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Medicago sativa has Reduced Biomass and Nodulation When Grown with Soil Microbiomes Conditioned to High Phosphorus Inputs

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ABSTRACT

Agricultural over-fertilization may adversely impact plant–microbial interactions affecting crop yield. It is unclear if soil microbiomes respond quickly to changes in fertilizer inputs once conditioned to specific nutrient regimes. We conducted a growth chamber study assessing the compositional and functional resilience of root-associated microbiomes of Medicago sativa to nutrient regime changes, and consequences for plant growth. Plants were grown with a common starting soil microbiome under four nutrient treatments: control (no fertilizer), organic phosphorus (compost added), low inorganic P (low triple superphosphate, TSP) and high inorganic P (high TSP). After several conditioning generations, in which microbiomes from rhizospheres of high biomass plants were transferred forward, microbiome composition was distinct across the four treatments. The resulting microbiomes were then transplanted into each of the nutrient treatments, leading generally to functional changes in hydrolytic enzyme activity and taxonomic convergence with other microbiomes transplanted into the same nutrient regime. However, high inorganic P-conditioned microbiomes were resistant to compositional change. Correspondingly, M. sativa grown with high inorganic P-conditioned microbiomes had lower biomass, fewer nodules, and lower %N than plants grown under the same nutrient regime with other microbiomes. These findings suggest that excessive inorganic P fertilization may change microbiomes such that they negatively affect plant growth.

Additional keywords: extracellular enzyme activity, nutrient amendments.

The availability of soil nutrients for biological uptake is strongly influenced by the activity of microorganisms, which are key drivers of soil biogeochemical cycles. Soil microorganisms are responsible for the transformation of nitrogen (N) to and from plant-available forms (Ghaly and Ramakrishnan 2015). Some bacteria (e.g., diazotrophs) can fix atmospheric nitrogen into soil, or within root nodules created by certain plant species (Bonilla and Bolanos 2009). Soil bacteria and fungi are critical for the phosphorus (P) cycle, as some solubilize fixed inorganic P and mineralize fixed organic P into the soil solution as plant-accessible phosphate ions (Condron et al. 2005; Pierzynski and McDowell 2005; Nannipieri et al. 2011; Sharma et al. 2013). Certain taxa, known as P-solubilizers, are especially active in this regard, releasing extracellular phosphatases or organic acids that free phosphate ions from complex organic molecules or from adsorption to soil minerals (Nannipieri et al. 2011; Sharma et al. 2013). Additionally, some fungi (e.g., arbuscular mycorrhizal fungi [AMF] and ectomycorrhizal fungi) can help plants scavenge available nutrients from the soil by forming root-connected hyphal networks that extend far beyond plant root hairs into the soil matrix (Jakobsen et al. 2005; Jansa et al. 2011).

Diazotrophs, P-solubilizers, and mycorrhizal fungi have long been studied for their individual impacts on plant growth. In contrast, the wide array of taxa within a soil microbiome may contribute additively to overall nutrient cycling, in ways that are less easily understood and that may vary through time. For example, the soil microbial pool can effectively compete with plants for limiting nutrients, leading to nutrient immobilization within microbial cells. (Jakobsen et al. 2005; Sharma et al. 2013). Upon cell turnover, these nutrients again become bioavailable, which can benefit plant growth. Thus, microbiome dynamics can...
substantially influence plant nutrient acquisition, and subsequently plant growth and health.

In agricultural systems, however, the impact of microbial nutrient cycling on plants can be disrupted by repeated soil nutrient amendments. These inputs, added to enhance crop yield and productivity, can chronically impact soil abiotic properties. For instance, soil P often accumulates in agricultural fields following repeated applications of inorganic P fertilizers, with 70 to 95% of applied P remaining adsorbed to soil particles following plant uptake (Hedley and McLauglin 2005). Additionally, organic amendments (e.g., manures or composts), when applied to provide adequate N nutrition, often increase soil P concentrations far beyond what is needed for plant growth (Nelson and Janke 2007). These fertilizer-induced changes in soil nutrients can impact the composition and function of soil microorganisms. For instance, high levels of soil inorganic P may suppress AMF root colonization and phosphatase activity, while organic P may have the opposite effect (Jakobsen et al. 2005; Jansa et al. 2011; Nannipieri et al. 2011; Zhang et al. 2015). Additionally, the abundance of P-solubilizing microbes was shown to decline under high P availability (Zheng et al. 2017). More broadly, studies from various plant–microbe systems show that microbiome composition and function can change substantially following nutrient additions, although the nature and extent of this change depends on the amendment (Eo and Park 2016; Hartmann et al. 2015; Leff et al. 2015; Li et al. 2017; Pan et al. 2014; Wang et al. 2017).

Different microbiomes can then have measurably different effects on plant traits (e.g., biomass, flowering time, drought tolerance, and disease resistance). These microbiome effects are separate from the direct impacts of abiotic soil properties on plant traits, as determined by the transfer of microbiomes to sterile soils (Lau and Lennon 2012; Mendes et al. 2011; Panke-Buisse et al. 2015; Swenson et al. 2000). In general, such experiments show that if different microbiomes in a comparable environment have measurably different functions, it can be inferred that microbial effects—and not just environmental effects—drive differences in ecosystem function (Allison and Martiny 2008). Thus, if two plants of the same genotype perform differently when grown in comparable soils with different microbiomes, much of that difference should be attributable to microbiome effects.

In this study, we determined whether nutrient-mediated changes to soil microbiome composition and activity were retained under alternate nutrient regimes, and whether this impacted the early growth of alfalfa (Medicago sativa). We conditioned a common starting microbiome to growing with M. sativa in one of four soil nutrient input treatments, transferring forward microbiomes from the highest biomass plants for each treatment into twice-autoclaved recipient soils containing the same nutrient treatment, for four generations of conditioning. Following conditioning, microbiomes were again transplanted into twice-autoclaved soils, but this time into each of the four nutrient treatments, yielding sixteen nutrient treatment and microbiome type crosses. We aimed to determine (i) the taxonomic legacy of agricultural nutrient inputs on the soil microbiome when transferred to an alternate nutrient regime, and (ii) the functional legacy of agricultural nutrient inputs on the soil microbiome, particularly with respect to plant performance and soil extracellular enzyme activity (EEA).

MATERIALS AND METHODS

Experimental design. We aimed to isolate the impacts of microorganisms from the impacts of soil nutrients on plant growth and soil function. We focused primarily on modifying soil P, and chose M. sativa (alfalfa), an agriculturally important crop that forms an association with N-fixing rhizobia in root nodules, to reduce the confounding effects of soil N limitation. We conditioned a common starting microbiome growing with alfalfa to different nutrient input regimes, by selecting microbiomes from pots with the highest aboveground plant biomass and collecting root-adherent soil. These microbiomes were then inoculated into twice autoclaved soil and distributed to the next set of pots, and this was repeated for four 3-week generations of plantings. After the fourth generation, the resulting microbiomes were transplanted into each of the four nutrient regimes, yielding 16 nutrient × microbiome crosses. Figure 1 outlines this design.

Nutrient treatments and growth conditions. Plants were grown in a diverse soil mixture derived from multiple farms and modified to have a more optimal soil texture. The mixture consisted of an 8:4:2:2:1 ratio of field soil (Williamson sandy loam, a mesic Typic Fragiaudept, obtained from the Caldwell Research Farm, Cornell University, Ithaca, NY, 42.45°N 76.46°W), sand, peat moss (Lambert, Rivière-Ouelle, QC, Canada), perlite, and fresh soil from the Helfer Research Farm (Niagara silt loam, a mesic Aeric Endoaqualf, Cornell University, 42.45°N 76.44°W), used as an additional microbial source. To this base soil mix, one of four nutrient treatments was applied: (i) control (Ct), with no nutrient amendments; (ii) organic P (OP), with a compost amendment; (iii) low inorganic P (LP), with a moderate input of triple superphosphate (TSP); and (iv) high inorganic P (HP), with a high input of TSP. Mixture ratios for each treatment are shown in Table 1.

These treatments were designed to differ most in soil P content while having the same basic physical structure. Samples of each treatment, postmixing, were sent to the Cornell Nutrient Analysis Laboratory (Ithaca, NY) to determine whether other soil parameters varied between treatments (Table 1). Organic matter was determined by loss on ignition, while pH was measured with a pH meter in 2:1 water/soil suspensions. Nitrate and ammonium soil concentrations were measured with a potassium chloride extraction. Soil P and K were determined from a modified Morgan’s solution extraction. Soil P levels were lowest in the control treatment and highest in the high inorganic P treatment (Table 1). Soil pH was not adjusted and therefore varied between treatments; pH was highest in
Eighty 6-inch pots (20 replicates per treatment) were filled with 1,400 ml of one of the four nutrient-amended soils. All pots contained a common soil microbiome, consisting of the communities from the two field soils in the base soil mix plus 2 ml of compost slurry (50:1 water/compost), using the compost integrated into the OP treatment as an additional microbial source for the Ct, LP, and HP treatments. M. sativa seeds (Nature’s Seed, Lehi, UT) from a common seed pool were surface-sterilized with a 10% bleach solution for 10 min, rinsed five times with distilled water, added to the pots, and incorporated lightly below the soil surface. Each pot was overseeded and subsequently thinned to 20 plants per pot 4 days after germination, with ongoing removal of any late germinating plants. Plants were grown in the Percival-Cornell University Weill Hall Growth Chamber Facility (Ithaca, NY). Conditions were controlled to a 16 h/8 h day and night cycle, with temperatures of 30°C/20°C, 30%/70% relative humidity, and a light level of 330 µE. Each pot received 275 ml of autoclaved water every day for 4 days following seeding, and every other day throughout the rest of each growth period.

Plant harvest and soil sampling. Three weeks following germination, all aboveground plant material was harvested and oven-dried at 75°C for 48 h, and then weighed. The corresponding soil media were stored in bags in a cold room at 4°C during this postharvest period to preserve the microbial communities for potential inoculation into the next generation (details provided in “Generation transfer”).

Meanwhile, five random pots from each treatment were selected for analyses, to represent the range of microbiome composition and activity in each treatment. Soil was shaken from the root systems of each pot and homogenized individually. A small amount of homogenized media from each sample was collected and frozen. Soil samples from conditioning generations one and four were extracted and sequenced to characterize community changes (details provided in “DNA extraction, PCR amplification, and high-throughput sequencing”), and soil samples from conditioning generation four were additionally assayed for extracellular enzyme activity to characterize community functional differences (details provided in “Soil extracellular enzyme activity”).

### Generation transfer

The four pots with the highest aboveground plant biomass (i.e., the top 20% of pots) were identified for each nutrient treatment. Soil from these pots was retrieved from storage, shaken free from plant roots, pooled, and homogenized. This served as the inoculum for the next conditioning generation of the same treatment (Fig. 1).

Dissolved TSP granules and twice-autoclaved compost were mixed into the twice-autoclaved base soil mix in the same ratios as before for each nutrient treatment (Table 1). The prepared soil inoculum was then mixed in at a 5% inoculum/recipient sterile soil ratio.

This method of microbiome transfer better represents the microbiome structure of the source in the next generation than other commonly used transfer methods, such as soil washes (Howard et al. 2017). Plant growth, harvest, and soil sampling were carried out as described above for each generation, for a total of four generations.

### Microbiome transplant

Following the fourth generation, microbiomes conditioned to a specific nutrient regime were transplanted into each of the four nutrient treatments, generating one home-home cross (microbiome introduced to nutrient regime it was adapted to) and three home-away crosses (microbiome introduced to alternate nutrient regimes) for each microbiome type (Fig. 1). Microbial inoculants were prepared for each treatment from the four highest aboveground plant biomass pots as before, but were each divided into four units. Twice-autoclaved soil mixtures were also prepared as before and divided into four units. For each type of inoculant, one unit was mixed with Ct mixture, another with OP mixture, another with LP mixture, and the last with HP mixture, in the same inoculum to soil ratio as before. That is, each treatment’s inoculant was mixed into each treatment’s autoclaved mixture, for a total of 16 microbiome-nutrient treatment crosses (Fig. 1). Each cross consisted of five replicate pots, for a total of 80 pots as before. All other aspects of growth and harvest were the same. Soil samples from all 80 pots of the microbiome transplant were extracted and sequenced (details provided in “DNA extraction, PCR amplification, and high-throughput sequencing”), and assayed for extracellular enzyme activity (details provided in “Soil extracellular enzyme activity”).

It should be noted that because of the method of inoculation, some soil nutrients were inevitably transferred, as in any direct soil

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**TABLE 1**

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Ct</th>
<th>OP</th>
<th>LP</th>
<th>HP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amount of base growth medium per pot</td>
<td>1,400 ml</td>
<td>1,400 ml</td>
<td>1,400 ml</td>
<td>1,400 ml</td>
</tr>
<tr>
<td>Form of P amendment</td>
<td>None</td>
<td>Compost</td>
<td>Dissolved triple superphosphate (TSP) granules</td>
<td>Dissolved triple superphosphate (TSP) granules</td>
</tr>
<tr>
<td>Amount of amendment added per pot</td>
<td>–</td>
<td>200 ml</td>
<td>2.56 g</td>
<td>12.79 g</td>
</tr>
<tr>
<td>Total amount of soil mix per pot</td>
<td>1,400 ml</td>
<td>1,400 ml</td>
<td>1,400 ml</td>
<td>1,400 ml</td>
</tr>
<tr>
<td>Water added per pot</td>
<td>150 ml</td>
<td>150 ml</td>
<td>150 ml</td>
<td>150 ml</td>
</tr>
<tr>
<td>pH</td>
<td>7.55</td>
<td>7.19</td>
<td>6.59</td>
<td>6.12</td>
</tr>
<tr>
<td>Organic matter content</td>
<td>3.97%</td>
<td>4.33%</td>
<td>3.40%</td>
<td>3.70%</td>
</tr>
<tr>
<td>Soil NO₃ + soil NH₄</td>
<td>38.70 ppm</td>
<td>68.27 ppm</td>
<td>40.00 ppm</td>
<td>43.53 ppm</td>
</tr>
<tr>
<td>Soil K</td>
<td>198.50 ppm</td>
<td>595.92 ppm</td>
<td>189.59 ppm</td>
<td>191.80 ppm</td>
</tr>
</tbody>
</table>

*At the start of generation 1 only, all pots received 2 ml of compost slurry (50:1 water/compost) as an additional microbial source.
Cornell Farm Services, Ithaca, NY.
VPG Inc., Bonham TX, NPK 0-45-0.
transfer experiment. That is, each soil mix in the microbiome transplant contained 95% volume of the recipient nutrient treatment mix plus 5% inoculant from another nutrient treatment mix with a different nutrient profile. Thus, soil parameters for each of the four crosses in one soil treatment would have been marginally different. We did not specifically measure these differences, but estimates of differing soil P, for which this effect would be most pronounced, can be found in Supplementary Table S1.

Analysis of nodulation. Following soil sampling, all roots from the microbiome transplant pots were rinsed clean and photographed for the purpose of quantifying the number of rhizobial nodules present. Each photograph included all roots from each pot. In Adobe Photoshop CC 2015, the images were manually examined for nodules by imposing a 1-inch grid over each photo, counting the number of nodules in each grid cell, and summing all grid cell counts (Supplementary Fig. S1).

Analysis of mineral concentration in plant tissues. Following imaging, roots harvested from the microbiome transplant pots were oven dried at 75°C for 48 h, then weighed. Dried roots and aboveground biomass from the same replicate were combined using a mortar and pestle, and ground to a fine powder with liquid nitrogen. Combination of root and shoot material was necessary to obtain enough biomass for these analyses. Samples were submitted to the Cornell Stable Isotope Laboratory (Ithaca, NY) for C and N analysis via combustion, and to the Pennsylvania State University Agricultural Analytical Services Lab (University Park, PA) for P analysis via acid digestion.

AMF root colonization. To determine if the AMF symbiosis influenced plant success in this experiment, plant roots from the first generation of conditioning were harvested, stained, and examined for mycorrhizal arbuscules. Cleaned root material (1 to 2 g) was cleared in 9 ml of 10% KOH at 121°C for 5 min. The roots were then rinsed three times and incubated with 10 ml of a trypan blue stain (200 ml each of distilled water, lactic acid, and glycerol, plus 0.15 g of trypan blue solid) for 24 h at 25°C. Stained roots were examined under a dissecting microscope under several magnifications, but no arbuscules were detected in any sample, as AMF colonization was not firmly established in the 3-week-old *M. sativa* roots. AMF colonization measurements were therefore not conducted for the remainder of the experiment. MiSeq sequencing of the fungal ITS region (details provided in “DNA extraction, PCR amplification, and high-throughput sequencing”) later revealed that a very low proportion of the recovered fungal community was classified as *Glomeromyces*: the highest relative abundance of *Glomeromyces* in any sample (generation 1, generation 4, and the microbiome transplant generation) was 0.07%, and the majority of samples had no detectable *Glomeromyces*.

Soil extracellular enzyme activity. Potential extracellular enzyme activities of five hydrolytic enzymes were measured using a standard fluorometric assay (Bell et al. 2013; Panke-Buissse et al. 2015). Briefly, soil slurries were prepared with 3 g of fresh soil in 150 ml of sodium acetate buffer (50 mM, pH 5), homogenized with an immersion blender, and added to black 96-well microplates (200 μl of slurry per well). A second assay was conducted with a sodium carbonate buffer at pH 7, as this more closely represented the in-situ pH of the Ctn and OP nutrient treatments. A standard curve was generated for each sample as well as for a buffer blank with known concentrations of 4-methylumbelliflone (MUB). To determine enzyme activity, 50 μl of an enzyme-specific substrate bound to MUB was added to the 200 μl of soil slurry. The substrates were 4-MUB-α-D-glucopyranoside, 4-MUB-β-D-glucopyranoside, 4-MUB-β-D-glucopyranoside, 4-MUB-N-acetyl-β-D-glucosaminide, and 4-MUB-phosphate, targeting α-glucosidase (AG), β-xylanosidase (BX), β-glucosidase (BG), N-acetyl glucosaminidase (NAG), and acid or alkaline phosphatase (AP) enzymes, respectively. Each substrate was also tested for autofluorescence. Following a dark incubation at 25°C for 3 h, plates were measured for fluorescence with a BioTek Synergy HT microplate reader (BioTek Industries, Inc., Winooski, VT), at an excitation wavelength of 365 nm and an emission wavelength of 450 nm.

Enzyme activity was calculated as in (Bell et al. 2013), with some modification. Specifically, sample variability was accounted for not by calculating at each sample’s MUB concentration from that sample’s standard curve, but by correcting the sample fluorescence values for quenching. The average reduction in fluorescence (caused by the interference of soil particles) between points on the sample’s standard curve and points on the blank buffer standard curve was calculated, and fluorescence values corrected accordingly. These corrected fluorescence values were then used to calculate MUB concentration in each well from the blank buffer standard curve. Calculations then proceeded normally. This method of calculation better preserved relative fluorescence patterns in the final calculated enzyme activities.

DNA extraction, PCR amplification, and high-throughput sequencing. DNA was extracted from 0.25 to 0.30 g of soil media using the MoBio PowerSoil DNA Isolation Kit (Mo Bio Laboratories Inc., Carlsbad, CA). Five replicates of the base soil mixture and five replicates of the compost were also extracted.

Initial 16S RNA gene PCR reactions were carried out as described in Howard et al. (2017), mainly following the 16S Metagenomic Sequencing Library Preparation guide (part no. 15044223 rev. B) with some modifications. The universal bacterial primers 341F (5’-CTTCAGGNNGGCCWGCAG-3’) and 805R (5’-GACTACHVGGGTATCTAATCC-3’) were used for 16S RNA gene amplifications (Herlemann et al. 2011), and for fungal amplifications, we used the primers ITS1F (5’-CTTGTCATTATGGAGAAATGA-3’) and 58AR2 (5’-CTCGGTCTCCTCATCGAT-3’) (Gardes and Bruns 1993; Martin and Rygiewicz 2005), and all primers were modified to include overhangs needed for indexing attachment, as described in Bell et al. (2016). For 16S RNA gene amplifications, reactions occurred in 20 μl volumes composed of 8 μl of 5 PRIME HotMasterMix (5 PRIME Inc., Gaithersburg, MD), 9 μl of H2O, 1 μl of each primer at 10 μM, and 1 μl of 1:10 DNA dilution. PCR cycling for 16S rRNA gene amplicons was performed using the following protocol: 94°C for 2 min; 25 cycles of 94°C for 20 s, 55°C for 20 s, and 72°C for 30 s; and 72°C for 5 min for the final elongation. For fungal ITS amplifications, the reaction mix was as described above, but with 1 μl of DMSO and only 8 μl of H2O per reaction. PCR cycling for fungal ITS amplicons was performed using the following protocol: 94°C for 3 min; 35 cycles of 94°C for 20 s, 55°C for 30 s and 72°C for 45 s; with a final elongation at 72°C for 5 min.

MagBio HighPrep PCR beads (MagBio Genomics, Gaithersburg, MD) were used to clean the initial amplicons in clear 96-well plates. In new 96-well plates, unique two-barcode index combinations were attached to the cleaned amplicons by placing in each well 5 μl of sample, 2.5 μl of forward and reverse primers with designated barcodes for overhang attachment, 2.5 μl of water, and 12.5 μl of Q5 High Fidelity 2× Master Mix (New England Biolabs Inc., Ipswich, MA). Index attachment PCR cycling conditions were as follows: 98°C for 1 min; 8 cycles of 98°C for 15 s, 55°C for 30 s, and 72°C for 20 s; and 72°C for 3 min for final elongation. The SequlnalPrep Normalization Kit (Thermo Fisher Scientific, Waltham, MA) was then used to normalize the amount of DNA retained from the barcoded amplicons. From each normalized sample, 5 μl was combined into separate pools for 16S rRNA and ITS genes, concentrated, and run on 1.2% agarose gels. The PureLink Quick Gel Extraction Kit was used to purify the relevant extracted bands, for a
final pool volume of 30 μl. Pools were sequenced at the Cornell Genomics Facility (Ithaca, NY) on the Illumina MiSeq. A 600-cycle MiSeq Reagent Kit v.3 was used for the 16S rRNA gene pool, and a 500-cycle MiSeq Reagent Kit v.2 for the fungal ITS pool.

Sequence processing and analysis. Initial sequence processing followed the Brazilian Microbiome Project Pipeline (Pylro et al. 2014) with some modifications. In Mothur v.1.36.1 (Schloss et al. 2009), paired-end sequences were merged with make.contigs, primers trimmed with trim.seqs (pdiffs=2, maxambig=0), and singleton sequences removed with split.abund. Using USEARCH v.7 (Edgar, 2010), 97% OTUs were clustered and chimeras removed (based on the rdp_gold.fa database). Using the GreenGenes v. 13.8 database for 16S rRNA gene sequences, and the UNITE v.7 database for ITS gene sequences, representative OTUs were classified in Mothur. OTUs suspected not to be of bacterial origin were removed with the command remove.lineage (taxon= Chloroplast-Mitochondria-unknown-Archaea-Eukaryota).

Further analysis of OTU and taxonomic tables was conducted in R v. 3.3.2. For all samples, the number of 16S rRNA gene sequences was rarefied to 1,004 reads, and the number of ITS sequences was rarefied to 1,421 reads, to normalize intersample comparisons for downstream analyses in R. There was a high proportion of fungal OTUs unclassified at the phylum level, but these were kept in the dataset for further analysis.

Principal coordinates analyses (PCoA) of Bray-Curtis dissimilarities between samples were carried out using R package vegan to visualize similarity in community composition. Abundance of taxonomic groups was determined using the phyloseq function in the R phyloseq package. Heatmaps and dendrograms of OTU relative abundance were generated with the function heatmap.2 in R package gplots.

Statistical analyses. The lsmeans R package was used to fit linear models in order to test the effects of nutrient treatment, generation, and the interaction between the two for the conditioning generations, or nutrient treatment, microbiome type, and the interaction between the two for the microbiome transplant. Where needed, data were log-transformed to meet the model assumptions of normality of residuals and homogenous variance. Significance of the models was tested with analysis of variance. Pairwise comparisons were then using Tukey’s honest significant difference test. Pearson correlations and significance of correlation were also tested between data sets.

Sequence availability. All sequences for this study were deposited in the SRA database under the project number SRP133237.

RESULTS

Bacterial composition post-conditioning. By the end of the four generations of conditioning under a specific nutrient treatment (either control [Ct], organic phosphorus [OP], low inorganic phosphorus [LP], or high inorganic phosphorus [HP]), distinct bacterial communities were observed in each of the four treatments (Supplementary Fig. S2). HP-conditioned microbiomes were most dissimilar from all other bacterial communities. In generation 1, all microbiomes were more similar to the compost slurry inoculum than the microbiome of the base soil mix, indicating that the compost inoculum was the dominant microbial source for the experiment. Shannon diversity in all treatments declined by generation 4.

Microbiome composition following microbiome transplant. Upon transplant, all microbiomes shifted away from their post-conditioning community composition and converged to a community profile that resembled others in the same recipient nutrient treatment (Fig. 2A). However, HP-conditioned microbiomes were resistant to change across all recipient nutrient treatments, and remained most dissimilar from other microbiomes (Fig. 2A). These compositional differences were relatively minor at the phylum/class level, but were clearly observable at the OTU level (Fig. 2B, Supplementary Fig. S3).

Overall, bacterial community composition seemed most influenced by whether or not inorganic fertilizer was applied, and to what extent, according to the global 16S rRNA gene heatmap (Fig. 2B). Based on unweighted pair-group method with arithmetic mean (UPGMA) hierarchical clustering of Bray-Curtis distances between samples (Fig. 2B), we observed four discrete clusters: (i) bacteria conditioned to the HP nutrient treatment; (ii) bacteria conditioned to other nutrients, but transplanted into the HP nutrient treatment; (iii) bacteria conditioned to the LP nutrient treatment, plus bacteria conditioned to the OP and Ct nutrient treatments but transplanted into the LP nutrient treatment; (iv) bacteria with no exposure to inorganic P fertilization at any point in the experiment. Similar clustering patterns were observed at lower taxonomic levels. Fungi showed similar but less resolved patterns in community composition in the microbiome transplant (Supplementary Fig. S4). Fungal communities transplanted into the HP nutrient treatment were very distinct, with a decrease in Basidiomycetes and an increase in Saccharomycetes.

Bacterial families known to contain either P-solubilizers or potential alfalfa N-fixing symbionts had varying abundance patterns in the microbiome transplant. Some groups had lower abundance in microbiomes conditioned to or transplanted into the HP nutrient treatment (Pseudomonadaceae, Comamonadaceae) (Fig. 3). Other groups had higher abundance in microbiomes conditioned to or transplanted into the HP nutrient treatment (Xanthomonadaceae, Burkholderiaceae), while some groups seemed fairly ubiquitous across treatments (Rhizobiaceae, Bacillaceae) (Fig. 3).

Extracellular enzyme activity. Microbially-produced extracellular enzymes in soil are important indicators of C, N, and P mineralization from soil organic matter, which allows inorganic N and P to be used by plants. As such, these measurements provide a sense of microbiome function and activity. AG, BX, and BG indicate C mineralization, while NAG indicates N mineralization and AP indicates P mineralization (Saiya-Cork et al. 2002). We hypothesized that enzyme activities would be greater under conditions where that nutrient was comparatively limiting in either the soil or the plant, as stoichiometry across soil-microbe-plant systems is often linked (Bell et al. 2014; Xu et al. 2017). Therefore, we expected enzyme activities to be fairly responsive to the change in nutrient treatment in the microbiome transplant. Extracellular enzyme activities were assayed at pH 5 (the standard assay pH) and at pH 7 because the soil pH of the nutrient treatments ranged from 6.12 in the high phosphorus treatment to 7.55 in the control treatment. We wished to measure activities closer to the in-situ pH for each sample, and hypothesized that activity patterns might be different in the two assays.

We found that all enzyme activities were significantly affected by microbiome origin and nutrient treatment (Fig. 4, Supplementary Fig. S5). That is, most microbiomes of the same origin had significantly different enzyme activities based on which soil they were transplanted into. We observed two general patterns in enzyme activity: (i) activity was highest for microbiomes transplanted into the HP nutrient treatment (Fig. 4A to B); and (ii) activity was highest for microbiomes transplanted into the OP nutrient treatment (Fig. 4C to D). Enzymes displaying pattern 1 included AP, NAG, and BG assayed at pH 5 (Fig. 4A to B). Enzymes displaying pattern 2 included AP, AG, and BX assayed at pH 7, and AG assayed at pH 5 (Fig. 4C to D). Enzymes not falling clearly into either pattern (i.e., displaying activity maxima in either HP or OP nutrient treatments) included BG and NAG assayed at pH 7, and BX assayed at pH 5.
Fig. 2. Soil bacterial community composition in the microbiome transplant. A, Principal coordinates analyses (PCoA) ordination of Bray-Curtis dissimilarities of relative operational taxonomic unit (OTU) abundances generated from 16S rRNA gene amplicons. Point colors reflect the microbiome origin, while point shapes reflect contemporary nutrient treatments. Boxes indicate the points that represent home-home crosses for a given nutrient treatment (e.g., control [Ct]-conditioned microbiomes in Ct treatment soil, organic P [OP]-conditioned microbiomes in OP-amended soil, etc.), to illustrate whether a transplanted microbiome is more similar to those from its native nutrient treatment or to those in the novel nutrient treatment. Shaded ellipses delineate groupings observed based on unweighted pair-group method with arithmetic mean (UPGMA) clustering, and correspond to clusters in panel B. B, Global heatmap of OTU relative abundances for 16S rRNA gene amplicons with UPGMA clustering of samples. UPGMA clustering is based on analyses of all OTUs in the dataset, but only bacterial OTUs with at least 2% relative abundance in at least one sample are shown in the heatmap. Colored bars to the right of the sample dendrogram reflect the microbiome origin and contemporary nutrient treatment of samples. Clusters of samples with similar communities are indicated with colored shading imposed over the sample dendrogram.
Activity patterns of some enzymes were quite responsive to the assay pH. AP activities switched from pattern 1 at pH 5 to pattern 2 at pH 7 (Fig. 4A and C), as did BG to some extent. Other enzymes, including NAG (Fig. 4B), AG (Fig. 4D), and BX were more consistent in both assays.

**M. sativa nodulation, growth, and tissue nutrients.** In the four generations of conditioning, there was variability in *M. sativa* aboveground biomass from generation to generation, but generally biomass was highest for plants grown in the OP nutrient treatment, followed by plants grown in the LP nutrient treatment, the Ct nutrient treatment, and the HP nutrient treatment, in that order (Supplementary Fig. S6). Plants grown in the HP nutrient treatment appeared shorter, had smaller leaves, and were yellowing by the time of harvest (Supplementary Fig. S7).

In the microbiome transplant, *M. sativa* biomass was significantly affected by nutrient treatment and microbiome origin (Fig. 5A to D). Plants grown in the same nutrient treatment with microbiomes conditioned to either Ct, OP, or LP amendments generally showed only minor or insignificant differences in biomass (Fig. 5A to D). Additionally, all plants grew equally well in the OP nutrient treatment regardless of microbial origin (Fig. 5B). However, in the other three nutrient treatments, plants grown with HP-conditioned microbiomes had significantly lower biomass than plants grown with one of the other microbiomes within the same nutrient treatment (Fig. 5A, C, and D).

*M. sativa* nodulation was also significantly affected by nutrient treatment and microbiome origin (Fig. 5E to H). In the OP, LP, and HP nutrient treatments, plants grown with OP- or LP-conditioned microbiomes had significantly lower nodule counts (Fig. 5F to H). In the Ct, OP, and LP nutrient treatments, plants grown with LP-conditioned microbiomes had significantly higher nodule counts (Fig. 5E to G). Nodule count was lower for all plants in the HP nutrient treatment, even when plants were grown with microbiomes with which they nodulated well in other nutrient treatments (Fig. 5H), suggesting some abiotic limitation on nodulation in the HP nutrient treatment. Nodule count was significantly, positively correlated with aboveground biomass from generation to generation, but generally biomass was signifi-
cantly lower than plants grown with other microbiomes in the same nutrient treatment. All plants grown in the HP nutrient treatment had lower tissue % N. Nodule counts and plant tissue % N were significantly and positively correlated (P < 0.05). Plant tissue % P was significantly and positively correlated with estimated soil P content (P < 0.05).

**DISCUSSION**

In this study, an initial soil microbiome (derived from agricultural soil and compost) was conditioned to grow with *M. sativa* under four nutrient treatments: no fertilizer (control), organic compost, low inorganic P, and high inorganic P. The four resulting microbiomes were then introduced to each nutrient regime, with *M. sativa* grown under each of the 16 conditions. We aimed to determine the taxonomic and functional resilience of the soil microbiome after different nutrient regimes, and impacts on subsequent plant growth and nodulation.

We note that while microbiomes were conditioned for four generations, they were assessed for legacy effects after only one 3-week generation. Thus, observed microbiome responses to new conditions were quite rapid, while observed legacy effects may have broken down over time.

**Taxonomic legacy of nutrient conditioning.** Soil microbiomes were strongly influenced by conditioning to nutrient treatments. It should be noted that the treatments were not controlled for soil pH and thus varied simultaneously in pH and soil P; increasing soil P was negatively correlated with soil pH (Table 1). Therefore, this study does not determine which of these properties is the driving factor behind the observed differences. However, fertilizer applications to agricultural soils will affect both pH and soil nutrients at once, and these results are still of practical interest.

Nutrient amendments have been shown to alter bacterial communities at the phylum level (Leff et al. 2015), but we observed clearer differences at lower taxonomic levels. For example, *Gammaproteobacteria* abundance did not vary substantially between treatments, despite large shifts within this class at the OTU level. We also observed large between-treatment shifts at the family level. We also observed large between-treatment shifts at the family level.
level for bacteria associated with P-solubilization or nodulation (Fig. 2). Although we cannot definitively assign function to these groups, we hypothesize that these changes also reflect differences in nutrient cycling between treatments.

Following the microbiome transplant, historical legacy—defined as the persistence of biotic conditions created by prior environments (Hawkes and Keitt 2015)—was strongest for HP-conditioned microbiomes, in which taxonomy remained distinct, irrespective of soil nutrient conditions (Fig. 2). The HP nutrient treatment substantially altered the abiotic environment, particularly with respect to soil pH and P (Table 1), potentially inhibiting microorganisms that thrive in the other treatments, while promoting those that tolerate high phosphorus loading and a more acidic environment. In contrast, the other treatments may have favored common taxa, while still maintaining a resilient microbial pool that can quickly respond to environmental shifts. We note that, like many microbiome transfer studies, soil compounds were transferred along with microorganisms, leading to slight differences in estimated soil P, and potentially impacting taxonomy to some extent.

We also note that while microbiomes were conditioned for four generations, they were assessed for legacy effects for only three weeks post-transplant. Thus, the time scale of microbiome responses to new conditions was quite short, while observed legacy effects may or may not have persisted in a longer experiment. This would depend on whether nutrient conditioning led to the exclusion of certain microorganisms, or whether it merely shifted the composition of dominant microbes, while allowing other microbes to survive at lower densities.

**Functional legacy of nutrient conditioning.** We also assessed the functional legacy of nutrient conditioning on soil microbiomes in two ways: (i) plant performance (growth, nodulation, and nutrition); and (ii) EEA.

M. sativa growth negatively affected by HP-conditioned microbiomes. Considered alone, soil abiotic conditions strongly influenced plant growth and nutrition. The Ct nutrient treatment was likely deficient in soil P for alfalfa growth; alfalfa biomass plateaus around 20 ppm soil P (Pang et al. 2010) and is considered P-deficient if plant tissue P is less than 0.25% (Lissbrant et al. 2009). Ct treatment soil contained 12.56 ppm P (Table 1), and plants grown in the Ct-Ct cross had 0.17% P (Table 2). OP and LP nutrient amendments therefore raised soil P and plant % P above deficiency thresholds (Tables 1 and 2), explaining increased biomass in these

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**Fig. 4.** Extracellular enzyme activities of generation 4 and microbiome transplant samples. Only a subset of enzymes assayed are displayed here, to demonstrate general activity patterns. Bars represent mean activity ± standard error. Results are grouped by microbiome origin to reflect microbiome responses to novel nutrient conditions. The home-home cross of the microbiome treatment (e.g., control [Ct]-conditioned microbiomes in Ct treatment soil, organic P [OP]-conditioned microbiomes in OP-amended soil, etc.) always follows the generation 4 bar, then the home-away crosses. Analysis of variance results for each enzyme are displayed at the top of the panel. Letters indicate significant groupings from Tukey’s post hoc honest significant difference test. Letters compare results within the same microbiome type, and are therefore not shared through a panel but are specific to the group of four bars they appear with. A, Acid or alkaline phosphatase (AP) activity assayed at pH 5. B, N-acetyl glucosaminidase (NAG) activity assayed at pH 5. C, AP activity assayed at pH 7. D, α-Glucosidase (AG) activity assayed at pH 7.
treatments. However, plant growth was reduced in the HP nutrient treatment. While some plants would experience P toxicity at the soil P levels of the HP nutrient treatment (Pang et al. 2010), the more likely explanation here for biomass reduction is adverse soil pH. The HP nutrient treatment, at a pH of 6.12, is close to the threshold pH at which rhizobia nodulation and N-fixation are impaired for alfalfa (Cheng et al. 2002; Correa et al. 2001; Rice et al. 1977), and plants grown in these soils had significantly reduced nodulation and %N. This was true regardless of microbiome type. Plants grown with the same microbiome had more nodules in the Ct, OP, and LP nutrient treatment than in the HP nutrient treatment. This strongly suggests an abiotic limitation on nodulation success.

Ultimately, however, we were interested in the microbiome effect on plant growth, which would indicate a functional legacy of nutrient conditioning. In the case of Ct, OP, and LP-conditioned microbiomes, community structure rapidly converged when these

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**Medicago sativa** growth in the Microbiome Transplant

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Aboveground Biomass (g)</th>
<th>Root Nodule Count</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ct Nutrient Treatment</strong></td>
<td><img src="image1" alt="" /></td>
<td><img src="image2" alt="" /></td>
</tr>
<tr>
<td><strong>OP Nutrient Treatment</strong></td>
<td><img src="image3" alt="" /></td>
<td><img src="image4" alt="" /></td>
</tr>
<tr>
<td><strong>LP Nutrient Treatment</strong></td>
<td><img src="image5" alt="" /></td>
<td><img src="image6" alt="" /></td>
</tr>
<tr>
<td><strong>HP Nutrient Treatment</strong></td>
<td><img src="image7" alt="" /></td>
<td><img src="image8" alt="" /></td>
</tr>
</tbody>
</table>

*Fig. 5. Alfalfa growth and nodulation in the microbiome transplant. A to D, Aboveground biomass, displayed by nutrient treatment. E to H, Root nodule raw count, displayed by nutrient treatment. Counts were of nodules on all roots present in a single pot. Bars are colored by microbiome origin and represent mean biomass or nodule count ± standard error (n = 5) for that nutrient treatment-microbiome origin cross. Home-home crosses (e.g., control [Ct]-conditioned microbiomes in Ct treatment soil, organic P [OP]-conditioned microbiomes in OP-amended soil, etc.) are bolded. Analysis of variance results for biomass and nodule count are displayed on the y-axes. Letters above bars indicate significant differences in biomass or nodulation by Tukey’s honest significant difference test.*
Soil extracellular enzyme activities showed adaptation to soil nutrient conditions. Though we found that HP-conditioned microbiomes had a functional legacy with respect to their impact on plant growth, we did not observe a functional legacy for any of our microbiomes related to extracellular enzyme activity. In contrast, our results indicate rapid microbial environmental filtering and/or adaptation in response to nutrient management and/or plant feedback.

As a technical note, our results demonstrate the importance of assay conditions relative to in situ sample conditions, as has previously been discussed (German et al. 2011). We assayed EEA in two pH buffers to reflect pH differences in our nutrient treatments. EEA for several hydrolytic enzymes varied by an order of magnitude between pH 5 and pH 7 for some samples, with large changes in relative activity between treatments for certain enzymes (Fig. 4A versus 4C). This reinforces the idea that enzyme activity optima can vary in a soil-specific manner (Turner 2010), which should be addressed by assay conditions.

We observed two general EEA patterns. For AP, NAG, and BG assayed at pH 5, microbiomes transplanted into the HP nutrient treatment had the highest enzyme activity (Fig. 4A and B). For AP, this pattern likely reflects soil pH. The substrate used for the phosphatase assay can be cleaved by both acid and alkaline phosphatase, which have pH optima around 4 to 6 and 8 to 10, respectively (Nannipieri et al. 2011; Turner 2010). Thus, we suspect our pH 5 assay mostly measured acid phosphatase activity, while our pH 7 assay suboptimally measured alkaline phosphatase. High AP activity measured at pH 5 in the HP nutrient treatment, regardless of microbiome origin, therefore indicates a rapid functional shift; environmental filtering and/or microbial adaptation selected

| TABLE 2 | Summary of plant tissue nutrient content in the microbiome transplant**w |
|---|---|---|---|---|---|
| Nutrient treatment | Microbiome origin | %C* | %N* | %P* | C:N | C:N:P |
| Ct | Ct | 42.82 ± 0.55 a | 2.91 ± 0.04 bc | 0.17 ± 0.004 d | 15:1 | 253:17:1 |
| Ct | OP | 42.64 ± 0.41 a | 3.12 ± 0.07 ab | 0.20 ± 0.005 c | 14:1 | 213:16:1 |
| Ct | LP | 43.89 ± 0.21 a | 3.44 ± 0.07 a | 0.29 ± 0.009 b | 13:1 | 150:12:1 |
| Ct | HP | 40.63 ± 0.97 b | 2.57 ± 0.10 c | 0.95 ± 0.113 a | 16:1 | 43:3:1 |
| OP | Ct | 43.57 ± 0.18 a | 3.34 ± 0.11 a | 0.47 ± 0.022 bc | 13:1 | 92:7:1 |
| OP | OP | 44.09 ± 0.26 a | 3.02 ± 0.13 ab | 0.44 ± 0.025 c | 15:1 | 101:7:1 |
| OP | LP | 43.82 ± 0.14 a | 3.31 ± 0.04 a | 0.52 ± 0.014 b | 13:1 | 85:6:1 |
| OP | HP | 43.15 ± 0.13 a | 2.65 ± 0.08 b | 0.63 ± 0.022 a | 16:1 | 69:4:1 |
| LP | Ct | 43.73 ± 0.25 a | 3.56 ± 0.14 ab | 0.83 ± 0.014 b | 12:1 | 53:4:1 |
| LP | OP | 43.72 ± 0.40 a | 3.32 ± 0.20 b | 0.74 ± 0.022 b | 13:1 | 59:4:1 |
| LP | LP | 44.00 ± 0.39 a | 3.83 ± 0.14 a | 0.80 ± 0.014 a | 11:1 | 55:5:1 |
| LP | HP | 39.53 ± 0.69 b | 1.61 ± 0.06 c | 1.13 ± 0.030 a | 25:1 | 35:1:1 |
| HP | Ct | 40.34 ± 0.29 b | 1.79 ± 0.05 a | 1.60 ± 0.045 a | 22:1 | 25:1:1 |
| HP | OP | 40.89 ± 0.96 ab | 1.91 ± 0.12 a | 1.68 ± 0.109 a | 21:1 | 24:1:1 |
| HP | LP | 42.39 ± 0.32 a | 1.89 ± 0.06 a | 1.72 ± 0.052 a | 22:1 | 25:1:1 |
| HP | HP | 40.84 ± 0.27 ab | 1.78 ± 0.06 a | 1.78 ± 0.040 a | 23:1 | 23:1:1 |

**w Nutrient concentration (% dry weight) in alfalfa plants (pooled shoots and roots) in the microbiome transplant experiment. %C, %N, and %P are displayed as mean ± SE (N = 5) for that nutrient treatment-microbiome origin cross. Crosses are grouped by nutrient treatment, and bolded crosses indicate the home-home cross for a given nutrient treatment (e.g., control [Ct]-conditioned microbiomes in Ct treatment soil, organic P [OP]-conditioned microbiomes in OP-amended soil, etc.). Letters denote significant differences from Tukey’s honest significant difference test. Letters compare results within a single nutrient treatment and are therefore not shared across a column, but are specific to the group of four crosses they appear beside. C:N and C:N:P were calculated from mean %C, %N, and %P for the cross and rounded to the nearest integers.

x %C: P microbiome × treatment = 0.00016.
y %N: P microbiome × treatment < 2.2e-16.
z %P: P microbiome × treatment < 2.2e-16.
within three weeks for a microbiome that produces phosphatases optimally suited for the in situ soil pH, rather than the pH of historic nutrient treatments. Mirroring this was AP activity measured at pH 7, as activity was highest in the OP nutrient treatment where phosphatases active around pH 7 would be most useful (Fig. 4C).

For NAG assayed at pH 5, high activity in the HP nutrient treatment may reflect adaptation of a different sort. Generally, legumes can exude soluble N compounds from their roots (Fustec et al. 2011), a readily available source of N for soil microorganisms. However, plants grown in the HP nutrient treatment had fewer N-fixing nodules and were N-starved (Fig. 5F, Table 2), and would not likely have deposited as much N to the rhizosphere as plants grown in other nutrient treatments, making N more limiting. In such situations, microorganisms may preferentially alter extracellular enzyme release to acquire relatively scarce nutrients (Bell et al. 2011). The increase in NAG activity may thus have been a response to soil N-limitation caused by fewer plant-derived soil N inputs. A similar explanation may apply to BG assayed at pH 5, as plants in the HP nutrient treatment showed lower biomass, which may have had decreased C inputs to the rhizosphere (Ikeda et al. 2014).

The second general pattern: for AP, AG, and BX assayed at pH 7, and AG assayed at pH 5, microorganisms transplanted into the OP nutrient treatment displayed the highest enzyme activity. AG and BX are carbon-mineralizing, and activity differences may reflect the fact that only the OP treatment added C to soil. Microorganisms transplanted to this environment could have increased enzyme production, allowing them to access nutrients that were less available during conditioning. The remaining enzymes do not fit cleanly into either pattern, and enzyme production was likely shaped by multiple factors, or factors not described here.

Conclusions. Soil nutrient amendments quickly and significantly altered the structure and activity of the soil microbiome of alfalfa in this growth chamber experiment. These changes were relatively reversible, especially with regards to extracellular enzyme activity, but a taxonomic legacy effect was observed for microorganisms conditioned to extreme inorganic P fertilization. *M. sativa* plants had lower biomass and fewer nodules in soils treated with high inorganic P, and when grown in the presence of microorganisms selected under high inorganic P under most other nutrient conditions. Although the degree of amendment in the high inorganic P treatment was extreme, this suggests that agricultural amendments resulting in excess soil P could have adverse impacts on the ability of the soil microbiome to support crop plants, even after changes to soil management.

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