



Variation in Root Exudate Composition Influences Soil Microbiome Membership and Function

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ABSTRACT Root exudation is one of the primary processes that mediate interactions between plant roots, microorganisms, and the soil matrix, yet the mechanisms by which exudation alters microbial metabolism in soils have been challenging to unravel. Here, utilizing distinct sorghum genotypes, we characterized the chemical heterogeneity between root exudates and the effects of that variability on soil microbial membership and metabolism. Distinct exudate chemical profiles were quantified and used to formulate synthetic root exudate treatments: a high-organic-acid treatment (HOT) and a high-sugar treatment (HST). To parse the response of the soil microbiome to different exudate regimens, laboratory soil reactors were amended with these root exudate treatments as well as a nonexudate control. Amplicon sequencing of the 16S rRNA gene illustrated distinct microbial diversity patterns and membership in response to HST, HOT, or control amendments. Exometabolite changes reflected these microbial community changes, and we observed enrichment of organic and amino acids, as well as possible phytohormones in the HST relative to the HOT and control. Linking the metabolic capacity of metagenome-assembled genomes in the HST to the exometabolite patterns, we identified microorganisms that could produce these phytohormones. Our findings emphasize the tractability of high-resolution multiomics tools to investigate soil microbiomes, opening the possibility of manipulating native microbial communities to improve specific soil microbial functions and enhance crop production.

IMPORTANCE Decrypting the chemical interactions between plant roots and the soil microbiome is a gateway for future manipulation and management of the rhizosphere, a soil compartment critical to promoting plant fitness and yields. Our experimental results demonstrate how soil microbial community and genomic diversity is influenced by root exudates of differing chemical compositions and how changes in this microbiome result in altered production of plant-relevant metabolites. Together, these findings demonstrate the tractability of high-resolution multiomics tools to investigate soil microbiomes and provide new information on plant-soil environments useful for the development of efficient and precise microbiota management strategies in agricultural systems.

KEYWORDS exometabolomics, metagenomics, rhizodeposition, root exudates, sorghum

The region where plant roots interface with soil, known as the rhizosphere, is one of the most intricate and diverse microbial ecosystems on earth (1). It is widely recognized that exudation of compounds from plant roots is key to assembling these rhizosphere microbial communities (2–4), which in turn can influence plant health and soil

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biogeochemistry. Plant roots exude up to 40% of their photosynthetically derived carbon, with these compounds ranging from small (e.g., carbohydrates, amino acids, organic acids, and phytohormones) to large (e.g., proteins and mucilage) molecules (5, 6). These exudates provide microbial substrates, as well as signaling molecules, that stimulate microbial activity. For instance, rhizosphere-enriched microorganisms can suppress plant pathogens and provide plant nutrients through nitrogen fixation and metal chelation, among other mechanisms (2, 7–9). Moreover, some members of the rhizosphere can produce plant hormones like indole-3-acetic acid (IAA), gibberellins, cytokinins, and abscisic acid, which modulate plant physiological health (10, 11). The importance of these various contributions highlights the clear need to understand the mechanistic link between plant root exudate chemistry and metabolic changes in the soil microbiome.

Recently there has been an increased focus on plant rhizodeposition and its impact on soil microorganisms. To enhance tractability in a complex system, many of these keystone studies selected model plants (e.g., *Brachypodium distachyon* and *Arabidopsis thaliana*) and evaluated their impact on known plant growth-promoting rhizobacteria (PGPR) (e.g., *Pseudomonas*) (12–15). However, the scaling of these concepts to more complex chemical profiles and soil microbial communities has been constrained by methodological challenges. For example, where exudates were chemically profiled, they were often limited to a small set of known metabolites (16–19), or when microbial communities were analyzed, they often relied on methods that tracked changes in membership (16S rRNA, phospholipid fatty acid profiles), but not metabolic potential over the course of exudation (20, 21). Therefore, there is a current need for studies that employ high-resolution microbial and chemical methods to begin to mechanistically describe the exudate-microbe interactions in agriculturally relevant crop species.

Here, we used a multiomics approach to characterize how root exudation from a model crop plant can drive soil microbial community structure and function. Using three common, but phenotypically diverse, sorghum (*Sorghum bicolor* (L.) Moench) genotypes, we determined the variation in root exudate chemical composition among these plant varieties. To narrow in on the microbial response to these different exudate profiles, we created plant-free soil microcosms and fed these with two sorghum-informed exudate amendments. We tracked these microcosms over 20 days using exometabolomics, 16S rRNA gene profiling, and genome-resolved metagenomics. Integrating these data, we show that different sorghum exudate treatments led to distinct microbial communities and metabolisms, including the production of three phytohormones. Harnessing this knowledge could support the growing need for sustainable agroecosystems by developing holistic agricultural management strategies that optimize the metabolic capabilities of the soil microbiome.

RESULTS AND DISCUSSION

Sorghum genetics influence root exudation patterns. Sorghum is one of the most widely produced agricultural crops in the world, serving as a grain, forage, and cover crop for use as human food, livestock feed, and as a biofuel feedstock (22, 23). Due to its diverse agronomic uses, we leveraged three sorghum genotypes known to have distinct aboveground phenotypes that we hypothesized would contribute to distinct root exudate profiles. We selected (i) the grain sorghum BTx623, (ii) the sweet sorghum type Leoti, and (iii) the bioenergy sorghum PI 505735 (23–25). We grew sorghum genotypes *in vitro* (hydroponically) for 7 days, and soluble exudates were collected in water and analyzed with nontargeted gas chromatography-mass spectrometry (GC-MS).

Metabolites from sorghum seedlings spanned known root exudate metabolite classes ranging from sugars to sugar alcohols, organic acids, and amino acids (Fig. 1A; see Fig. S1 in the supplemental material). Chemical differences were observed between genotypes BTx623 and Leoti, while PI 505735 represented an intermediate root exudate chemical profile (Fig. 1A). Among BTx623 and Leoti, statistical differences in exudate profiles were found (analysis of variance [ANOVA], $P < 0.001$), with BTx623

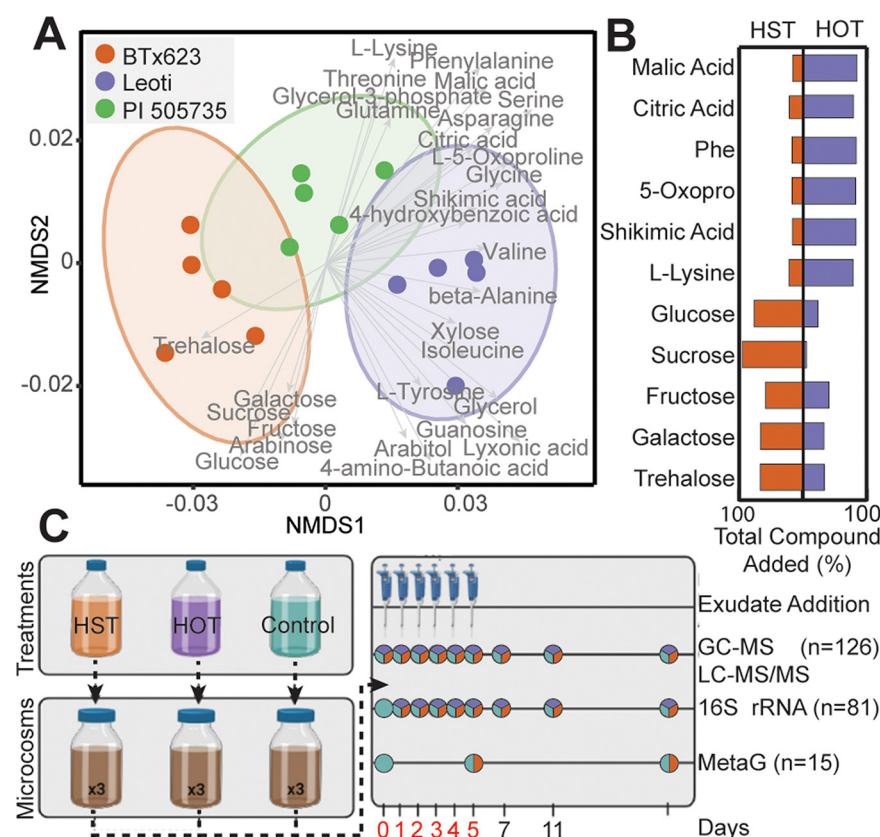


FIG 1 Root exudation metabolite profiles from three sorghum genotypes were used to design relevant amendments for soil microcosms. (A) Nonmetric multidimensional scaling (NMS) of Bray-Curtis distances across the metabolite abundance of Leoti (purple), BTx623 (orange), and PI 505735 (green) sorghum genotypes (stress = 0.06). Ellipses denote the 95% confidence interval for each treatment. Significant metabolite vectors are shown ($P < 0.05$) and labeled in gray. (B) Butterfly plots show the summed total amount of each compound added across the treatments (Phe, phenylalanine; 5-Oxopro, 5-oxoproline); orange represents the proportion added to high-sugar treatment (HST), and purple represents the proportion added to the high-organic-acid treatment (HOT). For example, of the sucrose added, 95% went to the HST and 5% to the HOT. For actual added concentrations, see Table S3. (C) The microcosm schematic depicts the treatment formulation, where HOT represents Leoti, HST represents BTx623, and Control was buffered medium lacking an exudate treatment. Triplicate microcosms with agricultural soil were maintained for a 20-day experiment, with the sampling schematic denoting when samples were obtained. Days with exudate addition are noted with pipettes, and the day number is in red. Time points with circles represent the samples taken each day, with the total number (n) of samples for each analysis listed. Colors within circles represent the type of sample (HST, orange; HOT, purple; control, teal) taken at that time point.

exudates enriched in monosaccharides and disaccharides, while Leoti was significantly enriched in organic and amino acids (Fig. 1A; see File S1 in the supplemental material). Given these sorghum genotypes released root exudates with distinct metabolic profiles, we next pursued how these different exudate regimens would alter microbiome membership and metabolic responses.

Laboratory microcosms reveal that exudate treatment structures agricultural soil microbial communities. We formulated root exudate solutions for laboratory-scale soil reactors (microcosms) using either a high-sugar treatment (HST) representing BTx623 or a high-organic-acid treatment (HOT) representing Leoti. These treatments were run in parallel with exudate-lacking control microcosms. Both HST- and HOT-amended microcosms received the same 11 root exudate compounds (glucose, galactose, fructose, sucrose, trehalose, malic acid, lysine, phenylalanine, 5-oxoproline, citric acid, and shikimic acid) (Table S3), but in various concentrations to model the relative concentrations observed in BTx623 and Leoti exudates (Fig. 1B). The soil microcosms were constructed using soil from semiarid agricultural plots and were amended with exudate treatments

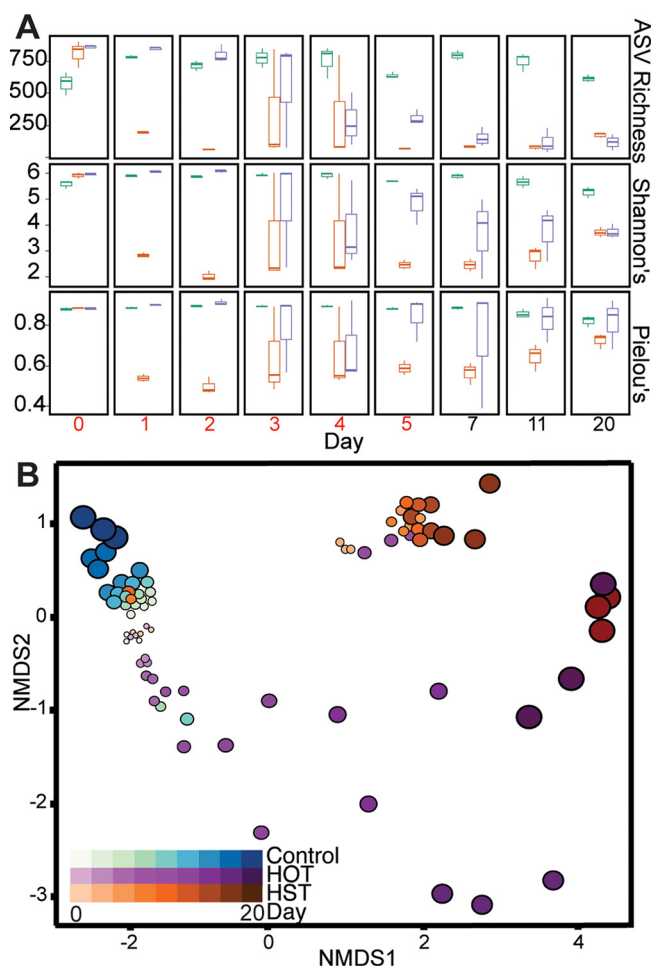


FIG 2 16S rRNA gene diversity metrics and membership changes with exudate amendment. (A) ASV richness (top), Shannon's diversity index (middle), and Pielou's evenness for each day highlighting differences in treatment richness, alpha diversity, and community evenness. Coloring corresponds to treatment: control, teal; HST, orange; and HOT, purple. Days where exudates were added are in red (days 0 to 5). (B) Nonmetric multidimensional scaling (NMDS) of Bray-Curtis distances of 16S rRNA amplicon communities showing changes in microbial community structure and membership over time, with colors representing treatment and the size and darkness of circles representing time (stress = 0.09).

daily for the first 5 days and sampled for chemical and microbiological analyses over 20 days (Fig. 1C).

Gene amplicon sequencing of the 16S rRNA gene was used to temporally profile the microbial diversity and membership across the three treatments (HOT, HST, exudate control) (Fig. 1C). We sequenced 81 samples, generating 2,148,831 reads with an average of more than 30,000 reads per sample (File S2). After denoising, a total of 9,818 amplicon sequencing variants (ASVs) were detected, representing 43 phyla (File S2). We first assessed alpha and beta diversity metrics to understand soil microbial community changes in response to exudates.

Over time, we observed distinct changes in response to amendments with either HST or HOT (Fig. 2A). For instance, after 1 day of exudate amendment, HST-amended microcosms had significant decreases in ASV richness ($P = 0.007$), Shannon's diversity index ($P = 0.019$), and Pielou's evenness ($P = 0.017$). In contrast, we did not detect significant changes in these metrics for HOT or control microcosms in this time period. This suggests HST enriched for select members of the microbial community. In support of this result, we observed an ASV belonging to the genus *Pseudomonas* (File S2) accounted for 59.4% of the HST community at day 1. Following 5 days of exudate

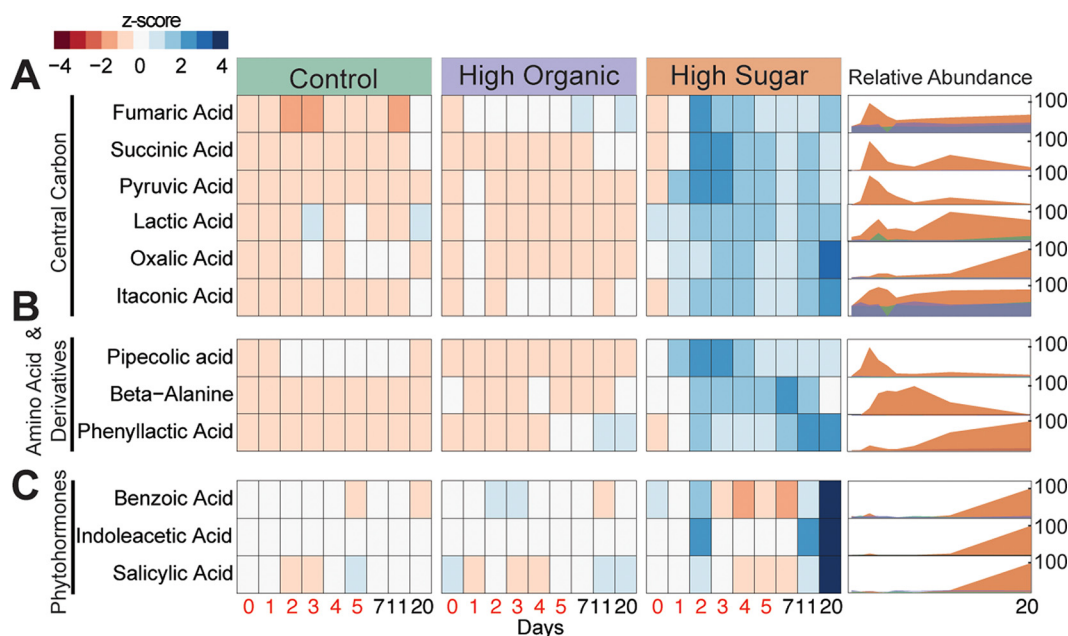


FIG 3 Exometabolite abundances across treatments. We categorized the detected exometabolites into three chemical classes: (A) central carbon metabolism, (B) amino acids and derivatives, and (C) phytohormones. Within each, heat maps (left) show the relative abundance (represented as a Z-score across samples) of each exometabolite detected. Time increases from left (day 0) to right (day 20) within each treatment, with days where exudate treatments were added in red (days 0 to 5). Ridgeline plots (right) show the relative abundance (0 to 100%) of these exometabolites over time. Coloring corresponds to treatment: control, teal; HST, orange; and HOT, purple.

addition, the HST-amended microcosms retained low diversity ($P = 0.038$), richness ($P = 0.042$), and evenness ($P = 0.017$) compared to the HOT and control microcosms (Fig. 2A). By the final day of the experiment, these diversity metrics were no longer significantly different between HST and HOT, highlighting the importance of these amendments in structuring the microbial communities.

Next, changes in the membership and structure of the microbial community across treatments were assessed using the Bray-Curtis dissimilarity matrix and visualized using nonmetric multidimensional scaling (NMDS) (Fig. 2B). We found that HST and HOT amendments significantly shifted microbial community composition and structure during and after periods of synthetic root exudate addition (multiresponse permutation procedure [mrpp] = 0.01, $P < 0.05$). A permutational multivariate analysis of variance (PERMANOVA) revealed that microbial communities were significantly altered across treatments ($P = 0.001$) and time ($P = 0.001$). Building on this, beta dispersion analyses indicated the control exudate communities were the most stable over time, while the HOT had the greatest across-time variation in microbial community membership (Fig. 2B). These microbial community alpha and beta diversity metrics highlight that both treatment and time were factors shaping the soil microbiome in exudate-amended microcosms. Scaling beyond these microcosms, this research highlights the need for understanding the composition and temporal dynamics of natural root exudates.

Exometabolites hint at potential microbial metabolisms stimulated by exudates.

In these soil reactors, we tracked the temporal dynamics of exometabolites (Fig. 1C), which are the extracellular fraction of molecules that are inferred to be produced and/or utilized by soil microorganisms. We classified and assigned the detected exometabolites to three chemical classes: (i) central carbon metabolism, (ii) amino acids and derived compounds, and (iii) phytohormones (Fig. 3; File S1). The exometabolites were coordinated to the microbial communities identified by 16S rRNA gene sequencing, and thus demonstrated differences by treatment and time (Fig. 3; Fig. S2). Relative to

the HOT and controls, the HST was enriched in most of the detected exometabolites, likely reflecting a greater metabolic stimulation from the sugar rich treatment (Fig. 3). This finding is not completely unexpected, as prior studies have noted sugar metabolism is more efficient than amino acid metabolism in soil microorganisms (26).

Metabolites relevant to microbial central carbon metabolism were some of the most enriched exometabolites in HST microcosms (Fig. 3A). We detected six organic acids that changed significantly over time in HST and were significantly enriched in HST compared to the control and HOT. For example, between days 2 and 3, succinic acid and fumaric acid increased 1.5- and 0.9-fold (\log_2), respectively, in the HST-amended microcosms relative to HOT and the control treatments (Fig. 3A). We also detected increases in itaconic acid (0.7-fold) and oxalic acid (1.6-fold) over time, both of which can be derived from tricarboxylic acid (TCA) cycle intermediates *cis*-aconitate and oxaloacetate, respectively (27). Furthermore, consumption of malate and citrate was inferred in HST-amended microcosms at the same time points due to their loss over time (Fig. S3). Perhaps indicating metabolite cross-feeding across the community, we also observed a spike in pyruvate at days 2 to 3 in the HST, followed by consumption concomitant with the production of lactic acid across the experiment (Fig. 3A). Collectively, these exometabolite data pointed to HST differentially altering soil carbon pools via the metabolism of the microbiome.

We also detected chemical evidence for production and consumption of amino acid and amino acid-derived compounds by these soil microbial communities. Like the central carbon metabolites, these organic nitrogen exometabolites were most enriched in HST-amended microcosms relative to the other treatments, a somewhat unexpected response given the HOT amendments were initially dosed with these types of compounds (Fig. 1B). Compared to control and HOT-amended microcosms, we observed the production of two nonproteinogenic amino acids (β -alanine and pipecolic acid [PA]) and one aromatic compound (phenyllactic acid [PLA]) in HST-amended microcosms (Fig. 3B). β -Alanine, which was produced over the first 7 days but removed by day 20, is an important amino acid precursor and also necessary for biosynthesis of coenzyme A, a critical cofactor of microbial metabolism in soils (28).

Next, we identified and tracked the production of pipecolic acid (PA), whose production peaks between days 2 and 4 in HST-amended microcosms. This amino acid is thought to be microbially synthesized from lysine (29), a component added to both treatments (Fig. 1B), that was consumed during this period (Fig. S3). PA synthesis has broader ramifications for the entire microbiome, as it is a required intermediate in secondary metabolite production of antibiotics and anthelmintics in many bacteria (29). PA dynamics, which indicate production and subsequent consumption by day 20, suggest its use as a public good (30). Finally, phenyllactic acid (PLA), a phenylalanine derivative, increased 2.6-fold in the HST microcosms over time, peaking at days 11 and 20 (Fig. 3B). Microbes that release amino acid and amino acid-derived compounds could be competitive root colonizers or act as antagonistic agents against target pathogens (31).

Most notably, we observed a significant increase in the abundance of three phytohormones. Salicylic acid (SA), benzoic acid (BA), and indole-3-acetic acid (IAA) significantly increased ($P < 0.01$) over the experiment in the HST relative to microcosms treated with HOT or the exudate-lacking control (Fig. 3C). In HST microcosms, BA and SA increased 1-fold (\log_2), while IAA increased the most, with a 3-fold (\log_2) increase from day 0 to day 20. In plants, SA, BA, and IAA are vital phytohormones integral to physiological processes like plant defense and development (32). Our detection of these metabolites over time highlights that certain root exudation chemical profiles can stimulate soil microbes to produce phytohormones critical for plant growth and defense.

Curation of a sorghum exudate-responsive microbial genome catalog. In light of the phytohormone production observed in our HST microcosms, we sought to identify microbial genomes capable of producing these compounds. Towards this goal, we

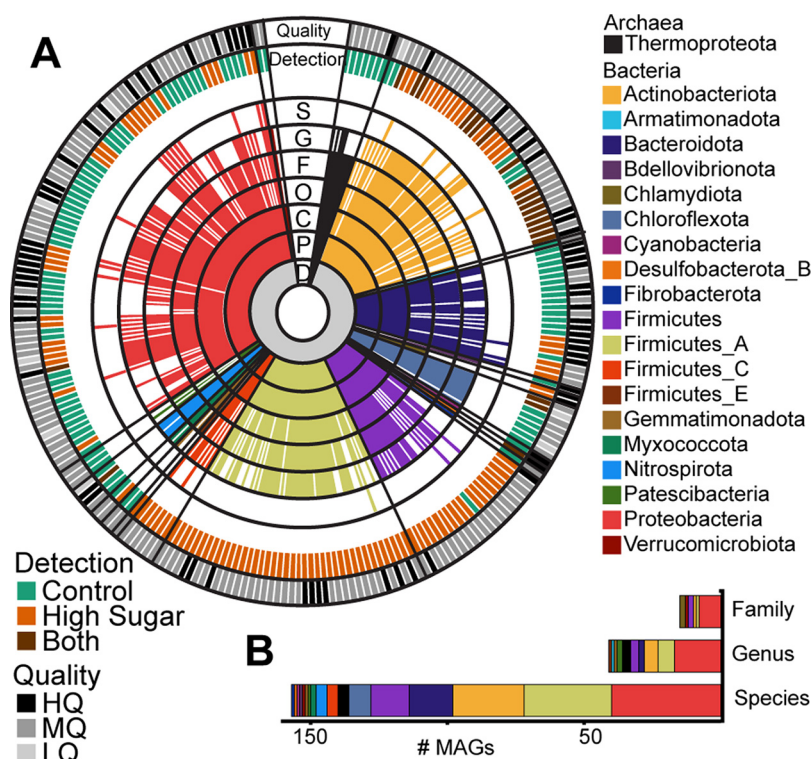


FIG 4 Taxonomy of the 243 dereplicated metagenome-assembled genomes in the genome database. (A) Sequential colored rings indicate the most-resolved taxonomic level that could be assigned by GTDB-tk (72). Taxonomic level is denoted in black with a single-letter abbreviation: D, domain; P, phylum; C, class; O, order; F, family; G, genus; and S, species. Ring color corresponds to phylum assignment, with the color legend at right. The treatment condition corresponding to MAG detection is illustrated in the outer ring labeled “Detection”; MAGs detected only in control metagenomes are indicated by teal, MAGs detected only in HST metagenomes are indicated by orange, and MAGs that were detected in both are indicated in brown (see Materials and Methods for detection thresholds). The MAG quality is shown in the outermost ring following the MIMAG standards (33): high quality (HQ; >90% complete and <5% contamination), medium quality (MQ; >50% complete and <10% contamination), and low quality (LQ; here defined as >48% complete and <10% contamination). (B) Stacked bar graph shows the number of dereplicated MAGs recovered that represent novel families, genera, or species according to taxonomy assignments from GTDB-tk. Coloring corresponds to MAG phylum.

constructed metagenome-assembled genomes (MAGs) representing the suite of soil microbes in the HST and control microcosms. To maximize MAG recovery, we obtained more than 365 Gbp total of sequencing from control and HST microcosms, at three different time points spanning the experiment (control, days 0, 5, and 20, and HST, days 5 and 20, each in triplicate; $n = 15$ metagenomes) (Fig. 1C). With these data, we reconstructed 371 MAGs that were dereplicated at 99% identity into 243 MAGs, of which 28% were high quality (33) (Fig. 4A; File S2). The genome recovery demonstrated here exceeds those from field-based studies, where assembling and binning are hampered by the vast complexity in agricultural soils (34), further validating the relevance of stimulated laboratory microcosms to increase the microbial genome tractability of soil systems (35, 36).

Of these MAGs, 49% ($n = 119$) were detected in HST-amended microcosm samples, while 41% ($n = 100$) were detected in the control microcosm samples (Fig. 4A; File S2). Notably, only 10% ($n = 24$) of these MAGs were shared across the two treatments, further supporting our 16S rRNA and exometabolite data showing HST amendments significantly altered the soil microbial community. Our MAG database contains representative soil microbes across 20 phyla, dominated by members of the *Actinobacteriota* and *Proteobacteria* (Fig. S4), and includes MAGs belonging to 15 previously unidentified families (6%), 41 previously unidentified genera (17%), and 157 previously

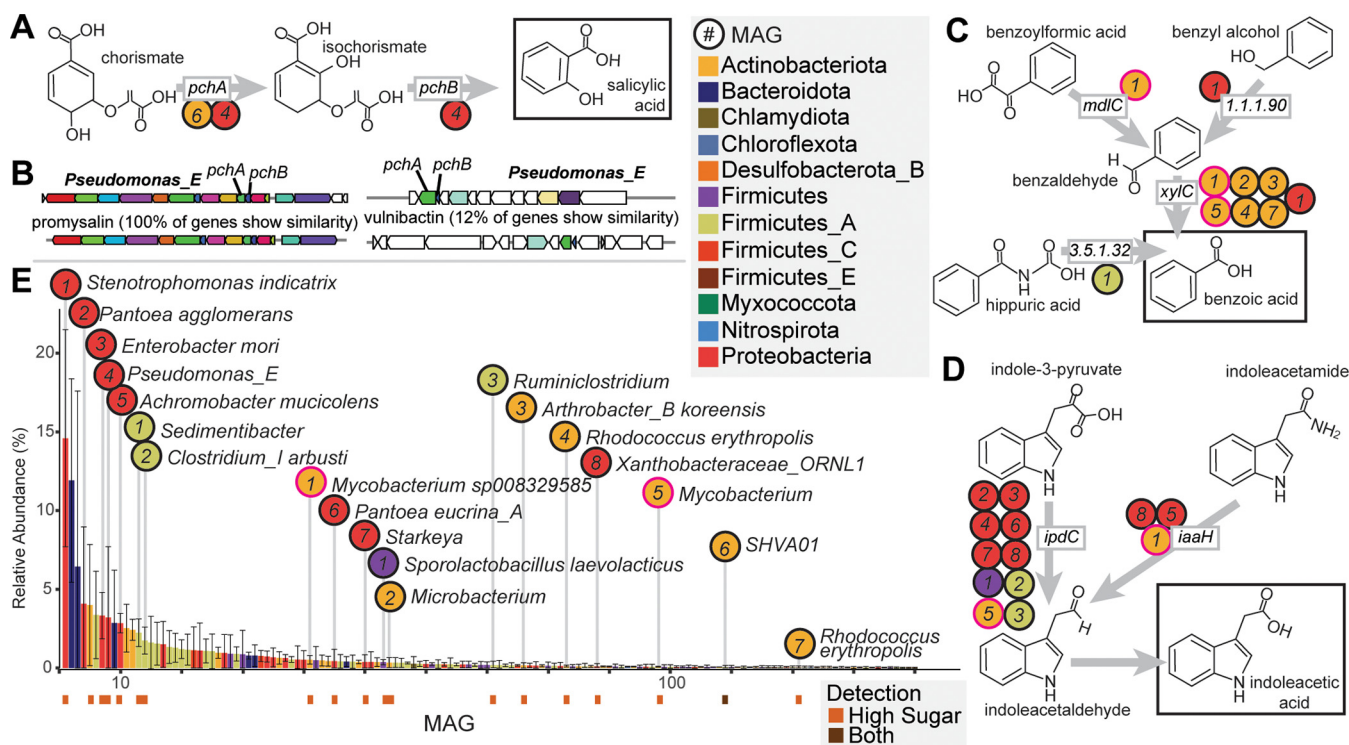


FIG 5 Diverse MAGs encode biosynthetic potential for salicylic acid, benzoic acid, and indoleacetic acid. (A) Salicylic acid production from chorismate by *pchAB*. (B) A *Pseudomonas* MAG carried two biosynthetic gene clusters containing *pchAB*, including one for the antimicrobial promysalin (left) and another for a predicted siderophore (right). The clusters on the bottom are the reference clusters from antiSMASH. (C) Benzoic acid production from multiple pathways. (D) Indoleacetic acid production pathways. (E) Rank abundance curve of MAGs detected in HST metagenomes at day 20. Bars represent the average relative abundance ($n = 3$), and error bars represent 1 standard deviation. Bars are colored by MAG phylum, and key phytohormone-producing MAGs are indicated. MAG detection is denoted below abundance bars for potential phytohormone-producing MAGs, with MAGs detected only in HST in orange ($n = 18$) and MAGs detected in both HST and the control in brown ($n = 1$). In panels A, C, and D, circles correspond to MAGs carrying each gene in panel E. The circles for the two *Mycobacterium* MAGs with BA and IAA potential are outlined in pink.

unidentified species (65%) (Fig. 4B). We recovered 26 MAGs with partial or full 16S rRNA genes and could directly link 19 of these MAGs to 16S rRNA amplicon sequencing-identified ASVs, providing metabolic blueprints for these taxa. These results highlight the advantage of coupling community profiling (16S rRNA gene) with high-resolution metagenomics to capture the functional potential encoded within the soil microbiome.

Bacterial salicylic acid production is assigned to a potentially new species of *Pseudomonas*. We examined which of our MAGs carried the two genes for SA production (*pchA*, encoding isochorismate synthase, and *pchB*, encoding isochorismate pyruvate lyase). The first gene, *pchA*, catalyzes the conversion of chorismate to isochorismate, with the second gene, *pchB*, catalyzing the subsequent conversion to salicylate (Fig. 5A) (37). Two of our HST-detected MAGs carried a *pchA* gene, while one of these MAGs also carried *pchB*. Notably, the MAG that carried both *pchA* and *pchB* likely represents a novel species within the genus *Pseudomonas_E* (L_E1_T20_B_bin.65) and had two copies of this gene set (Fig. 5B). Consistent with our exometabolite data for SA production (Fig. 3C), this *Pseudomonas_E* MAG was abundant in HST metagenomes at day 20 (Fig. 5E).

Previous research has shown that bacterial species such as those of the genera *Pseudomonas*, *Bacillus*, *Mycobacterium*, and *Azospirillum* synthesize this important compound, with pseudomonads, like those enriched in our reactors, exhibiting a propensity for SA production (38). It is recognized that microbially produced SA can act as a plant defense hormone providing pathogen resistance and modulating developmental signaling cascades (39–41). Beyond phytohormone roles, bacterially produced SA can function as a siderophore for metal acquisition from soils (38, 39, 42) or to support antibiotic production (43). Excitingly, further genomic analysis of the *Pseudomonas_E* MAG

revealed one set of *pchAB* genes, which occurred in a predicted siderophore biosynthetic gene cluster (Fig. 5B). Furthermore, the second *pchAB* cluster occurred in a 16-kb gene cluster with 100% gene similarity to the promysalin biosynthetic gene cluster from *Pseudomonas putida* (Fig. 5B). Promysalin is an antibiotic that contains SA and was shown to be selectively antagonistic to other closely related pseudomonads to enable rhizosphere colonization (44). Our data suggest the HST microcosms created conditions that stimulated the *Pseudomonas_E* MAG to produce SA, which could ultimately aid in its competition for limited resources in the complex soil microbiome. Additionally, the accumulation of SA in the enrichment leaves open the possibility that beyond enhanced microbial metabolism, this exometabolite could also be available to act as a plant defense hormone.

Diverse benzoic acid production pathways encoded by multiple MAGs. We next investigated BA production potential in our MAGs. Known BA production pathways derive from degradation of aromatic compounds (Fig. 5C) (45). The terminal step of most of these pathways is the oxidation of benzaldehyde to BA by benzaldehyde dehydrogenase (encoded by *xylC*). In these microcosms, seven MAGs carried *xylC*, including *Stenotrophomonas indicatrix*, *Arthrobacter koreensis*, two MAGs of *Rhodococcus erythropolis*, and two species of *Mycobacterium* (Fig. 5C). These MAGs were only detected in the HST microcosms at day 20 (Fig. 5E), matching when BA was observed in exometabolomic data (Fig. 3C).

The *S. indicatrix* MAG encoded an aryl-alcohol dehydrogenase (EC 1.1.1.90) adjacent to *xylC*, which would enable reduction of benzyl alcohol to benzaldehyde (Fig. 5C). This is also the most abundant MAG in HST metagenomes at day 20 (Fig. 5E), indicating it may be important to BA production in the HST microcosms. Beyond *S. indicatrix*, a MAG representing *Mycobacterium* sp008329585 encoded a benzoylformate decarboxylase (*mdlC*), which decarboxylates benzoylformic acid to benzaldehyde (Fig. 5C). Finally, an abundant MAG representing a novel species of *Sedimentibacter* carried a gene for hippurate hydrolase (Fig. 5E). This enzyme cleaves glycine from hippuric acid, producing BA (EC 3.5.1.32) (Fig. 5C). Consistent with the literature, these findings position BA as a central aromatic metabolite in these microcosms.

While there are limited reports today that microbial BA production can have a direct benefit to plants, it is well known that plant BA and derivatives are precursors to the production of secondary metabolites and hormones (46). BA is also known to improve plant stress tolerance and contribute to growth regulation (46). In support, exogenous application of BA has been associated with improved growth and yield in quinoa and drought-stressed soybean (47, 48). As such, BA synthesis by the soil microbiome and its relationship with plant health are an area of research that warrants more investigation.

Bacterial indoleacetic acid production encoded by diverse MAGs. Tryptophan (Trp) is the primary precursor for IAA biosynthesis in microorganisms (49), with five known biosynthetic pathways characterized in bacteria (50). We surveyed our MAG database for these genes and found 14 HST-detected MAGs capable of IAA production using two different pathways (File S3), consistent with reports that auxin biosynthesis is widely encoded in plants (43, 49). Nine MAGs carried the indole-pyruvate decarboxylase (*ipdC*) gene, the key gene for converting Trp to IAA via indole-3-pyruvate (Fig. 5D). These MAGs spanned four phyla, with six belonging to *Proteobacteria*, covering several of the most abundant MAGs at day 20 in the HST metagenomes (Fig. 5E). Beyond this pathway, three MAGs carried *iaaH* for IAA production via indoleacetamide (Fig. 5D). One of these MAGs, a novel species of *Xanthobacteraceae*, carried both *iaaH* and *ipdC*. This fits with previous observations of single organisms encoding redundant IAA biosynthesis pathways (51). Of note, two mycobacterial MAGs with IAA production potential also had BA production potential. One potentially novel *Mycobacterium* species carried *ipdC* and *xylC*, while *Mycobacterium* sp008329585 carried *iaaH* in addition to *mdlC* and *xylC*, indicating some soil microbes can produce multiple phytohormones.

IAA is a phytohormone in the auxin family that is necessary for proper plant development and can be used to stimulate root biomass growth (51, 52). From a microbial perspective, IAA production offers a competitive colonization strategy over non-IAA-

producing strains (53, 54), and it has been shown that microbially produced IAA in the rhizosphere can elicit positive plant growth promotion and disease suppression effects (18, 53–55). However, these effects are dose dependent, as high concentrations of IAA can also inhibit plant growth (56–58). The reduced complexity of our laboratory microcosm system allowed us to demonstrate that IAA can accumulate in soil microbiomes, and its production is redundantly encoded across 4 different phyla, offering new perspectives on the microbial contributions to soil IAA pools.

It is important to note that the metagenomic results presented here only represent the genomic capabilities of a hypothesized genome; thus, additional experimental evaluation of gene expression would be required to definitively confirm specific bacterial synthesis of SA, BA, and IAA and their roles as potential species of plant growth-promoting rhizobacteria (PGPR). Furthermore, gene expression could inform whether exometabolite production was a direct microbial response to exudates versus an indirect response via the priming effect (59). Yet, we find these MAG results, coupled with exometabolite evidence for these compounds through time, to be an exciting platform for targeted studies aimed at harnessing the power of microbial metabolisms for improving agroecosystems.

Conclusion. Coupling a reduced-complexity, tractable laboratory soil system with high-resolution multiomics approaches afforded new perspectives on the complex interactions between plant root exudation and the soil microbiome. We first showed that root exudate amendments modeled after two different sorghum genotypes drastically impacted overall soil microbial community diversity and membership. Second, shifts in microbial community structure mirrored differences in exometabolites. Finally, we linked interesting plant-relevant metabolites to bacterial genomes, highlighting new roles for specific microorganisms in phytohormone biosynthesis. We have provided a publicly available, genome-resolved microbial database for researchers interested in the sorghum rhizosphere. Our findings suggest root exudate composition influenced microbial production of possible phytohormones that could impact host plant physiology, as well as govern metal acquisition and antibiotic production from soils. Future studies can scale these hypotheses to the field to analyze the impact of plant genotype, root exudates, and associated microbial communities on overall plant performance. In summary, the results of this study represent an important step toward decoding the complex chemical language between plants and their rhizosphere microbial communities, a translation required to optimize these interactions for enhanced agricultural management and production.

MATERIALS AND METHODS

Sorghum root exudate collections. Mature seeds from three sorghum genotypes (Leoti, BTx623, and PI 505735) were first sterilized by placing seeds in sterile 50-mL conical tubes with 45 mL 95% ethanol. Tubes were vortexed, ethanol was removed, and 45 mL of Captan fungicide solution (0.2 g Captan fungicide in 45 mL sterile water) was added to remove any fungal components on the seeds, and the mixture was vortexed for 3 h. After removal of the Captan fungicide, 45 mL of 100% Clorox bleach (8.25% sodium hypochlorite active ingredient) and 3 droplets of dish soap (Ajax Triple Action Orange) were added for the final sterilization step, and seeds were shaken for 20 min. In a sterile tissue culture hood, the bleach was removed using sterile techniques, and the seeds were rinsed five times with sterile water. Seeds were blotted dry and placed on germination paper inside a clean 600-mL beaker with a solution of 1 mM CaCl_2 . After 7 days of growth, seedlings were removed from the paper and pooled into a 250-mL glass bottle filled with 80 mL of ultrapure water. Root exudates were collected for each genotype in separate bottles. The bottles were covered with aluminum foil to protect roots from the light and were placed on a rotary shaker. After 2 h, the roots were removed, blotted dry, and weighed. The root exudate suspensions containing the root exudates were filtered through a 0.2- μm -pore filter membrane to remove root detritus and microbial cells. Samples were frozen at -80°C . We recognize that the use of hydroponic growth chambers does not exactly replicate soil growth conditions of sorghum (60); however, this system provided a highly controllable, tractable, and sterile environment that eliminated confounding microbial or soil influences for more accurate downstream analytical detection. Samples were lyophilized completely before resuspension in 15 mL sterile high-performance liquid chromatography (HPLC)-grade water. Resuspended exudates were vortexed thoroughly to remove all residue from the bottle, transferred to a clean 50-mL Falcon tube, and dried completely under nitrogen gas. One milliliter of liquid chromatography-mass spectrometry (LC-MS)-grade 80% acetonitrile was added to the Falcon tube with dried exudates and vortexed until thoroughly resuspended. A 0.5-mL portion was

transferred to a clean 2-mL glass vial, and 5 μg of glucose- $^{13}\text{C}_6$, D-arabinose- $^{13}\text{C}_5$, sucrose- $^{13}\text{C}_{12}$, galactose- $^{13}\text{C}_6$, and fructose- $^{13}\text{C}_6$ was added to each vial and dried completely under nitrogen prior to derivatization.

Sample derivatization for sorghum root exudates. Dried samples were resuspended in 75 μL of 0.2 M methoxyamine hydrochloride in pyridine (Sigma), incubated at 60°C for 45 min, vortexed, sonicated for 10 min, incubated a second time at 60°C for 45 min, and allowed to cool to ambient temperature. Thirty microliters of each sample was then transferred to a separate 2-mL glass vial for MSTFA (*N*-methyl-*N*-trimethylsilyltrifluoroacetamide) derivatization and the other 30 μL for acetic anhydride (AA) derivatization. Ninety microliters of 100% AA was added to AA vials, and the mixture was vortexed for 30 s, incubated at 60°C for 60 min, and cooled for 10 min at ambient temperature. Thirty microliters of MSTFA plus 1% trimethylchlorosilane (TMCS) (Thermo Scientific) was added to the MSTFA vials, and the mixture was vortexed for 30 s, incubated at 60°C for 35 min, and cooled to ambient temperature before transferring to vial inserts. AA samples were dried completely under nitrogen, resuspended in 40 μL of 100% ethyl acetate, and then loaded into glass inserts for analysis.

Nontargeted and targeted GC-MS analyses for sorghum root exudates. Metabolites were analyzed and detected using a Trace 1310 gas chromatograph (Thermo) coupled to an ISQ mass spectrometer (Thermo). Samples (1 μL) were injected into an injection port at 285°C and a 1:10 split ratio. Separation was accomplished with a 30-m TG-5MS column (Thermo Scientific [0.25-mm inside diameter i.d., 0.25- μm film thickness]) and a helium gas at a flow rate of 1.2 mL/min. The oven temperature program started at 80°C for 30 s, was ramped to 330°C at 15°C/min, and then was held at the final temperature for 8 min. The transfer line and ion source were maintained at 300°C and 260°C, respectively. Masses between 50 and 650 m/z were scanned at 5 scans/s after electron impact ionization. A pooled quality control (QC) sample (made by combining an equal volume of all samples) was injected after every 6 samples to monitor instrument stability throughout the analysis.

Synthetic exudate preparation. Exudate treatments were formulated based on the results of the exudate profiles for the two most divergent sorghum genotypes (Leoti and BTx623). The high-sugar treatment (HST) was formulated to mimic BTx623, with higher sugar and lower organic acid composition, and the high-organic-acid treatment (HOT) was formulated to mimic the Leoti exudate profiles, with higher organic acid and lower sugar composition (see Table S3 and Files S5 and S6 in the supplemental material). The 11 synthetic root exudate compounds were chosen based on practical considerations and availability of standards. Organic acid root exudate relative quantification values were normalized to fructose peak area and scaled to the absolute quantification of fructose in that sample. Treatment calculations can be found in Files S5 and S6. The exudate-lacking medium control treatment (control) was a 10 mM phosphate buffer, consisting of ammonium chloride, disodium phosphate, and sodium dihydrogen at a pH of 6.5 (Table S1). Exudates found in each genotype were weighed out to their respective masses (Table S3) to supply equal exudates for 6 days of exudate addition (Fig. 1C, days 0 to 5) to the microcosms, homogenized, suspended in 10 mL 10 mM phosphate buffer, vortexed, aliquoted into 6 tubes for each day of exudate addition, and frozen at -80°C until use. Aliquots were removed 30 min before use, incubated at 24°C until thawed, and added to microcosms at the respective time point.

Soil samples. The soil (microbial inoculum) was collected from agricultural fields at the Colorado State University Agricultural Research and Education Center (CSU-ARDEC) near Fort Collins, CO, on 4 October 2019. The climate at the site is semiarid, with a mean annual precipitation of 408 mm and a mean annual temperature of 10.2°C (average from 1981 to 2010 [<https://usclimatedata.com/>]). The soil is classified as an Aridic Haplustalf. Three 2-cm-diameter soil core samples to a depth of approximately 15 cm were collected from each of seven different plots. The soil was stored at -20°C until microcosm construction. Twenty grams of soil from each replicate/plot was pooled and homogenized to create the representative bulk soil repository used in the following microcosms experiment.

Microcosm experimental setup. Microcosms were established and sampled as previously described (35, 60). Briefly, 5 g of homogenized soil and 35 mL of phosphate-buffered (pH 6.5) minimal medium (Table S1) were added to sterile 50-mL conical tubes to construct each microcosm. Microcosms were vortexed and allowed to settle for 5 min. Then, day 0 samples were taken by removing 1 mL of soil slurry for exometabolomics analysis and 1 mL of soil slurry for DNA extraction. After this initial sampling, 2 mL of exudate treatment was added to each microcosm and vortexed, and a second 1-mL aliquot was immediately taken for exometabolomics analysis. At this point, for each microcosm the bottle caps were removed and replaced with a sterile foam stopper for the rest of the experiment to maintain oxic conditions and prevent colonization by contaminating microbes. Microcosms were incubated in an orbital shaker set at 200 rpm at 24°C for 20 days. Each exudate treatment (i.e., HST, HOT, and control) was conducted in triplicate, and treatments were added (2 mL) to microcosms on days 0, 1, 2, 3, 4, and 5. After day 5, no additional exudate treatments were applied, but microcosms were maintained until day 20, which afforded additional samples taken at days 7, 11, and 20 (Fig. 1C). Samples were collected at roughly the same time each day and collected with aseptic techniques to ensure no additional microbial influence was introduced. All collected samples were immediately frozen at -80°C until processing.

Sample preparation and extraction for exometabolomics analysis. Samples were thawed at 4°C overnight and centrifuged for 20 min at $18,000 \times g$, and supernatant was transferred to a preweighed 1.5-dram vial and refrozen at -80°C for subsequent lyophilization. Samples were lyophilized for 24 h until all water was sublimated. Samples were weighed to calculate total exometabolite mass and then were resuspended in 4-mL sterile HPLC-grade water. A volume equivalent to 0.50 mg was transferred to a new, preweighed 2-mL glass vial and dried under N_2 . Finally, each sample was resuspended in 500 μL of sterile HPLC-grade water, vortexed for 1 min, and sonicated for 15 min. Two 250- μL aliquots were

transferred to new 2-mL vials, respectively, and samples were dried under N_2 . This yielded two 0.25-mg subsamples for analysis by GC-MS and ultraperformance liquid chromatography-tandem mass spectrometry (UPLC-MS/MS) as described below.

Targeted UPLC-MS/MS for phytohormone analysis of exometabolites. Subsamples of 0.25 mg were extracted in 75 μ L of a spiked methanol solution containing 100% methanol with 65.2 ng/mL abscisic acid-d6 (ABA-d6), 62.5 ng/mL salicylic acid-d6, and 90.0 ng/mL jasmonic acid-d5 (Sigma). After solvent addition, samples were placed on a shaker plate for 1 h at the highest speed setting, centrifuged at $3,500 \times g$ at 4°C for 5 min, and transferred to glass inserts. A final centrifugation step at $3,500 \times g$ for 15 min at 4°C was completed to ensure any precipitate was in the bulb of the vial insert. Five microliters of exometabolite samples was injected onto a PerkinElmer UPLC-MS/MS system, equipped with a PerkinElmer QSight LX50 solvent delivery module. An Acquity UPLC T3 column (1 by 100 mm, 1.8 μ m; Waters Corporation) was used for chromatographic separation. Mobile phase A consisted of LC-MS-grade water with 0.1% formic acid, and mobile phase B consisted of 100% LC-MS grade acetonitrile. The elution gradient was initially set at 0.1% B for 1 min, which was increased to 55.0% B at 12 min and further increased to 97.0% B at 15 min, then decreased to 0.1% B at 15.5 min. The column was reequilibrated for 4.5 min for a total run time of 20 min. The flow rate was set to 200 μ L/min, and the column temperature was maintained at 45°C. Samples were held at 4°C in the autosampler. Detection was performed on a PerkinElmer QSight 220 triple-quadrupole mass spectrometer in selected reaction monitoring (SRM) mode. The transitions monitored for each phytohormone compound can be found in Table S2. The MS was operated with an electrospray ionization (ESI) voltage of 4,500 V in positive mode and -3,500 V in negative mode. Nebulizer gas flow was set at 350 arbitrary units, and drying gas was set to 120 arbitrary units. The source temperature was 315°C, and the hot-surface induced desolvation (HSID) temperature was set to 200°C.

Nontargeted GC-MS analysis of exometabolites. Sample preparation was conducted as previously described (23, 61) and as described above for sorghum root exudates. Briefly, dried samples were resuspended in 50 μ L of pyridine containing 25 mg/mL methoxyamine hydrochloride (Sigma), centrifuged, incubated at 60°C for 45 min, vortexed for 30 s, sonicated for 10 min, centrifuged briefly, and incubated a second time for 45 min at 60°C. Samples were cooled to room temperature and centrifuged for 2 min. Then, 50 μ L of MSTFA plus 1% TMCS (Thermo Fisher) was added, and samples were vortexed for 30 s, centrifuged, and incubated a third time at 60°C for 35 min. Samples were cooled to room temperature and centrifuged, and 80 μ L of supernatant was transferred to glass vial inserts within glass vials. Samples were centrifuged a final time for 10 min before analysis. Metabolites were separated with a 30-m TG-5MS column (Thermo Scientific [0.25-mm i.d., 0.25- μ m film thickness]) and detected using a PerkinElmer Clarus 690 gas chromatograph coupled to a Clarus SQ 85 mass spectrometer (PerkinElmer). Samples (1 μ L) were injected at a 10:1 split ratio onto the column with a 1.0-mL/min helium gas flow rate. The gas chromatography inlet was held at 285°C, the transfer line was held at 300°C, and the source temperature was held at 260°C. The GC oven program started at 80°C for 30s, followed by a ramp of 15°C/min to 330°C, followed by an 8-min hold. Masses between 50 and 620 m/z were scanned at 4 scans/s under electron impact ionization. Injection of QCs was analyzed after every 6th sample to ensure proper instrument function and to detect any analytical variation.

GC-MS and LC-MS data analysis. Nontargeted GC-MS data (both sorghum exudates and exometabolites) were processed within the R statistical software (62) using methods previously described (63). For GC-MS samples, .cdf files were processed through the following workflow: (i) XCMS software was used for preprocessing to identify molecular features (64), (ii) features were further normalized to total ion current (TIC), and (iii) the package RAMClustR (65) was used for clustering features into spectra and prepared for subsequent identification of spectra in RAMSearch (66) using external databases, including Golm (<http://gmd.mpimp-golm.mpg.de/>) and NIST (<http://www.nist.gov>). For exometabolite data, prior to TIC normalization, features were normalized by linearly regressing run order versus QC feature intensities to account for instrument signal intensity drift. For root exudate data, relative quantitation was also normalized by root weight. Targeted quantification of sugars from the GC-MS data was performed using Chromeleon 7.2 (Thermo Fisher). The integrated peak area for each sugar was normalized to its corresponding internal standard, with the exceptions that arabinose- $^{13}C_5$ was used for xylose, glucose- $^{13}C_6$ was used for mannose, and sucrose- $^{13}C_{12}$ was used for trehalose and maltose. Quantification was determined using a linear regression of an 8-point standard curve for each sugar. Final concentrations were normalized to root weight. For phytohormone analysis of exometabolites, LC-MS data were processed using Simplicity 3Q (v1.5) (PerkinElmer) bioinformatics software for sample processing. Briefly, the peak area for each phytohormone compound was normalized to internal standard peak area, and quantification was assessed using a linear regression against an external calibration curve. Exudate and exometabolite data are provided in File S1.

Soil DNA extraction and library preparation. Total genomic DNA was extracted from the microcosms using the Zymo Quick-DNA fecal/soil microbe microprep kit. 16S rRNA gene amplicon sequencing was performed on the Illumina MiSeq using 251-bp paired-end reads and the Earth Microbiome Project primers 515F/806R (1), for an average of more than 30,000 reads per sample (see File S2 for individual sample data). The 16S rRNA partial gene reads were analyzed, and reads were demultiplexed using QIIME2 (2019.10) (67). Using DADA2 (68), demultiplexed reads were denoised to produce an amplicon sequence variant (ASV) table and filtered to remove noisy sequences, chimeras, and singletons. Feature classification was completed by comparing the ASV table against the trained full-length SILVA classified (silva132.250) database for taxonomic classification. The ASV table was filtered to contain ASVs that were observed in at least 2 samples, and the output files were visualized in QIIME2. Samples L_E1_T5_A (HST day 5 rep A), L_E2_T11_A, and L_E2_T20_A (HOT days 11 and 20 rep A) yielded insufficient

sequencing results and were excluded from subsequent analyses. The ASV feature table is provided in File S2.

Metagenomics analysis. Metagenomic DNA from day 0 (control) and day 5 (control and HST) metagenomes ($n = 9$) was sequenced at the Genomics Shared Resource at the University of Colorado Cancer Center using the NovaSeq6000 platform. Metagenomic DNA from day 20 (control and HST, $n = 6$) was prepared for metagenomic sequencing using the Nextera XT low-input Illumina library creation kit, and samples were sequenced at the Department of Energy Joint Genome Institute on the Illumina NovaSeq 6000. FastQ files were trimmed using Sickle (v1.33) (69). Day 0 and day 5 reads were concatenated within each of the time points/treatments (i.e., triplicate controls at day 5) for coassembly. Day 0 and day 5 (both control and HST) coassemblies and the day 20 individual assemblies were assembled with IDBA-UD (70). Within each assembly, scaffolds greater than 2.5 kb were binned into metagenome-assembled genomes (MAGs) using MetaBAT2 (v2.12.1) (71). MAGs were assessed for completion and contamination using checkM (72). A MAG was retained if it was >48% complete with <10% contamination and was assigned quality following MIMAG guidelines (33). Using dRep (73), MAGs were dereplicated to 99% identity. MAG taxonomy was assigned using GTDB-tk (v1.5.0, R06-RS202), and taxonomic novelty was defined as the first unnamed taxonomy level (74). To obtain MAG abundance, trimmed metagenomic reads from individual samples were mapped to the dereplicated MAG set using bbmap (75) (v38.70) at $\text{minid} = 95$ and output as sam files, which were converted to sorted bam files using samtools (76) (v1.9). CoverM (v0.3.2) was used to determine MAG relative abundance as described by McGivern et al. (35). MAGs were annotated using DRAM (77). Biosynthetic gene clusters were detected using the antiSMASH webserver with default parameters (v6.0) (78). See File S2 for MAG quality, taxonomy, and mapping, File S3 for DRAM annotations, and see <https://doi.org/10.5281/zenodo.5639650> for raw annotations.

Statistical analyses. Sorghum genotype ordinations were generated using metaMDS function from the vegan (79) package in R (v4.0.2) (62) and visualized with ggplot2 (80). Metabolite loadings were calculated with the envfit function in vegan. Alpha diversity of microbial communities was calculated using the diversity function from the vegan package (79) in R (v4.0.2) (62) with Shannon's (H), Pielou's, and richness indices. To estimate beta diversity across treatments, we utilized the Bray-Curtis dissimilarity matrix, which was visualized by nonmetric multidimensional scaling (NMDS) in R with the ggplot2 package with stress of the nonparametric fit for the goodness of fit for the model. Significance of compositional differences across treatments was quantified using mrpp and the betadisper commands from the vegan package, with an ANOVA model in R. Significance between communities across time and treatment was quantified using beta-group-significance commands in QIIME2 (2019.10) (66) with a PERMANOVA model. Fold changes for exometabolite dynamics were calculated and converted to \log_2 abundances via the log function. Two-way ANOVA and pairwise comparisons were completed in GraphPad Prism (v8.2.1) using grouped analyses with Sidak's multiple comparison testing with $\alpha = 0.05$.

Data availability. The MAGs, metagenomes, and 16S rRNA amplicon sequencing reads from this data set have been deposited in National Center for Biotechnology Information under BioProject accession no. [PRJNA725542](https://www.ncbi.nlm.nih.gov/bioproject/PRJNA725542) (for biosample accession numbers, see File S2). MAGs are also provided in archive (<https://doi.org/10.5281/zenodo.5142207>). Metabolomics (exometabolites and root exudates) data have been deposited in the EMBL-EBI MetaboLights database with the identifier MTBLS3164 (81). The complete data set can be accessed at <https://www.ebi.ac.uk/metabolights/MTBLS3164>.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

SUPPLEMENTAL FILE 1, XLSX file, 0.1 MB.

SUPPLEMENTAL FILE 2, XLSX file, 2.7 MB.

SUPPLEMENTAL FILE 3, XLSX file, 2.8 MB.

SUPPLEMENTAL FILE 4, PDF file, 0.3 MB.

SUPPLEMENTAL FILE 5, XLSX file, 0.1 MB.

SUPPLEMENTAL FILE 6, XLSX file, 0.2 MB.

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