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Authors

Ginnan, Nichole

De Anda, N

Campos Freitas Vieira, Flavia

et al.

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Microbial Turnover and Dispersal Events Occur in Synchrony with Plant Phenology in the Perennial Evergreen Tree Crop *Citrus sinensis*

 Nichole A. Ginnan,^a  N. Itzel De Anda,^a  Flavia Campos Freitas Vieira,^a  Philippe E. Rolshausen,^b  M. Caroline Roper^a

^aDepartment of Microbiology and Plant Pathology, University of California, Riverside, California, USA

^bDepartment of Botany and Plant Sciences, University of California, Riverside, California, USA

ABSTRACT Emerging research indicates that plant-associated microbes can alter plant developmental timing. However, it is unclear if host phenology affects microbial community assembly. Microbiome studies in annual or deciduous perennial plants face challenges in separating effects of tissue age from phenological driven effects on the microbiome. In contrast, evergreen perennial trees, like *Citrus sinensis*, retain leaves for years, allowing for uniform sampling of similarly aged leaves from the same developmental cohort. This aids in separating phenological effects on the microbiome from impacts due to annual leaf maturation/senescence. Here, we used this system to test the hypothesis that host phenology acts as a driver of microbiome composition. *Citrus sinensis* leaves and roots were sampled during seven phenological stages. Using amplicon-based sequencing, followed by diversity, phylogenetic, differential abundance, and network analyses, we examined changes in bacterial and fungal communities. Host phenological stage is the main determinant of microbiome composition, particularly within the foliar bacteriome. Microbial enrichment/depletion patterns suggest that microbial turnover and dispersal were driving these shifts. Moreover, a subset of community shifts were phylogenetically conserved across bacterial clades, suggesting that inherited traits contribute to microbe-microbe and/or plant-microbe interactions during specific phenophases. Plant phenology influences microbial community composition. These findings enhance understanding of microbiome assembly and identify microbes that potentially influence plant development and reproduction.

IMPORTANCE Research at the forefront of plant microbiome studies indicates that plant-associated microbes can alter the timing of plant development (phenology). However, it is unclear if host phenological stage affects microbial community assembly. Microbiome studies in annual or deciduous perennial plants can face difficulty in separating effects of tissue age from phenological driven effects on the microbiome. Evergreen perennial plants, like sweet orange, maintain mature leaves for multiple years, allowing for uniform sampling of similarly aged tissue across host reproductive stages. Using this system, multiyear sampling, and high-throughput sequencing, we identified plant phenology as a major driver of microbiome composition, particularly within the leaf-associated bacterial communities. Distinct changes in microbial patterns suggest that microbial turnover and dispersal are mechanisms driving these community shifts. Additionally, closely related bacteria have similar abundance patterns across plant stages, indicating that inherited microbial traits may influence how bacteria respond to host developmental changes. Overall, this study illustrates that plant phenology does indeed govern microbiome seasonal shifts and identifies microbial candidates that may affect plant reproduction and development.

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Address correspondence to M. Caroline Roper, mcroper@ucr.edu.

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Plant phenology, the periodic timing of plant life cycle events, is innately linked to exogenous climatic variables that affect plant development, such as temperature, photoperiod, and nutrient and water availability, as well as other abiotic and biotic factors (1, 2). Additionally, endogenous genome-encoded factors such as dynamic internal photosynthate source-sink pathways, intricate phytohormone signaling networks, and other developmental regulatory processes mediate the transition between phenological stages (3, 4). The timing of specific developmental stages, such as flowering, can determine a plant's geographic distribution range as well as determine crop yield and productivity (5, 6). Alterations in plant phenology can also have a cascading effect on the fitness of organisms that depend on those specific plants for nutrient acquisition, such as pollinator species (7–9).

Citrus is a significant economic crop and provides several health benefits because of the myriad of nutrients, antioxidants, vitamins, minerals, and dietary fiber found in fresh and juiced citrus fruits (10–12). Citrus varieties are grown across the globe, and because of this, citrus phenology is well characterized to guide management strategies of different varieties for specific climatic conditions. Phenological modeling of citrus has focused primarily on buds, flowers, and fruit and is used to predict bloom time across different growing regions (13). This has implications for protecting flowers from floral pests and pathogens by allowing growers to time spray applications in an informed manner (14). Citrus flowers are a significant source of nectar related to honey production, particularly in California's Central Valley. As such, bloom timing models are also important for the beekeeping industry (15). In addition, bloom time and duration models can be extrapolated to predict fruit set (16), and these performance models can provide yield predictions.

Soil and rhizosphere microbiomes can drive changes in flowering time in the herbaceous perennial plant *Boechea stricta*, a wild relative of *Arabidopsis* (17, 18), and affect other aboveground plant traits in the annual plant system *Brassica rapa* (19, 20). However, questions about microbiome shifts associated with transitions between phenological stages have not been addressed in perennial trees, particularly domesticated evergreens like citrus (21). Citrus phenology models primarily take into account temperature and number of degree days above a certain threshold temperature (16) but, to the best of our knowledge, have not incorporated studies on the microbial communities associated with transitions across phenological stages. The citrus microbiome is an emerging prototype for understanding microbial contributions to plant health in a perennial arboreal crop system (22–26). Due to its well-defined phenology, citrus is an ideal system to investigate the interplay between host phenology and microbial community composition.

Several seminal studies in annual and short-lived perennial plants have characterized changes in rhizosphere and root microbiome composition across plant developmental cycles, suggesting that host phenology drives these alterations (27–32). However, Dombrowski et al. suggests that initially microbiota are sequentially acquired resulting in community changes as the host ages but eventually the microbiome matures and stabilizes, functioning independently from host development (33). Another recent study supports the idea that time is a stronger predictor of microbiome composition than plant developmental stage (34). This prompts discussion on whether these community shifts are a consequence of tissue age and a microbiome maturation process or if these changes are driven by plant phenology. In addition to producing and maturing leaves and roots throughout the year, long-lived evergreen perennial plant systems retain mature leaves for 1 to 3 years (35, 36), which allows for selection of leaf tissues of similar age and developmental cohort across phenophases. Because of these features, we utilized this system to help decouple tissue age from host

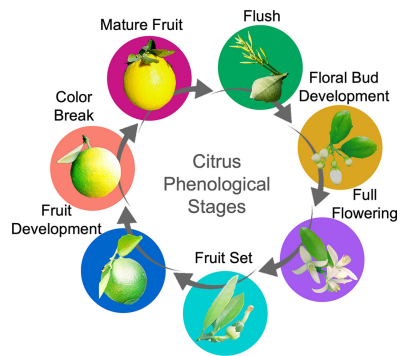


FIG 1 Citrus phenological stages. Cyclic seasonal development of *Citrus sinensis*.

phenological effects and tested the hypothesis that host phenology acts as a driver of community compositional shifts within the aboveground (foliar) and belowground (root) microbiomes of citrus. Indeed, we determined that the significant shifts in both diversity and composition of the microbial community structure were driven primarily by host phenological stages and not exogenous environmental factors such as rainfall, hours of irrigation, or temperature. Foliar communities were more affected by host phenology than root microbiomes, which were comparably more stable. Interestingly, major alterations in foliar microbial community composition correlated with the shifts in source-to-sink pathways of carbohydrate transit, namely, during the transition from floral bud development to full flowering to fruit set. More specifically, subsets of these taxa displayed temporal turnover patterns indicating that specific taxa were enriched as trees shifted to reproductive growth associated with fruit production. We also observed taxa typically associated with pollinator species that were substantially enriched only during flowering, suggesting that these microbes were introduced into the foliar microbiome as microbial immigrants via an insect-mediated dispersal mechanism.

In agricultural plant systems, comprehensive microbiome studies allow researchers to place an emphasis on how the microbiome as a whole functions to promote overall plant health by a variety of mechanisms, such as enhancing nutrient uptake or resisting pathogen ingress to promote a sustainable agroecosystem. Uncovering links between plant phenology and shifts in microbiome structure is the first step toward a mechanistic understanding of microbiome resilience over cyclical development in a perennial plant host. In addition, this can further serve as the foundation to understanding how the microbiome responds to changes in host development and, in turn, if microbiome community structure can influence host phenological transitions.

RESULTS

Significant shifts in alpha diversity occur across phenological stages. We focused our study on seven citrus phenophases that included spring vegetative shoot flush, referred to as “flush” (F), early floral bud break and development (FB), full flowering (FF), fruit set (FS), exponential fruit growth and development, referred to as “fruit development” (FD), color break (CB; initiation of fruit maturation), and mature fruit (MF). Citrus phenological stages can overlap on individual trees, and some stages span multiple months; thus, some stages include multiple months of sampling (Fig. 1 and see Fig. S1 in the supplemental material). Initially, we assessed alpha diversity or the diversity within a sample, in this case the number of operational taxonomic units (OTUs). Overall, bacterial and fungal leaf microbiomes had the most significant shifts in alpha diversity across phenological stages compared to those of the root microbiomes. Specifically, alpha diversity in both the leaf bacteriome and mycobiome remained consistent as trees transitioned from leaf flush to flowering (floral bud development and full flowering). Following full flowering, there was a significant increase in alpha diversity in the leaf bacteriome and mycobiome at fruit set (Fig. 2a and b). Species richness

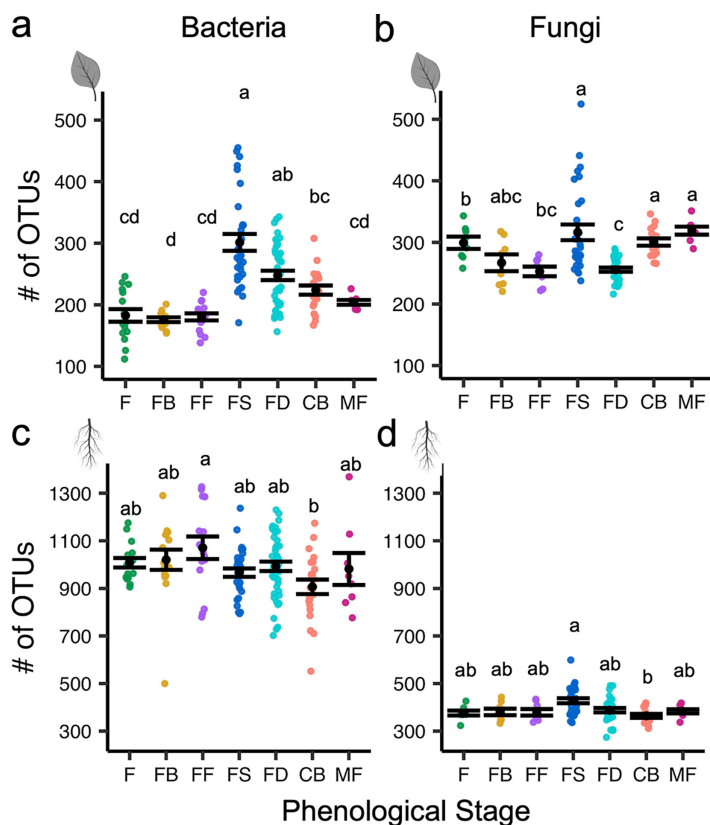


FIG 2 Species richness varies across phenophases. Alpha diversity plot of richness (no. of OTUs) for individual sampling events for bacterial leaf (a), fungal leaf (b), bacterial root (c), and fungal root (d) communities. Black points, “•,” represent the mean, and error bars represent standard error. Letters indicate a significant difference of $P \leq 0.05$, determined using a Kruskal-Wallis test, with a pairwise Dunn’s test and correcting for multiple comparisons with Holm’s method. Phenological stages on the x axis include flush (F), floral bud break (FB), full flowering (FF), fruit set (FS), fruit development (FD), color break (CB), and mature fruit (MF).

within the leaf bacteriome significantly decreased when trees transitioned from fruit growth and development to color break and mature fruit stages.

Despite being relatively stable across the study, root bacteriome alpha diversity peaked during full flowering. Similar to the overall leaf microbiome, the root mycobiome had the highest alpha diversity during fruit set (Fig. 2c and d). Our study did not discriminate between rhizoplane and endophytic root microbiota, nor was it possible to select feeder roots of a specific age cohort. Future work that separates these compartments in similarly aged roots may reveal more finely resolved shifts in species richness associated with these root environments.

Host phenological stage is a major determinant of community composition.

Although climatic variables can be difficult to uncouple from plant development variables, the greatest amount of the variation in the data was attributed to the host phenological stage for all four communities (permutational multivariate analysis of variance [PERMANOVA], $P \leq 0.001$, $r^2 = 0.062$ to 0.187) (Table 1, Fig. 3). Time (i.e., sample year) had less impact than phenology on beta diversity (variation in taxa abundance and composition between samples) across all communities (Table 1, Fig. S2). Interestingly, the community composition (beta diversity) of leaf bacteriome and mycobiome was influenced by host phenology more than that of root communities, indicating that changes in host phenology had a larger influence on diversity within foliar microbiomes than in root microbiomes. Specifically, a principal coordinate analysis of weighted UniFrac distances indicated significant clustering of individual microbial communities by phenological stage (Fig. 3). In a pairwise comparison of community

TABLE 1 Beta diversity analysis (PERMANOVA) results for all variables^a

Characteristic	Leaf bacteria				Leaf fungi			
	df	F	r ²	P	df	F	r ²	P
Phenology	6	6.706	0.187	0.001	6	3.893	0.163	0.001
Sample yr	1	6.205	0.029	0.001	1	11.065	0.077	0.001
Fertilizer	1	5.854	0.027	0.001	1	2.579	0.018	0.003
Avg. temp	1	5.984	0.028	0.001	1	1.762	0.012	0.041
Hours of irrigation	1	2.866	0.013	0.008	1	3.787	0.026	0.001
Total rain	1	3.802	0.018	0.001	1	2.072	0.014	0.023
Phenology: sample yr	5	3.557	0.082	0.001	1	2.141	0.015	0.012
Avg. temp: hours of irrigation	1	1.972	0.009	0.038	1	1.402	0.010	0.119
Residuals	130		0.605		95		0.664	
Characteristic	Root bacteria				Root fungi			
	df	F	r ²	P	df	F	r ²	P
Phenology	6	1.676	0.062	0.001	6	2.038	0.098	0.001
Sample yr	1	3.749	0.023	0.001	1	4.044	0.032	0.001
Fertilizer	1	1.452	0.009	0.036	1	3.890	0.031	0.001
Avg. temp	1	0.962	0.006	0.497	1	2.151	0.017	0.008
Hours of irrigation	1	2.077	0.013	0.001	1	2.072	0.017	0.011
Total rain	1	1.326	0.008	0.061	1	2.523	0.020	0.005
Phenology: sample yr	5	1.299	0.040	0.002	1	2.193	0.018	0.008
Avg. temp: hours of irrigation	1	0.986	0.006	0.483	1	1.174	0.009	0.211
Residuals	136		0.834		95		0.758	

^aTest was performed using the `vegan::adonis2` function and square-root transformed weighted UniFrac distances. Bold text indicates the variable with the greatest r² value (phenology).

compositional differences between each phenological stage, the leaf bacteriome had the greatest number of significant adjusted *P* values, with 21 of the 21 pairwise comparisons being significantly different, followed by the leaf mycobiome (19 of 21 comparisons) (Table S2). Root communities had fewer significantly different pairwise comparisons (root mycobiome = 5/21, root bacteriome = 8/21).

Rainfall, fertilizer applications, temperature, and irrigation hours fluctuated across our sampling period (Fig. S1, Table S3). Rainfall was sparse in this sample location (the Central Valley of California), ranging from 0.00 to 2.55 inches each month (Fig. S1b, Table S3), and total rainfall was a minor determinant of community structure across all four communities, explaining only 0.8 to 2.0% of the variation (PERMANOVA, $P \leq 0.001$ to 0.061, $r^2 = 0.008$ to 0.020) (Table 1). Similarly, fertilizer application describes a small percentage (0.9 to 3.1%) of the variation in the data for all four communities examined. We evaluated temperature based on the average temperature and interactive effects it might have with water availability (hours of irrigation) in order to capture the full range of conditions that could affect microbial community composition. Temperature had a minor impact on communities, as this factor describes only 0.6 to 2.8% of the variation in the data that include temperature as an interaction factor. In addition to phenology, interactions between phenology and sample year were a driving factor of leaf bacterial community composition, explaining 8.2% of the changes across the data (PERMANOVA, $P \leq 0.001$, $r^2 = 0.082$).

Taken together, these beta diversity analyses indicate that plant phenological stage was the major driving factor in community composition for bacterial and fungal communities associated with leaves and roots. Significant compositional shifts were also visible at the phylum level, particularly in the leaf bacterial community (Fig. S3). Other covariates tested (irrigation, mean temperature, fertilizer applications, rainfall, and sample year) were minor or insignificant contributors to citrus-associated leaf and root microbiome composition.

Stable, phylogenetically conserved microbial signatures across phenophases.

We identified core microbial taxa for each of our seven phenological stages. Our core bacterial and fungal leaf and fungal root microbiomes include genera that were greater than 0.01%, and core root bacteriome included genera greater than 0.1%, relative abundance in at least 75% of the samples within a phenological stage. All of our

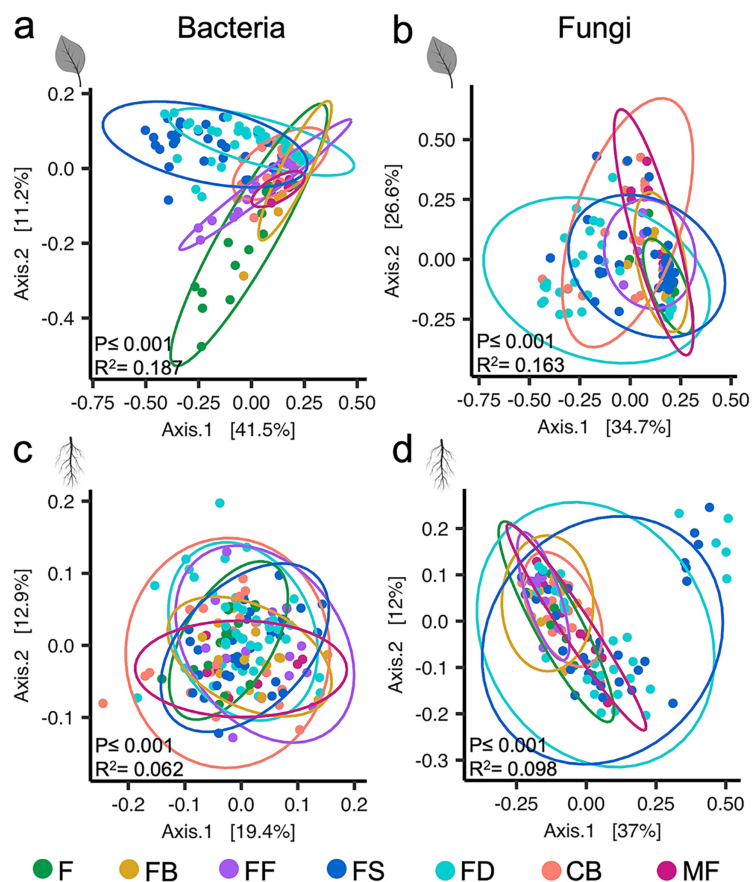


FIG 3 Host phenology affects community diversity and composition. Beta diversity PCoA plots of bacterial leaf (a), fungal leaf (b), bacterial root (c), and fungal root (d) communities. Points are colored by phenological stage and represent a complete community from a single leaf or root sample. Ellipses represent 95% confidence intervals. The P values and r^2 values were obtained using a PERMANOVA (Adonis). Phenological stages include flush (F), floral bud break (FB), full flowering (FF), fruit set (FS), fruit development (FD), color break (CB), and mature fruit (MF).

downstream analyses used genera that met our core taxa cutoffs in at least one phenophase. We assessed our core taxa and separated them into three categories: (i) high stability, defined as core member of six or more phenophases, (ii) medium stability, core member of three, four, or five phenophases, or (iii) low stability, core member of two or fewer phenophases. We determined that of the identified core there were 3 (5.2%) leaf bacterial, 8 (30.7%) leaf fungal, 62 (70.4%) root bacterial, and 22 (61.1%) root fungal core genera that had high stability across phenophases (Fig. 4, Fig. S4). This suggests that both bacterial and fungal root communities have a substantially greater number of consistent or stable microbial features across the developmental cycle. However, our experimental design did not differentiate between endophytes versus epiphytes and, thus, may have missed some fine resolution microbial community shifts occurring between the endosphere and episphere. There were two bacterial (*Pseudomonas* and *Sphingomonas*) and one fungal (*Aureobasidium*) genera that were highly stable in both roots and leaves (Fig. 4c and d).

A phylogenetic analysis of the core genera indicates that both bacterial and fungal root communities were rich in highly stable and phylogenetically diverse core taxa (Fig. S4). Root core genera from the bacterial clade *Alphaproteobacteria* (class) and the fungal family *Pleosporomycetidae* were all or nearly all binned as highly stable, indicating that genera in these clades were consistently high in relative abundance across all phenological stages. Medium- and low-stability core genera appear randomly dispersed across the root community phylogeny, with no obvious patterns.

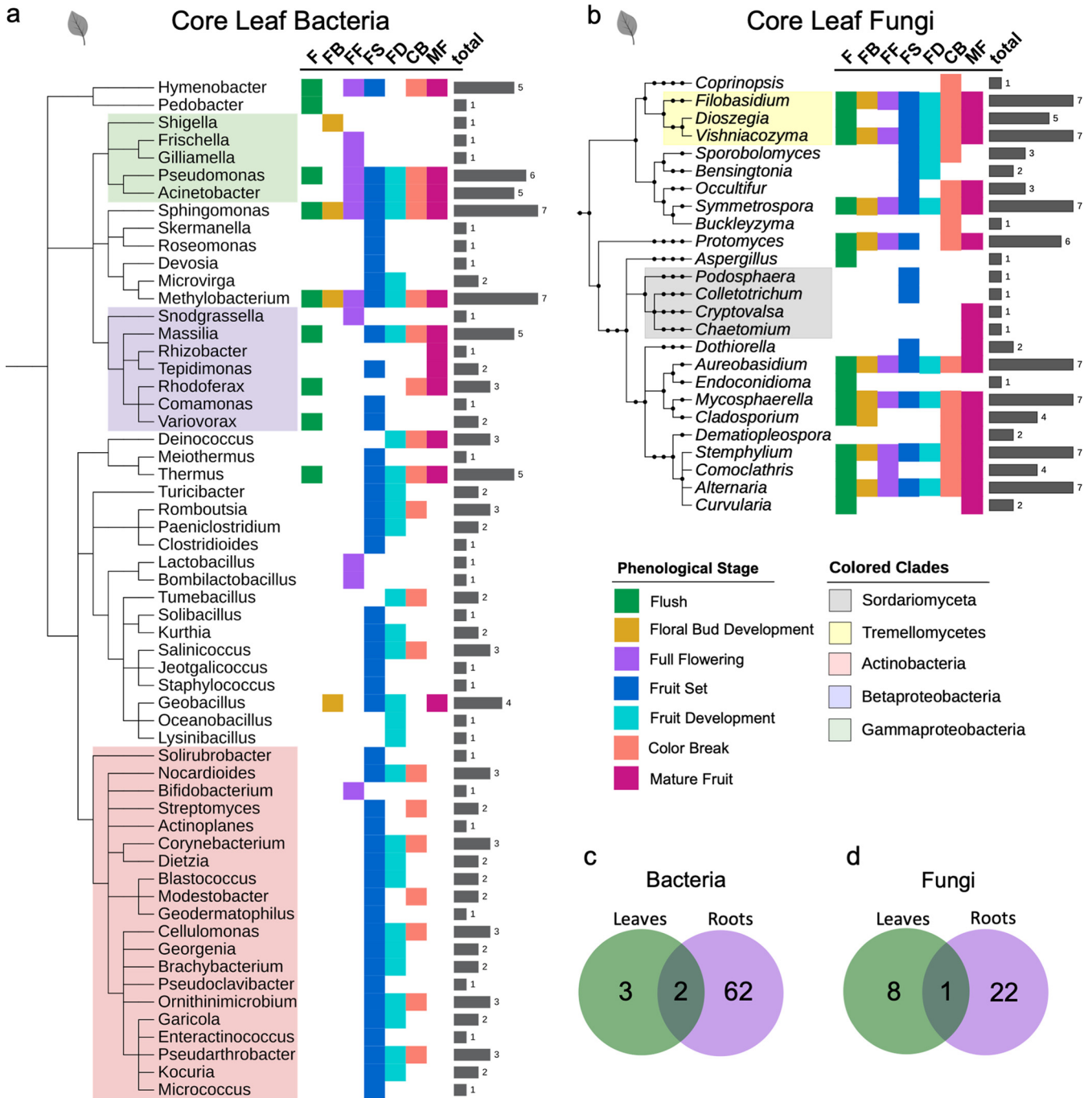


FIG 4 A subset of leaf bacterial and fungal core genera have phylogenetically conserved phenological association patterns. Phylogenetic trees of leaf bacterial (a) and fungal (b) genera that were core ($\geq 0.01\%$ relative abundance and $\geq 75\%$ prevalence) to one or more stages. Colored squares indicate a genus is core to flush (green), floral bud development (gold), full flowering (purple), fruit set (blue), fruit development (light blue), color break (salmon), and/or mature fruit (magenta). Gray bars indicate the total number of phenological stages during which each genus is core. Venn diagrams show the number of highly stable core bacterial (c) and fungal (d) genera in leaf and root communities.

However, leaf bacterial and fungal core community phylogenetic trees contained high, medium, and low stability patterns at the class and phylum levels (Fig. 4a and b). All core genera in the fungal class Tremellomycetes had medium to high stability. In contrast, all core genera in the fungal class Sordariomycetes had low stability across phenophases and met the defined core cutoffs only during fruit set or mature fruit stages. The leaf taxa within the bacterial class Gammaproteobacteria consisted of genera with high, medium, and low stability across the phenophases. Interestingly, all the

Gammaproteobacteria were core members of the full flowering or floral bud development microbiomes regardless of their stability in other phenophases. Another distinct phylogenetic pattern observed in the leaf community was genera in the bacterial phylum *Actinobacteria* that had low or medium stability across all phenophases. However, 95.0% of core genera in the *Actinobacteria* clade were core during fruit set and/or fruit development. The only exception to this within the *Actinobacteria* clade was *Bifidobacterium*, which was associated only with full flowering and was not a core member of fruit set or fruit development microbiomes (Fig. 4a). Lastly, the leaf bacterial class *Betaproteobacteria* contains low- to medium-stability core genera with the most dispersed stage associations.

Overall, these data indicate that root bacterial and fungal communities have greater stability across phenophases than those of leaves (Fig. 4c and d). Additionally, core taxa had phylogenetically related trends within the high-, medium-, and low-stability classifications, indicating that conserved, vertically descended microbial traits may play a role in determining bacterial and fungal associations across phenophases, particularly in aboveground leaf tissue.

Specific taxa were enriched in the foliar microbiome across the flowering phenophases. We completed a genus-level differential relative abundance analysis on our list of core taxa that were $\geq 0.01\%$ relative abundance and $\geq 75\%$ prevalence in one or more phenophases. Our differential relative abundance analysis can determine finer-scale phenophase associations beyond just classification as a core microbiome member by looking for increases in relative abundance, proportionate to other members of the microbial community (enrichments), during specific phenophases. Ecologically dominant taxa (taxa with high abundance relative to that of other species in the community) are predicted to have a proportionately larger contribution to community function. Among all the phenophases, those associated with flowering (floral bud development and full flowering) had striking microbial enrichments, particularly among the leaf bacteria. *Acinetobacter* was a core member of five phenophases but was significantly enriched during full flowering compared to other phenophases (Fig. 5a). *Acinetobacter* had a gradual enrichment from flush and floral bud development to full flowering. This gradual enrichment signature indicates that *Acinetobacter* was present throughout the year but has a high temporal turnover rate that is in sync with the transitions from flush to floral bud development and then to full flowering.

We also observed bacteria that were sharply enriched during full flowering rather than undergoing gradual enrichments over the phenophases that lead up to full flowering (flush and floral bud development). These include *Snodgrassella*, *Frischella*, *Gilliamella*, and *Bifidobacterium* (Fig. 5b to e). The sharp enrichment patterns during full flowering suggest that these taxa were introduced into the community via a dispersal event.

Foliar microbial depletions associated with flowering. We also identified bacterial leaf genera that had significant depletions (reduction in abundance relative to that of other taxa) during floral bud development and/or full flowering (Fig. 6a to h). Four *Actinobacter* genera, *Corynebacterium*, *Dietzia*, *Georgenia*, and *Ornithinimicrobium*, were significantly depleted during floral bud development and full flowering (Fig. 6a to d). *Bacillus*, *Methylobacterium*, *Romboutsia*, and *Sphingomonas* also significantly decreased in relative abundance during floral bud development and/or full flowering (Fig. 6e to h). For all differentially abundant genera, including bacteria and fungi, across all phenophases, see Fig. S5 and Table S4.

Microbe-microbe interactions contribute to phenophase-specific community structure. We performed a network analysis on the foliar bacterial communities from all samples. We focused on significantly enriched and/or depleted populations and populations with direct connections or putative first-degree interactions (neighbors). The goal of this approach was to give a broad overview of bacterial interactions across phenophases and identify taxa that potentially interact with specific phenophase-enriched taxa and potentially play a role in observed seasonal community compositional shifts. *Rhizobium*, *Sphingomonas*, an unknown bacterium, an unknown *Bacillaceae* (family), *Acinetobacter*,

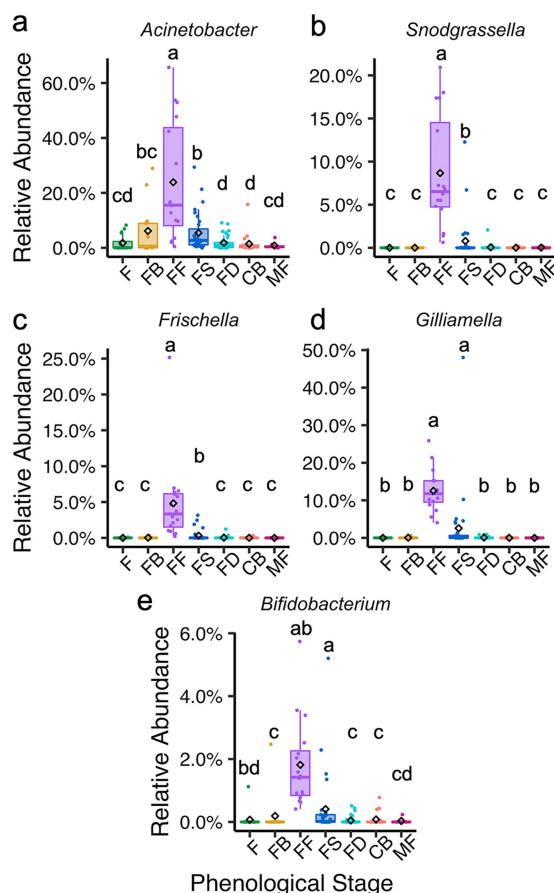


FIG 5 Core leaf bacteriome members enriched during full flowering. Gradually (a) and suddenly (b to e) enriched core ($\geq 0.01\%$ relative abundance and $\geq 75\%$ prevalence) taxa during full flowering. The diamond symbol indicates the mean relative abundance. Letters indicate significant differences of $P \leq 0.05$, determined using DESeq2 GLM, Wald test with FDR adjustment. Phenological stages on the x axis include flush (F), floral bud break (FB), full flowering (FF), fruit set (FS), fruit development (FD), color break (CB), and mature fruit (MF).

and *Romboutsia* have the highest normalized betweenness centrality scores ranging from 0.110 to 0.187. Betweenness centrality is a proxy for influence within a network because it measures how often a particular node (i.e., taxon) is the shortest connection or bridge between two other nodes. These high betweenness centrality scores and placement within the network indicate that these genera are potentially keystone taxa that may perform a stabilizing role in the microbial communities across phenological transitions and events (Fig. 7, red nodes). Groups of taxa connected by putative positive interactions cluster together to form distinct modules. These modules are separated by putative negative interactions. Our analysis organized bacterial taxa that were enriched in fruit set and fruit development into a single highly connective community module (cluster) (Fig. 7, blue nodes). This suggests that fruit set- and fruit development-associated microbiomes are compositionally similar and few microbe-microbe interactions change during the transition from fruit set to fruit development. Leaf bacteria associated with flowering also formed a module within the network (Fig. 7, purple nodes). Specific bacteria within the fruit set/development and flowering modules also interact with taxa that were enriched in the other four phenophases, which cluster together into a third module (Fig. 7, gray nodes). Overall, these predicted positive interactions represent inter- or codependent microbe-microbe relationships, and the putative negative interactions indicate potential direct (e.g., antibiosis) or indirect (e.g., resource exclusion) competition. These predicted microbe-microbe interactions within the microbiome likely affect community composition

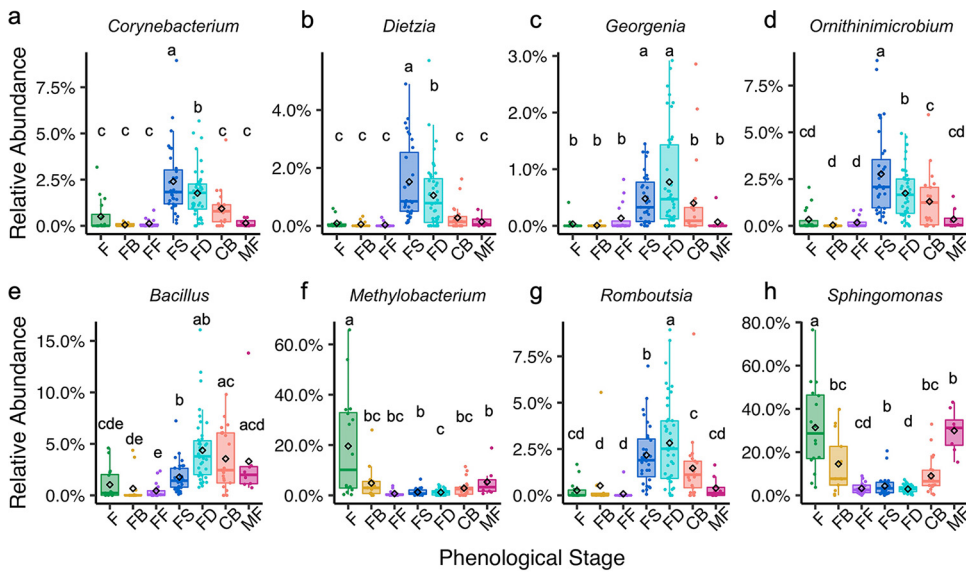


FIG 6 Core leaf bacteriome members depleted during full flowering. The diamond symbol indicates the mean relative abundance of core ($\geq 0.01\%$ relative abundance and $\geq 75\%$ prevalence) taxa across stages. Letters indicate significant differences of $P \leq 0.05$, determined using DESeq2 GLM, Wald test with FDR adjustment. Phenological stages on the x axis include flush (F), floral bud break (FB), full flowering (FF), fruit set (FS), fruit development (FD), color break (CB), and mature fruit (MF).

in addition to the exogenous influences of abiotic environmental conditions and biotic host physiological factors (e.g., carbon availability).

DISCUSSION

The majority of studies examining how plant developmental stage affects the plant's microbiome have focused on bacteria associated with the rhizosphere of annuals or herbaceous perennials such as maize (37, 38), rice (28), sorghum (27, 32), wheat (39), *Arabidopsis* (29), and *Boechera* (30). These important studies indicate that rhizosphere-associated microbiomes can shift in association with plant developmental stages in both domesticated and wild plants that have short-lived aboveground tissues. Studies of the endophytic xylem sap microbiome in grapevine, a deciduous perennial, also showed that microbial shifts were linked to changes in phenological stage (40). However, much less is known about how overall plant phenology affects above- and belowground microbiomes of evergreen woody perennials that have lifespans that can be decades long and can retain their leaves for multiple years, compared to annuals or deciduous perennials that produce and shed all their leaves each season. Here, we investigated microbiome compositional dynamics in above- and belowground tissues of mature 20-year-old *Citrus sinensis* trees to determine if temporal microbiome fluctuations were associated with host phenological events. The unique contribution of our research was the separation of leaf development from tree phenology. We did this by analyzing the changes in the foliar microbiome on fully mature leaves, which developed in the leaf cohort from the previous year, in relation to the phenological stages of the current year. Thus, the leaves were exposed to the same starting inoculum, minimizing the bias of any potential priority effects (i.e., order of arrival).

Our results indicate that the phyllosphere microbiome has an active and dynamic relationship with host phenology. More specifically, microbial shifts occurred as trees transitioned from the spring leaf flushing stage and entered flowering. The transition from spring flush to floral bud development and full flowering aligns with important transitions in source-to-sink transport of photosynthate in the tree (4). During foliar flushing periods, young leaves are a primary carbohydrate sink as they rapidly expand and mature. This source-to-sink transport of photosynthate shifts during floral bud break and development, when mature leaves

taxa in the community, as well as ecological mechanisms that contribute to plant microbiome assembly, such as microbial species turnover and dispersal (41–43). Interestingly, microbial enrichment and depletion patterns of specific taxa suggest that microbial species turnover and dispersal events within the citrus microbiome occur in sync with phenological stage transitions. These enrichment/depletion patterns for specific taxa were more apparent in leaves than in the root compartment. Specifically, the bacterial genus *Acinetobacter* was enriched in leaves as trees transitioned from spring flush to floral bud development and peaked in relative abundance during full flowering, which is when mature leaves shift toward becoming source tissues for developing flowers and fruit. This may create a microenvironment that selects for an increase in relative abundance of these taxa when carbohydrate is translocating out of the leaves. Plant-associated *Acinetobacter* spp. have plant growth-promoting properties that include antagonism toward fungi (44) and the ability to solubilize phosphate and produce the plant hormone gibberellic acid (45, 46). *Acinetobacter* spp. are highly abundant in the floral nectar microbiome of *Citrus paradisi* and other plant species (47, 48) and were identified as a core member of the grapevine xylem sap microbiome (40). Their significant increase in relative abundance in the leaf microbiome at the time of flowering in citrus suggests a potential synergy between the foliar and floral microbiomes. *Acinetobacter* was also predicted to be a keystone taxon and was a major link between the flowering community and fruit set/development community clusters in our network. This enrichment in *Acinetobacter* may be simply due to selection imposed on the microbial community by the local plant environment, but it is tempting to speculate that *Acinetobacter* spp. provide an exogenous service to the plant by producing gibberellic acid and biologically available phosphorus to promote flowering that is in phase with its host's phenological development. This hypothesis that the plant environment selects for taxa within its foliar microbiome that, in turn, promote its own reproductive growth warrants future inquiry. Specific bacterial enrichments also occurred at bloom time in grapevine, further supporting the speculation of a microbial role in plant growth development that is in sync with plant developmental stage (40).

We also observed signatures that indicate that specific taxa were depleted in relative abundance during flowering but enriched during fruit set. Phylogenetic reconstruction of these taxa indicated that the majority of the taxa belonging to the *Actinobacteria* phylum (19 of the 20 genera) were significantly depleted during flowering but subsequently enriched when trees began to set fruit. This phylogenetic conservation of depletion/enrichment patterns within the *Actinobacteria* clade indicates that this is a nonrandom fluctuation within the microbiome structure associated with the transition from flowering to fruit production. As citrus trees set fruit, the fruits themselves begin exporting and importing hormones, such as indole acetic acid (IAA) and cytokinins, respectively (3). This results in a change in hormone levels in leaves as well. Phytohormones can affect microbiome composition by being directly utilized as a carbon source or through other undefined mechanisms (49, 50). We speculate that these hormonal shifts may place selective pressure on the foliar microbial community and lead to significant enrichments of *Actinobacteria* during fruit set and development. Specific differentially abundant taxa within the *Actinobacteria* clade that followed this pattern included *Corynebacterium*, *Dietzia*, *Georgenia*, and *Ornithinimicrobium*. Members of these genera can fix nitrogen and produce IAA, both of which are important supporters of fruit development (51–54). Thus, it is tempting to speculate that these taxa could play a role in coregulating fruit development in a manner that is synergistic with the host's production of reproductive hormones. The biological role of *Actinobacteria* in the foliar microbiomes of plants is not well understood, but overall species richness was conserved across all phenological stages, except for fruit set, indicating that this phenological stage allows for microbial enrichments of specific taxa, particularly those belonging to the *Actinobacteria* phylum.

Genera outside the *Actinobacteria* clade were also depleted in leaves during floral

bud development and full flowering, including *Bacillus*, *Methylobacterium*, *Romboutsia*, and *Sphingomonas*. Notably, *Romboutsia* and *Sphingomonas* were predicted to be keystone taxa in our microbe-microbe interaction network analysis, and all are in the top 20% highest betweenness centrality scores. Keystone taxa play a stabilizing role in microbial communities. Depletion of these taxa during flowering may have cascading effects that influence microbial species turnover by allowing other taxa, such as the *Actinobacteria*, to flourish during subsequent developmental stages, like fruit set. This suggests that microbial turnover in the foliar microbiome is mediated by selective pressures imposed by the plant developmental stage in conjunction with microbe-microbe interactions to modulate community diversity and composition. Further experimentation is needed to confirm these interactions.

Microbial dispersal events can drive microbial turnover and influence the relative abundance of endogenous taxa in the community. Full flowering is a dynamic phenophase in plant development where there are frequent interactions between plants and pollinator species that rely on floral resources, like nectar and pollen. These macro-level interactions can also have effects at the microorganismal level. Pollinator (e.g., bee) visitation alters flower surface, nectar, and subsequent seed microbial community composition (55–58). During flowering, we observed striking microbial enrichments of bacteria taxa belonging to the *Betaproteobacteria* and *Gammaproteobacteria* clades that include *Gilliamella*, *Snodgrassella*, *Bifidobacterium*, and *Frischella*. These enrichments in these anaerobic taxa were unique to the flowering phenophase and quickly declined following flowering, suggesting they are immigrants to the community and not endogenous members of the native microbiome. Moreover, these anaerobic taxa are prevalent in the bee gut microbiome (59). We hypothesize that these taxa immigrated into the citrus microbiome via a dispersal event during bee visitation. We consider this external influence host phenology-associated because phenophase-specific plant morphology (i.e., flowers) regulate this diffuse interaction. Bacteria can be introduced to plants by bees and potentially migrate from the flower to the vascular bundles resulting in systemic movement within the plant (60–62). Leaf carbohydrate content is highest during flowering, which may promote the growth of these fermenting bacteria (4). Notably, *Bifidobacterium* was the only core leaf genus from the *Actinobacteria* phylum that was enriched during flowering, whereas the other 19 *Actinobacteria* taxa were depleted during flowering. This further supports the hypothesis that *Bifidobacterium* was introduced via a dispersal event and is not part of the endogenous microbiota like the other taxa in the plant-associated *Actinobacteria* clade. Nectar-inhabiting bacteria can influence nectar volatile profiles that, in turn, influence pollinator visitation preferences (63), and it would be interesting to determine if these putative immigrants contribute to shifts in nectar volatile profiles that affect bee feeding behaviors.

The next frontier in microbiome research is to determine the functional roles that microbes play in microbe-microbe and host-microbiome interactions. Martiny et al. found that conservation of microbial traits was linked more strongly to vertical phylogenetic relatedness of the microorganisms within a microbiome than to traits that are shared among taxa by horizontal gene transfer (64). Similarly, we also observed phylogenetic conservation within microbial enrichments, suggesting that those groups of related organisms play similar functional roles during specific phenophases or across several phenophases. We speculate that taxa with high stability across phenophases may serve a community-stabilizing function, while low stability or phenophase-specific core microbes likely have more specialized, transient roles in the community. Because microbes can alter host phenology (18, 21, 65–68), which is a critical factor in plant health and productivity, incorporation of microbial presence/absence and patterns of enrichment into plant phenological models may improve phenophase timing predictions once the functional roles of these microbes are determined. This information could also lead to the commercialization of biofertilizers for horticultural purposes that could be applied at specific plant life stages to enhance crop productivity.

MATERIALS AND METHODS

Field site and study design. Our longitudinal study included eight Late Navel Powell sweet orange trees (*Citrus sinensis* L. Osbeck, cv Powell) on Carrizo rootstock grown at the University of California (UC) Lindcove Research and Extension Center (LREC) in Exeter, CA. Trees were planted in 1997 (20 to 22 years old at time of sampling) and managed with conventional farming strategies similar to those of industry orchards. Sampling occurred at 20 time points (monthly for first sample year, targeted sampling for second year) from July 2017 to April 2019. Prior to each monthly sampling, tagged trees were visually assessed and developmental stages were recorded. We recorded the dominant stage occurring during each sampling event, but note that phenophases can overlap, particularly during transitions. We avoided sampling during transitions between stages when possible. Seven major phenological stages were used for categorization in this study. These included (i) spring vegetative shoot flush (soft “feather” flush, early elongation and leaf emission [phase 2: stages V2 to V3; see reference 69]), referred to as “flush” (F; February to March), (ii) early floral bud break and development, referred to as “floral bud development” (FB; March), (iii) full flowering (FF; April), (iv) fruit set (FS; May to July), (v) exponential fruit growth and development, referred to as “fruit development” (FD; August to October), (vi) color break (CB; initiation of fruit maturation; November to December), and (vii) mature fruit (MF; January) (Fig. 1) (70). In California, sweet orange trees can have foliar flushing events in the spring, summer, and fall. However, spring flush events are robust and synchronized among trees, whereas summer and fall flushes are more sporadic and weaker. Therefore, we focused on the spring vegetative flush for this study and sampled when all parts of the canopy for all trees had new flush. At the time of all other sampling events, there was only 0 to 10% new flush intermittently dispersed across trees and individual canopies. Citrus root flushes are known to occur in June and November in California but are difficult to observe in large field trees because they are under the soil; therefore, the exact timing of root flush events was not included as metadata in this study. Mature fruits were harvested between our April FF and May FS sampling events. Fertilizer treatments and/or amendments and number of hours irrigated were collected as monthly metadata variables.

Leaf and root sampling. Leaves sampled were fully mature, with a fully hardened cuticle, and dark green (no yellowing or signs of senescence). Leaves were from one of the previous year’s flush events and, thus, in a developmental cohort separate from that of the reproductive structures on the trees at the time of sampling. During sampling, each tree was divided into 4 quadrants (north, south, east, and west), and two stems with at least two leaves of the target age attached (petiole attached) were collected from each quadrant with sterilized hand clippers and pooled into a sterile 24-oz. standup whirl-pak bag (Nasco B01401, Fort Atkinson, WI). Both fruit-bearing and nonbearing branches were collected at random, and all stems/leaves from a single tree were pooled and immediately placed on ice in a cooler for transit to the laboratory.

Fibrous or feeder roots were sampled from two sides of the tree approximately 0.5 m away from the base of the trunk near the irrigation line. A small hand shovel was used to remove the topsoil until roots were visible. Approximately 5 g of fibrous roots were gently pulled from the ground and shaken until all loose soil was removed. Then, roots were sealed in a sterile 24-oz. standup whirl-pak bag (Nasco B01401, Fort Atkinson, WI) and immediately placed on ice in a cooler for transit to the laboratory.

Gloves were changed and clippers and hand shovels were sterilized with 30% household bleach between trees. All samples were transferred to the LREC onsite laboratory where they were immediately frozen at -20°C . Samples were inspected by the California Department of Food and Agriculture according to California citrus quarantine protocols prior to overnight shipment to UC Riverside on dry ice.

Tissue processing and DNA extractions. At UC Riverside, samples were kept frozen on dry ice during processing for downstream DNA extractions. For leaf samples, 1 to 2 leaves were removed from each of the eight stems in the pooled sample bag (total of 8 to 10 leaves per tree sample). Leaf tissue was chopped into 1- to 2-cm pieces and put in 50-mL conical tubes (Falcon 352098, Corning, Glendale, AZ). Roots were rinsed with sterile water to remove surface soil and placed in a 50-mL conical tube (Falcon 352098, Corning, Glendale, AZ). All leaf and root samples were stored at -80°C and then lyophilized with a benchtop freeze dryer (Labconco FreeZone 4.5L, Kansas City, MO) for 16 to 20 h. Dried samples were crushed into <0.5 -cm fragments and homogenized in the 50-mL conical tubes using a flame-sterilized metal spatula. Leaf and root DNA extractions were performed according to published protocols (22). Briefly, 100 mg of freeze-dried crushed tissue was transferred to 2-mL microcentrifuge tubes (Eppendorf Safe-Lock tubes; Eppendorf, Hamburg, Germany) containing a 4-mm stainless-steel grinding ball (SPEX SamplePrep, Metuchen, NJ, USA). Samples were chilled at -80°C for 15 min and then pulverized into a powder using a 2010 Geno/Grinder (SPEX SamplePrep) at 1,680 rpm for 20 s ($\times 2$). Then, 1 mL of 4 M guanidine thiocyanate was added to the pulverized leaf and root samples. All samples were incubated at 4°C for 15 min. Next, samples were centrifuged for 1 h at $17,500 \times g$. DNA was isolated from the supernatant using the MagMAX-96 DNA multi-sample kit (Thermo Fisher Scientific) on a MagMAX Express-96 deep well magnetic particle processor. DNA was eluted in 100 mL of DNA elution buffer and stored at -20°C until utilized for bacterial and fungal Illumina library construction.

High-throughput sequencing library preparation. Leaf and root bacterial communities were sequenced from all leaf ($n = 159$) and root ($n = 159$) samples (20 time points). Bacterial Illumina Miseq libraries were built by amplifying the bacterial rRNA 16S V4 region using the 515FB/806RB primer set and the standardized Earth Microbiome Project protocol, which can be found online (<https://earthmicrobiome.org/protocols-and-standards/16s/>) (71). To limit the amplification of plant mitochondrial and plastid 16S regions, we used pPNA and mPNA clamps (PNA Bio, Newbury Park, CA), which bind to these plant sequences and block binding of the bacterial 515FB/806RB (72, 73). Leaf sample reactions received $0.75 \mu\text{M}$ pPNA and mPNA clamps, and $0.75 \mu\text{M}$ mPNA was added to each root sample reaction.

The first 14 consecutive monthly leaf and root samples were included in fungal community libraries. Our preliminary analyses shifted our main focus to bacterial communities, and therefore we did not sequence the fungal communities of leaf/roots collected at the last 6 time points in the second sampling year. Fungal Illumina MiSeq libraries were built by amplifying the fungal ITS1 region using the ITS1f/ITS2 primer set and the standardized Earth Microbiome Project protocol.

Triplicate PCRs for each sample were pooled, and amplification was verified on a 1% agarose gel. Amplicon samples were quantified for DNA concentration using Quant-iT PicoGreen (Invitrogen). Equal amounts of amplicons (240 ng) from each sample were pooled and AMPure XP (Beckman Coulter) beads were used to clean the sample library. The cleaned library was then quantified using Qubit (Qiagen; 260/280). The libraries were diluted to 20 to 30 $\mu\text{g}/\text{mL}$, and a final quality assessment was done with a Bioanalyzer at the UCR Genomics Core facility. Paired-end sequencing (2×300) was performed on an Illumina MiSeq platform with a 20% PhiX spike included before sequencing.

Data processing and statistics. Demultiplexed, PhiX reads-removed, and Illumina adapter-trimmed sequences were received from the UCR Genomics Core. Amplicon sequencing raw reads for 16S rRNA genes and fungal ITS2 region are available on the NCBI SRA database under BioProject accession number [PRJNA685913](https://www.ncbi.nlm.nih.gov/bioproject/PRJNA685913). Bacterial and fungal reads were preprocessed using a USEARCH (v9.1.13)/VSEARCH (v2.3.2) pipeline. Forward and reverse sequencing files were joined with USEARCH allowing for staggered ends and up to 10 mismatches. After quality filtering using VSEARCH, there were 65.5 million 16S reads and 21.1 million fungal reads. Sequences were dereplicated, singletons were removed, and OTUs were formed using USEARCH with a 97% similarity cutoff. The bacterial library produced 19,626 OTUs, which were assigned to 4,183 taxonomic names using the RDP 16S database, v18. The fungal library produced 44,447 OTUs, which were assigned to 31,056 taxonomic names using the UNITE fungal reference database, v02.02.2019. On average, 12.7% of the 16S library reads from each leaf and root sample were assigned as bacteria. See Table S1 in the supplemental material for all 16S bacterial read counts of individual samples. The remaining sequences, which were removed, were attributed to chloroplast, plant mitochondria, or *Archaea* or could not be assigned to a kingdom. Our fungal libraries did not have non-specific binding issues. USEARCH was also used to create phylogenetic tree files in Newick format.

Preprocessed taxonomically assigned OTU tables were imported into R (v3.6.0). Samples with fewer than 1,000 reads were removed. Reads were rarified to even depth for each alpha diversity comparison. Alpha diversity was compared using the number of OTUs observed (74). A ranked sums analysis of variance statistical test, Kruskal-Wallis, followed by a pairwise Dunn's test with Holm's correction for multiple comparisons, was used to calculate *P* values.

Beta diversity analyses were performed using R packages phyloseq (v1.28.0) and vegan (v2.5.6) (74, 75). Preprocessed reads were transformed using total sum scaling normalization. Using the `ordinate()` and `plot_ordination()` functions, a principal coordinate analysis (PCoA) was done on weighted Unifrac distances, which accounts for relative relatedness and quantitative variance of communities. Ninety-five percent confidence ellipses were added to further examine groups using the `stat_ellipse()` function. A permutational multivariate analysis of variance (PERMANOVA) statistical test was performed on weighted Unifrac distances that were square-root transformed using the `vegan::adonis2()` function, including the following covariates and interaction terms in the model: `adonis2(dist.matrix ~ phenology*Sample_year + Fertilizer + Mean_Temp*hrs_irrigated + Total_rain, permutations = 999, method = "wunifrac," sqrt.dist = TRUE)`. Pairwise PERMANOVA with false-discovery rate (FDR) correction was accomplished with `RVAideMemoire::pairwise.perm.manova()` (v0.9.74) (76). Core microbiota identifications were performed using `microbiome::core()` (v1.6.0) with prevalence set at 0.75 and detection set at 0.01/100 (77). The core root bacteriome was also defined with prevalence set at 0.75 and detection set at 0.1/100, which did not affect the interpretation of the results but greatly improved readability of the phylogenetic tree.

To understand if phylogenetic relationships affect microbial associations with phenological stages, core genera taxonomic assignments were input into a phylogenetic tree generator, phyloT v2 (<https://phyloT.biobyte.de/index.cgi>), to generate a Newick format phylogenetic tree. Tree and metadata were visualized using the interactive tree of life visualization program, iTOL (<https://itol.embl.de/>) (78). Genera with fewer than 50 reads were filtered out, and differentially abundant populations at the genus level were identified using DESeq2 (v1.24.0) to run a parametric fit for dispersion on a negative binomial generalized linear model, followed by a Wald test with FDR adjustment to produce *P* values (79). Input root and leaf bacterial OTU tables used in DESeq2 analysis were rescaled using a pseudocount of 1. All possible phenological stage pairwise combinations were tested using `results()` and the `contrast` option. Using `ggplot2` (v3.2.1) and `phyloseq::subset_taxa()` function, the relative abundance of specific species were plotted as boxplots (74, 80).

The top 300 most abundant leaf bacterial OTUs were input into a network analysis using Sparse Inverse Covariance Estimation for Ecological Association Inference, Spiec-Easi (v1.0.7), which infers interactions using neighborhood selection and the concept of conditional independence, rather than a standard correlation or covariance estimation (81). With `set.seed(1244)`, neighborhood modeling (mb) was executed with the `nlambda` set to 70 and `rep.num` set at 99; standard settings were used for all other parameters. Spiec-Easi results were converted to `igraph` format and imported into Gephi (v0.9.2). The network was visualized using a Yifan Hu layout, and taxa were filtered to focus on differentially abundant taxa and immediate neighbors. Betweenness centralities were calculated using Gephi network diameter statistics, and centralities were normalized to a 0 to 1 scale. Taxa were further filtered by abundance, with the final figure showcasing the 155 most abundant OTUs within the above parameters. This significantly increased readability without affecting the interpretation of the results.

Data availability. The data that support the findings of this study are openly available in the NCBI SRA database under BioProject [PRJNA685913](https://www.ncbi.nlm.nih.gov/bioproject/PRJNA685913).

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

FIG S1, PDF file, 1.7 MB.

FIG S2, TIF file, 1.6 MB.

FIG S3, PDF file, 0.9 MB.

FIG S4, PDF file, 2.3 MB.

FIG S5, TIF file, 1.3 MB.

TABLE S1, XLSX file, 0.02 MB.

TABLE S2, DOCX file, 0.01 MB.

TABLE S3, XLSX file, 0.04 MB.

TABLE S4, XLSX file, 0.2 MB.

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We declare that we have no competing interests.

N.A.G. and M.C.R. designed the project. N.A.G. and P.R. conducted fieldwork. N.A.G. and I.D.A. processed field samples. N.A.G. and F.C.F.V. built sequencing libraries. N.A.G. performed data analyses. N.A.G. and M.C.R. wrote the manuscript. All authors contributed to the editing of the manuscript.

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