⁵ collected soil samples at different stages of the experiment and characterized the microbial commu-¹³⁶ nity by high-throughput sequencing. Plant competitive outcomes were predicted by measuring ¹³⁷ plant biomass performance at the end of the experiment.

138 Conditioning phase

To cultivate soil microbes associated with each species, we grew three high-density monocultures 139 $(8 \text{ g viable seed}/\text{m}^2)$ of each species in bleach-sterilized 1-gallon pots (Fig. 1). We first filled each pot 140 with 2.60 L sterilized potting mix (equal parts sand, clay, peat, perlite, and vermiculite; autoclaved 141 twice, each 2 hours with a 24-hour resting period in between). We then added 0.30 L of field soil to 142 each pot and topped it with a 0.10 L layer of sterilized potting mix to achieve a 10% volume of live 143 inoculum. Into each pot, we sowed 0.141 grams of seeds of a single species, which we had surface-144 sterilized by soaking in 1% bleach for 2 minutes and washing with ultrapure water twice for 1 145 minute each. We stored pots at 4°C for five days to trigger germination, after which we moved pots 146 to a growth chamber (25°C, 60% humidity, 10:14 hour day:night cycle) for 80 days, approximately 147 the length of a complete growing season. In addition to the 9 large conditioning pots, we grew 148 10 replicate individuals of each species in sterilized potting mix to serve as phytometers between 149 the different phases of the experiment (3 species \times 10 replicate individuals = 30 pots). We rotated 150 control plants (30 pots) and conditioning monoculture pots (9 pots) weekly within the growth 151 chamber. 152

The conditioning phase of the experiment concluded in December 2020. At this time, we 153 randomly chose soil from one monoculture pot for each species to serve as the inoculum source 154 for the "immediate" response treatment (green pots in Fig. 1). We designated the remaining two 155 monoculture pots per species for the two time-lagged response treatments, and left these in the 156 growth chamber (25°C, 10% humidity, 10:14 hour day:night cycle) for an extra six-month dry period 157 to mimic the temporal gap between two consecutive seasons. From one of these, we removed all 158 aboveground biomass of the conditioning plant (grey pot in Fig. 1), whereas in the other we left 159 all plant tissue intact (brown pot in Fig. 1). Thus, for each species, we were able to evaluate 160 the effects of soil conditioning on subsequent plant growth without any time lag ("immediate" 161 treatment), and could also evaluate how the presence of litter interacts with time lags to affect 162 the plant performance during the "delayed" response phase (Fig. 1). Before using the conditioned 163 monoculture pots for their corresponding response phase, we collected soil samples from each pot 164 to characterize how seasonal time lag and plant litter influenced the soil microbial community (see 165 section DNA sequencing of the microbial community). 166

167 Response phase

To create soil inocula for the "immediate" response phase, we removed the aboveground biomass 168 from one conditioning monoculture pot per species, and sifted the soil through a 2 mm sieve to 169 remove roots and homogenize the soil. We autoclaved half of this soil to create the sterilized 170 inocula; the other half served as the live inoculum. We grew 10 replicate individuals of each 171 species with either live or sterilized inoculum from each of the three species' freshly conditioned 172 monoculture pot (i.e., 3 plant species \times 3 soil inoculum \times 2 sterilization treatments \times 10 replicates 173 = 180 pots). These plants in immediate response treatment correspond to the typical experimental 174 procedure in plant-soil feedback experiments, which use soil inoculum collected at the end of the 175 conditioning phase. To quantify the impact of time lag and litter presence, we started another 176 round of response phase (i.e., the two delayed treatments) in June 2021 using soil inoculum that 177 experienced an additional six months of dry period after the conditioning phase, thereby more 178 closely mimicking the natural temporal dynamics of these grasslands. For these treatments, we 179 grew 10 replicates of each species with live and sterile inoculum from the corresponding "delayed" 180 and "delayed with litter" monoculture pots of each of the three species (i.e., 3 plant species \times 3 soil 181 inoculum \times 2 delay treatments \times 2 sterilization treatments \times 10 replicates = 360 individuals). 182

For the response phase pots, we filled 125 mL Deepots (Stuewe & Sons, Inc.) with sterilized 183 potting mix and 10% volume of soil inoculum, and covered the inoculum with a thin layer of steril-184 ized potting mix. We surface-sterilized and pre-germinated seeds, and transplanted seedlings into 185 the pots so that each pot had a single individual. As evaluating microbially mediated coexistence 186 outcomes requires growing plants in a reference soil (sensu Kandlikar et al., 2019), we also grew 187 10 replicate individuals of each species using unconditioned field soil as inoculum (previously 188 collected and stored at 0°C; blue pots in Fig. 1). To control for batch effects between the two 189 rounds of response phases (i.e., the immediate response treatment in December 2020 and the two 190 delayed response treatments in June 2021), during each round we grew 10 replicate individuals 191 of each species with sterilized potting mix (i.e., batch effect controls with no inoculum). In total, 192 the response phase of our experiment included 660 pots (i.e., 180 pots for the immediate response 193 treatment + 360 pots for the two delayed response treatments + 3 plant species \times 10 replicates \times 2 194 rounds of field reference soil treatments + 3 plant species \times 10 replicates \times 2 rounds of batch effect 195

controls). We grew the plants in a growth chamber for 80 days (same conditions as the conditioning phase, rotated weekly), after which we harvested, dried (72 hours at 60°C), and weighed plant
aboveground biomass. At the end of all response phases, we collected soil samples from each pot
to characterize the soil microbial community.

²⁰⁰ DNA sequencing of the microbial community

As described earlier, we collected soil samples from conditioned monoculture pots (before using 201 them as soil inoculum) as well as response phase pots at the end of the experiment. The former was 202 meant to characterize how seasonal time lag and plant litter affected the soil microbial community, 203 while the latter was meant to see if these changes in the inoculum triggered long-lasting impacts 204 (Hannula *et al.*, 2021). We mixed soils within each pot well and subsampled 0.25 g of soil. To 205 each sample, we added 8 ng of P and F synthetic chimeric DNA spike for the quantification 206 of prokaryotic and fungal absolute abundance prior to DNA extraction (Tkacz et al., 2018). We 207 extracted DNA using Qiagen DNeasy PowerSoil Pro Kit according to the manufacturer's manual, 208 with a 65°C water bath incubation (10 minutes) prior to bead-beating to improve yield. Using two-209 step PCR, we amplified the V4 region of bacterial 16S ribosomal RNA gene (Caporaso et al., 2012) 210 and fungal internal spacer 1 region (ITS1) (White et al., 1990) with index primers. We purified, 211 normalized, and pooled amplicon libraries and sequenced using 2×300 bp paired-end Illumina 212 MiSeq (see Supporting Methods S1). 213

214 Data analysis

215 Microbial community

We converted raw binary base call (BCL) files to fastq files and demultiplexed with Illumina Bcl2fastq2 (v2.20). We trimmed adapter sequences from reads using cutadapt (Martin, 2011) with python (v3.10.9). Using the Divisive Amplicon Denoising Algorithm (dada2 v.1.28.0) in R (v4.3.0), we quality filtered and trimmed the fastq files, and inferred amplicon sequence variants (ASV) (Callahan *et al.*, 2016a) following the published workflow (Callahan *et al.*, 2016b). Specifically, we discarded low-quality ends of reads by trimming the bacterial forward reads to 250 bp and the reverse reads to 210 bp, discarding any reads shorter than these lengths. We chose not to trim

fungal read lengths due to the varying size of the ITS gene region. We used phyloseq (McMurdie & Holmes, 2013) for downstream analysis. We discarded any ASV that was only detected in ≤ 5 samples; we also removed samples with extremely small or large read counts (i.e., more or less than 5x the average number of reads across all samples). We rarefied samples to 5000 sequencing reads for downstream analyses.

We transformed the community matrix from raw reads to relative abundance. We calculated 228 absolute abundance as in Tkacz et al. (2018). Briefly, we divided the total number of environmental 229 reads by the number of synthetic reads in each sample and then multiplied it by the number of 230 gene copies in the 8 ng of synthetic spike-in, which was calculated by multiplying the number 231 of gene copies in 1 ng of spike-in (i.e., 3.5E+07 for 16S and 1.2E+07 for ITS, respectively) by 232 eight. To understand the starting soil microbial species pool that plants experience in the response 233 phase, we explored differences in absolute abundances of 16s and ITS ASVs of the conditioned 234 soil (immediately after the conditioning phase, after the six-month delay without litter, after the 235 six-month delay with litter, and field reference soil). We first agglomerated the sequences in 236 the conditioned soil samples to the class taxonomic level using tax_glom phyloseq function, and 237 then visualized the total count for each class present in the samples. We identified compositional 238 differences (i.e., beta diversity) with the Bray-Curtis dissimilarity metric and compared with a 230 permutational multivariate analysis of variance using the vegan (Oksanen et al., 2017) and stats R 240 packages. 241

242 Plant biomass performance and competitive outcomes

To test for differences in plant biomass when grown with conspecific-conditioned versus reference 243 soil (i.e., unconditioned field soil and sterilized potting mix) microbes, we conducted a series 244 of linear models with log-transformed biomass values as the response variable, and soil source 245 as the predictor. We fit separate models per species (ACWR, FEMI, and PLER) and treatment 246 (immediate, delayed with litter, delayed without litter) to facilitate model interpretation. Prior to 247 biomass analyses, we filtered out outliers within each experiment phase \times species \times soil \times plant 248 combination. Outliers were identified as individuals with biomass lower than Q1-1.5*IQR or higher 249 than Q3+1.5*IQR, where Q1 and Q3 were the 25th or 75th quartiles, respectively, and IQR represents 250 the difference between Q1 and Q3. We evaluated statistical significance at $\alpha = 0.05$. 251

To predict how different response treatments modified the effects of plant-soil feedback on 252 plant coexistence, we calculated microbially mediated stabilization and fitness differences for each 253 treatment separately. We first quantified the effects of plant *j*-conditioned microbial community 254 on the biomass performance of plant *i*, denoted as m_{ij} (*i* and j = 1 or 2). This microbial effect is 255 defined as $m_{ij} = \ln(\text{biomass of plant } i \text{ in soil } j) - \ln(\text{biomass of plant } i \text{ in reference soil})$, i.e., the 256 rate of plant *i* biomass accumulation when grown in soils with the plant *j* microbial community 257 relative to that in an unconditioned reference soil. We then compare pots with conspecific versus 258 heterospecific soil inoculum to calculate microbially mediated stabilization and fitness differences 259 following the theoretical derivations in Kandlikar et al. (2019) (see also Kandlikar et al., 2021, Yan 260 *et al.*, 2022):

Stabilization =
$$-\frac{1}{2}(m_{11} - m_{12} - m_{21} + m_{22}) = \left(\frac{m_{21} + m_{12}}{2}\right) - \left(\frac{m_{11} + m_{22}}{2}\right),$$
 (1)

Fitness difference =
$$\frac{1}{2}(m_{11} + m_{12} - m_{21} - m_{22}) = \left(\frac{m_{11} + m_{12}}{2}\right) - \left(\frac{m_{21} + m_{22}}{2}\right).$$
 (2)

Here, microbially mediated stabilization (eqn. 1) quantifies how plants condition the soil to impact 261 heterospecific relative to conspecific competitors. Positive values favor coexistence, as conditioned 262 soils more negatively (or less positively) impact their hosts. Negative values (i.e., destabilization) 263 indicate that conditioned soils more positively (or less negatively) impact host plants, and can 264 drive priority effects. On the other hand, microbially mediated fitness difference quantifies how 265 conditioned microbes disproportionately impact one plant species over the other: in the form of 266 eqn. 2, a positive value indicates that plant 1 is favored by soil microbes because they benefit more 267 from mutualistic microbes and/or suffer less from pathogenic microbes, and vice versa. We can 268 thereby predict microbially mediated competitive outcomes based on these two metrics. If the 269 absolute value of fitness difference overwhelms the absolute value of stabilization, then the plant 270 with higher fitness will outcompete the other plant (i.e., plant 1 wins if eqn. 2 > 0 while plant 2 271 wins if eqn. 2 < 0). However, if the absolute value of stabilization exceeds that of fitness difference, 272 then the theory predicts coexistence if eqn. 1 > 0 and priority effects if eqn. 1 < 0. Comparing 273 eqns 1 and 2 across the three response treatments allows us to evaluate our hypothesis regarding 274 how seasonal time lag and litter decomposition alter the coexistence consequences of plant-soil 275 feedback. 276

As an alternative predictor of coexistence, we also calculated the invasion growth rate (IGR)
of each species within a competing species pair. Specifically, the IGR of species 1 when growing
in the monoculture equilibrium of plant 2 and its corresponding soil microbe, and vice versa, are as follows (Kandlikar *et al.*, 2019):

$$IGR_1 = m_{12} - m_{22},\tag{3}$$

$$IGR_2 = m_{21} - m_{11}. \tag{4}$$

If the two IGRs differ in their signs, then the species with a positive IGR will outcompete the 280 species with a negative IGR. If both IGRs are positive (i.e., eqns 3 and 4 > 0), the two species 281 are predicted to coexist as both can recover from low density; alternatively, the theory predicts 282 priority effects if both IGRs are negative (i.e., eqns 3 and 4 < 0). Furthermore, each species' IGR 283 only depends on how the resident soil microbe impacts the invader (e.g., m_{12}), relative to their 284 impact on the resident host (e.g., m_{22}). Therefore, compared to eqns 1 and 2, which are aggregated 285 metrics that incorporate the effects of both conditioned soil communities, it is easier to identify 286 the key microbial impacts that are driving the changes in competitive outcome across response 287 treatments. 288

We used a sampling approach to propagate the uncertainty when estimating m_{ij} through 289 to the predictions of plant competitive outcome (Yan et al., 2022, Terry & Armitage, 2023). Based 290 on eqns 1–4, six different biomass terms are needed to predict the competitive outcome (i.e., the 291 biomass of each plant species growing in soil 1, 2, and in the reference soil). For each sample draw, 292 we randomly sampled one value for each of the six biomass terms from a normal distribution, 293 with a mean equal to the empirical mean biomass and a standard deviation equal to the empirical 294 standard error (SE). We repeated this procedure 1000 times for each species pair and calculated 295 the stabilization, fitness difference, and invasion growth rates. This approach, compared to the 296 commonly used orthogonal error bars (e.g., Kandlikar et al., 2021), better propagates uncertainty 297 as it captures the interdependence between parameter estimations (Terry & Armitage, 2023). All 298 analyses were conducted in R (v4.3.0) (R Core Team, 2021). 290

300 Results

301 Microbial community

We first present results of the microbial community in the soil inocula, which revealed that sea-302 sonal time lag and plant litter decomposition had a clear impact on bacterial and fungal abundance 303 (Fig. 2). The bacterial abundance was highest when sequenced immediately following the con-304 ditioning phase (i.e., inoculum used for the immediate response treatment). However, for all 305 three plant species, it decreased consistently after a six-month delay period (i.e., inoculum used 306 for the two delayed treatments; (Fig. 2A). The fungal community, while of much lower absolute 307 abundance, showed the opposite trend where total abundance increased after the six-month de-308 lay period compared to that immediately after soil conditioning (Fig. 2B). The exception to this 309 trend was soil conditioned by F. microstachys without the litter intact during the delay, where no 310 fungal reads aside from the synthetic spike-in emerged after filtering. These results suggest that 311 the responding seedlings interacted with different microbial communities in the three response 312 treatments, due to the differences in microbial species pool in the soil inocula. 313

To evaluate whether different soil inocula led to divergent microbial composition, we also 314 sequenced the microbial community at the end of each response phase (Fig. 3). As absolute 315 abundances were quantified, we combined bacterial and fungal communities when analyzing 316 differences in microbial community composition. For all soil sources conditioned by different 317 plant species, microbial composition varied between response treatments (A. wrangelianus: R^2 = 318 0.211; *F. microstachys*: $R^2 = 0.223$; *P. erecta*: $R^2 = 0.210$; Reference soil: $R^2 = 0.133$; all P < 0.05). The 319 response treatment was still the significant predictor for all soil inocula when the bacterial and 320 fungal communities were analyzed separately (Fig. S1–S2). 321

322 Plant biomass

In the immediate response phase treatment, each of our focal species had lower aboveground biomass when grown with conspecific-conditioned soil microbes, relative to their growth in unconditioned field microbial community (ACWR: $F_{1,17} = 4.89, P = 0.041$; FEMI: $F_{1,16} = 10.64, P =$ 0.049; PLER: $F_{1,17} = 30.68, P < 0.001$; Fig. 4; see also Fig. S3 for all biomass results). *F. microstachys*

and *P. erecta* grew worse in soils with a conspecific soil community than in sterilized potting mix 327 (FEMI: $F_{1,16} = 7.21, P = 0.016$; PLER: $F_{1,16} = 9.53, P = 0.007$), while the opposite was true for 328 A. wrangelianus ($F_{1,16} = 4.53, P = 0.048$). Plants generally grew poorer in both delayed treat-329 ments (i.e., with/without litter present in conditioned soils during the time lag) compared to the 330 immediate treatment (Fig. 4). Specifically, in the delayed treatments, A. wrangelianus grew substan-331 tially better when inoculated with any live soil microbial community than with sterilized potting 332 mix, but its growth in conspecific-conditioned microbes was not significantly different than with 333 an unconditioned field community (Fig. 4A). In contrast, F. microstachys growth in the delayed 334 treatments was substantially lower with any live soil community than in sterilized potting mix. 335 When litter was removed at the end of plant conditioning, conspecific-conditioned soil microbes 336 resulted in lower plant biomass relative to unconditioned field microbes in F. microstachys and P. 337 erecta (FEMI: $F_{1,14} = 6.47, P = 0.02$; PLER: $F_{1,18} = 27.57, P < 0.001$). This effect was diminished 338 when plant litter was left intact after the soil conditioning phase. 339

340 Plant coexistence outcomes

The aforementioned biomass differences across the three response treatments resulted in shifts in 341 competitive outcomes for all three species pairs (Fig. 5). Although each species pair responded 342 differently, the two delayed treatments mostly decreased the strength of stabilization (i.e., destabi-343 lization), with the exception being the delayed without litter treatment for the A. wrangelianus–P. 344 erecta pair (Fig. 5C). The legume plant A. wrangelianus was predicted to outcompete F. microstachys 345 and *P. erecta* in the immediate response treatment (green points in Fig. 5B–C). However, fitness 346 differences shifted toward a direction that disfavor A. wrangelianus in the two delayed treatments 347 (grey and brown points in Fig. 5B–C). Correspondingly, the IGR of A. wrangelianus became negative 348 (Fig. 5E–F). As a result, A. wrangelianus loses its competitive dominance, exhibiting priority effect 349 with *P. erecta* and being outcompeted by *F. microstachys* in the delayed with litter treatment. 350

³⁵¹ We can also examine each species pair in more detail. For *F. microstachys* and *P. erecta*, the ³⁵² most common competitive outcome shifted from coexistence in the immediate treatment to *F.* ³⁵³ *microstachys* outcompeting *P. erecta* in the delayed with litter treatment (Fig. 5A). In addition to ³⁵⁴ destabilization, this shift in competitive outcome resulted from an increase in fitness difference in

favor of F. microstachys. The corresponding decrease in the IGR of P. erecta (Fig. 5D) indicates that the 355 shift in competitive outcome was mainly driven by changes in the soil microbes conditioned by F. 356 microstachys. For A. wrangelianus and F. microstachys, the most common competitive outcome shifted 357 from A. wrangelianus dominance in the immediate treatment to F. microstachys dominance in the 358 two delayed treatments (Fig. 5B). This change in competitive outcome mostly resulted from the flip 359 in the competitive hierarchy between the two plants (i.e., a decrease in fitness difference in favor of 360 *F. microstachys*). A corresponding flip in the sign of the two species' IGR can also be seen in Fig. 5E. 361 For A. wrangelianus and P. erecta, the dominance of A. wrangelianus is the most common competitive 362 outcome in the immediate treatment (Fig. 5C and E). The pair shifted towards coexistence (i.e., both 363 plants have positive IGR) in the delayed treatment, but destabilization strengthened and resulted 364 in priority effect (i.e., both plants have negative IGR) in the delayed with litter treatment. 365

366 Discussion

Typical two-phase plant-soil feedback greenhouse experiments grow the responding plant im-367 mediately after soil conditioning in the greenhouse (Brinkman et al., 2010). When transplanted 368 immediately, the predicted effects of soil microbes on species coexistence were consistent with 369 those of a previous study using the same system — the legume A. wrangelianus benefited from 370 microbially mediated fitness advantage and was predicted to outcompete the other two species 371 (Kandlikar et al., 2021) (light green points in Fig. 5). Yet during naturally occurring time lags of 372 discrete growing seasons, the microbial community originally conditioned by plants may shift 373 due to stochastic drift or biotic interactions between microbes. Moreover, the presence of litter can 374 introduce microbes from other plant parts into the soil (Whitaker et al., 2017, Fanin et al., 2021) and 375 the decomposition of litter can change the soil abiotic environment, thereby altering the soil micro-376 bial community (Veen et al., 2021, Minás et al., 2021). Thus, the microbial community encountered 377 by a responding plant after the time delay may no longer resemble that of the original conditioning 378 (Fig. 2). By quantifying stabilization and fitness differences, we show that microbially mediated 379 plant coexistence outcomes change with the presence of a temporal delay and litter decomposition 380 (grey and brown points in Fig. 5). Our study is a reminder that natural history, particularly in 381 the form of temporal lags between consecutive generations, should be considered when designing 382

³⁸³ and inferring long-term coexistence from plant–soil feedback experiments.

Our sequencing results showed an increase in fungal abundance but a decrease in bacterial 384 abundance between the initial conditioning phase and delayed phases (Fig.2). For both bacterial 385 and fungal communities, the field reference soil contained the highest abundance of microbes. This 386 is in line with the expectation that the field soil acts as a microbial species pool, which is thinned out 387 to a smaller species-specific subset after plant conditioning. However, the high bacterial abundance 388 in A. wrangelianus soil suggests that plants can also enrich specific microbes (Fig.2A). We speculate 389 that the initial low abundance of fungi could be attributed to their slower growth rates, while the 390 enrichment of saprotrophic Ascomycota fungi in the litter addition treatments results from their 391 greater ability to utilize plant litter (Challacombe et al., 2019). On the other hand, the decrease in 392 bacterial abundance after the six-month delay could be attributed to drought-induced physiological 393 stress, the lack of dormancy ability, and intensified resource competition with fungi (Shade et al., 394 2012, Schimel, 2018, Lennon et al., 2021). When comparing the microbial community at the end of 395 the response phases, we found that the total fungal and bacterial community diverges depending 396 on response treatments (Fig. 3). This suggests that the temporal delay of previously conditioned 397 soil will have a long-lasting impact on the microbial species pool and their corresponding impact 398 during the next generation. 390

Our results are the first to demonstrate the influence of a temporal delay by leveraging recent 400 advancements in plant–soil feedback theory (Kandlikar et al., 2019). Moreover, our results highlight 401 nuances when adopting modern coexistence theory to interpret empirical results. First, we show 402 that quantifying invasion growth rates, in addition to the more commonly used stabilization and 403 fitness difference metrics, yields a more nuanced understanding of the mechanisms through which 404 microbial effects on plant coexistence arise. One example is the competition between F. microstachys 405 and *P. erecta*, where the shift in competitive outcomes in the delayed treatments is caused by a loss 406 of stabilization and an increase in fitness difference favoring F. microstachys (Fig. 5A). Examining 407 the invasion growth rates suggests that the shift was primarily due to a decrease in the invasion 408 growth rate of *P. erecta* (Fig. 5D), signifying a change in the microbial effects imposed by the soils of 400 *F. microstachys.* Further investigation of plant biomass responses suggests that the observed shifts 410 in coexistence outcomes arise because conditioned soils of F. microstachys substantially decrease 411

conspecific growth (relative to unconditioned reference) in the immediate treatment, but this 412 negative impact of conditioned soils is minimal after the time lag (Fig. 4B). Second, instead of 413 representing the uncertainty of stabilization and fitness difference as orthogonal error bars, we 414 visualized the distribution of random samples (i.e., using the same approach as in Yan et al., 2022 415 but without the summary pie chart). Our results show that visualizing the spread of predicted 416 outcomes reveals informative patterns: for example, in the case of A. wrangelianus and P. erecta, 417 the diagonal distribution in the delayed-with-litter treatment arises due to the larger variation in 418 the impact imposed by *P. erecta* soils (Ke & Wan, 2020). We echo recent calls for more careful 419 considerations when calculating and visually presenting the uncertainty of predicted competitive 420 outcomes (Terry & Armitage, 2023). 421

We incorporated litter dynamics in our experiment by preserving dead plant individuals 422 in the pot during the temporal gap between the conditioning and response phases. This design 423 corresponds well with our annual plant system, where litter input comes from natural plant senes-424 cence after the growing season. In other systems, different modes of plant death can generate 425 litter dynamics that shape soil microbial communities differently. For instance, wind disturbances 426 that uproot entire plants generate a substantial pulsed input of litter, potentially benefiting sapro-427 trophic microbes but adversely affecting obligated root-associated microbes (Cowden & Peterson, 428 2013, Nagendra & Peterson, 2016). Conversely, herbivores and anthropogenic activities (e.g., clear-429 cutting) primarily remove aboveground parts while leaving belowground components intact in 430 the soil. Such dynamics result in less aboveground litter, yet the remaining root tissues may con-431 tinue to support arbuscular mycorrhizal fungi (Pepe et al., 2018). Recent studies have highlighted 432 that aboveground and belowground litter differentially impact plant-soil microbe interactions. 433 For example, Aldorfová et al. (2022) showed that root litter negatively affected plant performance 434 due to enhanced pathogen transmission while shoot litter modified soil nutrient levels without 435 significantly affecting plant growth. This complicated interplay between different belowground 436 processes underscores the importance of including litter dynamics in microbially mediated plant-437 soil feedback studies (Ke et al., 2015, Veen et al., 2019). 438

We speculate that the impact of a temporal delay extends beyond that shown in our experi ment. First, while we measured plant biomass performance following common plant-soil feedback

studies, the temporal delay may modify microbial effects on other plant demographic rates. For 441 example, seed survival and germination of Californian annuals take place in dry soil after the 442 Mediterranean summer. Following recent calls for studying microbial impact beyond biomass, 443 we encourage future research to study the impact of microbial persistence on the early-stage seed-444 to-seedling transition (Dudenhöffer et al., 2018, Miller et al., 2019, Krishnadas & Stump, 2021). 445 Furthermore, how persistent conditioned microbial effects are after plant death is an important 446 question for many systems, not limited to systems with strong seasonality and non-overlapping 447 generations. For instance, in systems with sparse vegetation cover (e.g., foredunes), a conditioned 448 patch may be left vacant for extended periods due to dispersal limitation. In more complex systems 449 with vertical structures (e.g., forests), one may argue that soil microbes mostly impact seedling 450 survival on conditioned soil beneath the canopy of a living adult. However, the persistence of 451 microbial effects following adult death and canopy opening can influence the performance of 452 seedlings, thereby determining which species can successfully reach the canopy. Therefore, in-453 vestigating the persistence of conditioned microbial effects is likely more critical to community 454 dynamics than previously recognized (Ke & Levine, 2021). 455

456 Conclusion

In our effort to bridge the gap between greenhouse experiments and natural ecosystems, we 457 demonstrate the feasibility and importance of adjusting experimental schedules to provide a more 458 realistic representation of natural systems. In light of our findings, we propose that in annual 459 plant systems with non-overlapping generations, the intricate interplay of natural seasonality and 460 litter dynamics prevent the direct extrapolation of plant-soil microbe interactions from one grow-461 ing season to the next. Our results reveal that the modification of plant-soil feedback following 462 plant death is complex and varies between species pairs, thereby hindering generalizations based 463 on studies that did not consider these factors. With the ongoing shifts in plant phenology and 464 seasonal patterns due to climate change (Rudgers et al., 2020), predicting plant community dynam-465 ics requires the explicit incorporation of the temporal aspects and natural history elements into 466 plant-soil feedback research (Ke et al., 2021). 467

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478 Authorship Contribution

SXO, GSK, and PJK conceptualized and designed the study. SXO performed the experiment and
sample processing. All authors contributed to data analyses and writing.

481 Data Availability Statement

All primary data and computer scripts will be deposited on Github and Dryad upon submission.
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Figure legends

Figure 1. Schematic diagram of the two-phase plant-soil feedback experiment with three fully factorial response treatments: the immediate transplant treatment (light green pots), the delayed without litter treatment (grey pots), and the delayed with litter treatment (brown pots). The two rounds of response phases are six months apart, each including a treatment with field unconditioned soil as references (blue pots). Note that the sterilized soil treatments and batch control pots are not included in this illustration (see Methods).

Figure 2. Absolute abundance of (A) bacterial and (B) fungal taxa (aggregated at the Class level) 638 within the soil inocula used for different response treatments. Each stacked bar represents the ab-630 solute abundance of microbial taxa (x-axis and colors) within a specific soil inoculum (y-axis). The 640 inocula are differentiated by their conditioning host species (i.e., Acmispon wrangelianus (ACWR); 641 Festuca microstachys (FEMI); Plantago erecta (PLER)) and response treatments (i.e., immediate re-642 sponse, delayed without litter, and delayed with litter). The top row, labeled as field reference, 643 depicts soil samples collected from Sedgwick Reserve at the experiment's outset (i.e., prior to the 644 growth of any conditioning individual). 645

Figure 3. Principal coordinates analysis (PCoA) for the combined soil microbial community 646 composition (i.e., bacterial 16S and fungal ITS) sequenced at the end of the response phase. Each 647 panel represents a different inoculum source (conditioning host plant). From left to right: Acmispon 648 wrangelianus (ACWR); Festuca microstachys (FEMI); Plantago erecta (PLER); unconditioned Sedgwick 640 Reserve field soil as reference soil (REF). Each point represents the microbial community sampled 650 from a seedling at the end of the response phase and the shape represents its species identity. Colors 651 represent the three response treatments: immediate (light green), delayed without litter (grey), 652 and delayed with litter (grown). As the two delayed treatments shared the same reference soil 653 controls, we obmitted one of the delayed treatment in the rightmost panel. Purple circles (labeled 654 as SW) represent soils collected from Sedgwick Reserve at the beginning of the experiment (i.e., 655 without the growth of any conditioning or responding individual) and were added for visualization 656 purposes. 657

Figure 4. Effects of soil microbial inocula on plant biomass for (A) Acmispon wrangelianus (ACWR), 658 (B) Festuca microstachys (FEMI), and (C) Plantago erecta (PLER). Each panel shows the aboveground 659 biomass (log-scale x-axis) of the focal plant, grown with a soil microbial community that had been 660 conditioned by conspecifics (blue) or heterospecifics (grey circles); unconditioned communities 661 from field soil (green); or sterilized potting mix (brown). Note that the two delayed treatments 662 shared the same field reference and sterilized potting mix controls. The three plant-conditioned 663 soil inocula are ordered (from bottom to top) as follows: ACWR, FEMI, and PLER. Larger symbols 664 indicate the mean biomass, error bars show $2 \times SEM$, and small points show each individual 665 biomass. 666

Figure 5. Predicted competitive outcomes between pairs of plants: (A & D) Festuca microstachys 667 (FEMI) and Plantago erecta (PLER); (B & E) Acmispon wrangelianus (ACWR) and F. microstachys; (C & 668 F) A. wrangelianus and P. erecta. For each panel, the first and second species listed on the facet label 669 correspond to species 1 and 2 in eqns 1–4, respectively. (A–C) The parameter space of stabilization 670 (x-axis) and fitness difference (y-axis) for the three species pairs. Each region represents different 671 predicted competitive outcomes: the right and left grey triangular regions represent coexistence 672 and priority effect, respectively. The upper and lower white triangular regions represent the 673 dominance of species 1 and 2, respectively. For each species pair, the three response treatments 674 are plotted on the same panel and are indicated by different colors: immediate (light green), 675 delayed without litter (grey), and delayed with litter (brown). Each translucent point represents 676 a random draw (see Methods) and the open black circle represents the mean stabilization and 677 fitness difference of 1000 random draws. (D-E) Invasion growth rates (IGR, y-axis) for the three 678 species pair under different response treatments (x-axis). Different colors represent different plant 679 species: ACWR (green), FEML (orange), and PLER (purple). 680

681 Supporting Information

- ⁶⁸² The following Supporting Information is available for this article:
- 683 Methods S1. Supporting methods for soil microbial community characterization
- ⁶⁸⁴ Fig. S1. Principal coordinates analysis for the bacterial community composition
- ⁶⁸⁵ Fig. S2. Principal coordinates analysis for the fungal community composition
- ⁶⁸⁶ Fig. S3. Effects of soil microbial inocula on plant biomass in all response and control treatments

687 Figures



Figure 1 Schematic diagram of the two-phase plant–soil feedback experiment with three fully factorial response treatments: the immediate transplant treatment (light green pots), the delayed without litter treatment (grey pots), and the delayed with litter treatment (brown pots). The two rounds of response phases are six months apart, each including a treatment with field unconditioned soil as references (blue pots). Note that the sterilized soil treatments and batch control pots are not included in this illustration (see Methods).



Figure 2 Absolute abundance of (A) bacterial and (B) fungal taxa (aggregated at the Class level) within the soil inocula used for different response treatments. Each stacked bar represents the absolute abundance of microbial taxa (x-axis and colors) within a specific soil inoculum (y-axis). The inocula are differentiated by their conditioning host species (i.e., *Acmispon wrangelianus* (ACWR); *Festuca microstachys* (FEMI); *Plantago erecta* (PLER)) and response treatments (i.e., immediate response, delayed without litter, and delayed with litter). The top row, labeled as field reference, depicts soil samples collected from Sedgwick Reserve at the experiment's outset (i.e., prior to the growth of any conditioning individual).



Figure 3 Principal coordinates analysis (PCoA) for the combined soil microbial community composition (i.e., bacterial 16S and fungal ITS) sequenced at the end of the response phase. Each panel represents a different inoculum source (conditioning host plant). From left to right: *Acmispon wrangelianus* (ACWR); *Festuca microstachys* (FEMI); *Plantago erecta* (PLER); unconditioned Sedgwick Reserve field soil as reference soil (REF). Each point represents the microbial community sampled from a seedling at the end of the response phase and the shape represents its species identity. Colors represent the three response treatments: immediate (light green), delayed without litter (grey), and delayed with litter (grown). As the two delayed treatments shared the same reference soil controls, we obmitted one of the delayed treatment in the rightmost panel. Purple circles (labeled as SW) represent soils collected from Sedgwick Reserve at the beginning of the experiment (i.e., without the growth of any conditioning or responding individual) and were added for visualization purposes.



Figure 4 Effects of soil microbial inocula on plant biomass for (A) *Acmispon wrangelianus* (ACWR), (B) *Festuca microstachys* (FEMI), and (C) *Plantago erecta* (PLER). Each panel shows the aboveground biomass (log-scale x-axis) of the focal plant, grown with a soil microbial community that had been conditioned by conspecifics (blue) or heterospecifics (grey circles); unconditioned communities from field soil (green); or sterilized potting mix (brown). Note that the two delayed treatments shared the same field reference and sterilized potting mix controls. The three plant-conditioned soil inocula are ordered (from bottom to top) as follows: ACWR, FEMI, and PLER. Larger symbols indicate the mean biomass, error bars show $2 \times SEM$, and small points show each individual biomass.



Figure 5 Predicted competitive outcomes between pairs of plants: (A & D) Festuca microstachys (FEMI) and Plantago erecta (PLER); (B & E) Acmispon wrangelianus (ACWR) and F. microstachys; (C & F) A. wrangelianus and P. erecta. For each panel, the first and second species listed on the facet label correspond to species 1 and 2 in eqns 1–4, respectively. (A-C) The parameter space of stabilization (x-axis) and fitness difference (y-axis) for the three species pairs. Each region represents different predicted competitive outcomes: the right and left grey triangular regions represent coexistence and priority effect, respectively. The upper and lower white triangular regions represent the dominance of species 1 and 2, respectively. For each species pair, the three response treatments are plotted on the same panel and are indicated by different colors: immediate (light green), delayed without litter (grey), and delayed with litter (brown). Each translucent point represents a random draw (see Methods) and the open black circle represents the mean stabilization and fitness difference of 1000 random draws. (D-E) Invasion growth rates (IGR, y-axis) for the three species pair under different response treatments (x-axis). Different colors represent different plant species: ACWR (green), FEML (orange), and PLER (purple).

Supporting Information for

Realistic time-lags and litter dynamics alter predictions of plant–soil feedback across generations

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² Supporting Methods S1

3 Synthetic spike-in

Using the synthetic spike-in method from Tkacz et al. (2018), we bought plasmids with p-Spike P 4 for prokaryotic 16S (https://www.addgene.org/101172/) and p-Spike F for fungal communities 5 (https://www.addgene.org/101174/). We plated the plasmids on Luria Broth (LB) media with 6 carbenicillin for the ampicillin selection marker and incubated overnight in 25°C. We then picked a 7 colony and inoculated 10 mL of LB broth containing carbenicillin and incubated at 30° C / 120rpm 8 measuring CD on a nanodrop machine at 2 hour intervals until CD concentration = 1. Using 9 zymoPURE Plasmid miniprep kit, we eluted plasmid DNA and measured DNA concentration 10 using High Sensitivity dsDNA Qubit Assay (Thermofisher, Waltham, MA). We loaded a subsample 11 of the eluted plasmid DNA for gel electrophoresis to check for the correct plasmid size. 12

13 Amplicon sequencing

For bacterial metabarcoding, we amplified the highly variable (V4) region of the 16s rRNA gene
 using primers 515F (5'- TCG TCG GCA GCG TCA GAT GTG TAT AAG AGA CAG GTG YCA GCM

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GCC GCG GTAA -3') and 806R (5'- GTC TCG TGG GCT CGG AGA TGT GTA TAA GAG ACA 16 GGG ACT ACN VGG GTW TCT AAT -3'). For fungal metabarcoding, we amplified the fungal 17 ITS1 region using primers based on the ITS1F (5'- AAT GAT ACG GCG ACC ACC GAG ATC TAC 18 ACG GCT TGG TCA TTT AGA GGA AGT AA -3') and ITS2 (5'- CAA GCA GAA GAC GGC ATA 19 CGA GAT - [INDEX] - CGG CTG CGT TCT TCA TCG ATGC -3'), where [INDEX] is a sample-20 specific 12-nt error-correcting Golay barcode. Illumina adapters on each 5' end of the primers were 21 used to attach unique Nextera XT indexes for sample identification. First step PCR consisted of 22 3.2μ L of PCR-grade water, 5μ L of Meridian Bioscience MyTaq HS Red Mix (Bioline, Tunton, MA), 23 0.4μ L each of forward and reverse primers, and 1μ L of extracted DNA. PCR cycles were: 95°C 24 for 2 min, 35 cycles of 95°C for 20 sec, 50°C for 20 sec, 72°C for 50 sec, and a final extension at 25 72° C for 10 min with storage at 4°C. We confirmed amplification by gel electrophoresis. Second 26 step PCR consisted of 3.2μ L of PCR-grade water, 5μ L of Meridian Bioscience MyTaq HS Red Mix 27 (Bioline, Tunton, MA), 0.4 μ L each of Nextera XT index primers 1 and 2, and 1 μ L of first step PCR 28 product. We confirmed amplification by gel electrophoresis and purified amplicons using Sera-29 Mag Speedbeads (Sigma-Aldrich, St. Louis, MO). We quantified DNA concentration using High 30 Sensitivity dsDNA Qubit Assay (Thermofisher, Waltham, MA) and pooled evenly across samples 31 to a concentration of 4nM. The final DNA concentration was quantified using BioAnalyzer and 32 sequenced on an Illumina MiSeq sequencer (2 X 300 cycle sequencing kit, Illumina, San Diego, CA) 33 with a 15% PhiX spike-in at the Stanford Genomic Sequencing Service Center. 34

35 Metabarcoding analysis

Reads were demultiplexed and assigned to samples using Illumina bcl2fastq conversion software. 36 We processed ITS1 and 16S samples separately. We trimmed raw amplicon sequences using 37 Cutadapt (Martin, 2011). We used the DADA2 pipeline (Callahan et al., 2016a) to merge paired-end 38 sequences, quality filter, remove chimeric reads, and cluster sequences into amplicon sequence 39 variants (ASVs). We used the SILVA database (Quast et al., 2012) for 16S taxonomic assignment and 40 the UNITE database (Nilsson et al., 2019) for ITS taxonomic assignment. We removed any ASV that 41 was present in ≤ 5 samples or whose relative abundance was < 0.01 across all samples. We also 42 removed samples with extremely small or large read counts (i.e., more or less than 5x the average 43

⁴⁴ number of reads across all samples). We rarefied samples to 5000 sequencing reads.

45 Supporting Figures



Figure S1 Principal coordinates analysis (PCoA) for the bacterial community composition sequenced at the end of the response phase. Each panel represents a different inoculum source (conditioning host plant). From left to right: *Acmispon wrangelianus* (ACWR); *Festuca microstachys* (FEMI); *Plantago erecta* (PLER); unconditioned Sedgwick Reserve field soil as reference soil (REF). Each point represents the microbial community sampled from a seedling at the end of the response phase and the shape represents its species identity. Colors represent the three response treatments: immediate (light green), delayed without litter (grey), and delayed with litter (grown). As the two delayed treatments shared the same reference soil controls, we obmitted one of the delayed treatment in the rightmost panel. Purple circles (labeled as SW) represent soils collected from Sedgwick Reserve at the beginning of the experiment (i.e., without the growth of any conditioning or responding individual) and were added for visualization purposes.



Figure S2 Principal coordinates analysis (PCoA) for the fungal community composition sequenced at the end of the response phase. Each panel represents a different inoculum source (conditioning host plant). From left to right: *Acmispon wrangelianus* (ACWR); *Festuca microstachys* (FEMI); *Plantago erecta* (PLER); unconditioned Sedgwick Reserve field soil as reference soil (REF). Each point represents the microbial community sampled from a seedling at the end of the response phase and the shape represents its species identity. Colors represent the three response treatments: immediate (light green), delayed without litter (grey), and delayed with litter (grown). As the two delayed treatments shared the same reference soil controls, we obmitted one of the delayed treatment in the rightmost panel. Purple circles (labeled as SW) represent soils collected from Sedgwick Reserve at the beginning of the experiment (i.e., without the growth of any conditioning or responding individual) and were added for visualization purposes.



Figure S3 Effects of soil microbial inocula on plant biomass in all response and control treatments for (A) *Acmispon wrangelianus* (ACWR), (B) *Festuca microstachys* (FEMI), and (C) *Plantago erecta* (PLER). Capital "S" indicates sterilized soils and "L" indicates live unsterilized soils. Colors represent different soil inocula: sterilized potting mix (brown), unconditioned field soil (green), soil conditioned by conspecifics (blue), soil conditioned by conspecifics but sterilized (light blue), soils conditioned by heterospecifics (dark grey), and soils conditioned by heterospecifics but sterilized potting mix controls. The three plant-conditioned soil inocula are ordered (from bottom to top) as follows: ACWR, FEMI, and PLER. Larger symbols indicate the mean biomass, error bars show $2 \times SEM$, and small points show each individual biomass.