Genotypic variation in floral volatiles influences floral microbiome more strongly than interactions with herbivores and mycorrhizae in strawberries

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Abstract

The floral microbiome is of significant relevance to plant reproduction and crop productivity. While plant genotype is key to floral microbiome assembly, whether and how genotypic variation in floral traits and plant-level mutualistic and antagonistic interactions at the rhizosphere and phyllosphere influence the microbiome in the anthosphere remain little known. Using a factorial field experiment that manipulated biotic interactions belowground (mycorrhizae treatments) and aboveground (herbivory treatments) in three strawberry genotypes, we assessed how genotypic variation in flower abundance and size and plant-level biotic interactions influence the bidirectional relationships between floral volatile organic compounds (VOCs) and the floral microbiome using structural equation modeling. We found that plant genotype played a stronger role, overall, in shaping the floral microbiome than biotic interactions with mycorrhizae and herbivores. Genotypic variation in flower abundance and size influenced the emission of floral VOCs, especially terpenes (e.g. α- and β-pinene, ocimene isomers) and benzenoids (e.g. p-anisaldehyde, benzaldehyde), which in turn affected floral bacterial and fungal communities. While the effects of biotic interactions on floral traits including VOCs were weak, mycorrhizae treatments (mycorrhizae and herbivory + mycorrhizae) affected the fungal community composition in flowers. These findings improve our understanding of the mechanisms by which plant genotype influences floral microbiome assembly and provide the first evidence that biotic interactions in the rhizosphere and phyllosphere can influence the floral microbiome, and offer important insights into agricultural microbiomes.

Introduction

The floral microbiome that consists of thousands or more taxa of bacteria and fungi mediates processes that are essential for plant reproduction and crop productivity [1–5]. While we are just beginning to understand the principles and functions of microbiome assembly in flowers [2, 3], studies have revealed that the floral microbiome is governed by plant genotype [1], and it overlaps with the leaf microbiome harboring a subset of the microbiota hosted by the root microbiome following a source–sink gradient [5]. Thus, plant genotypic variation in floral traits and species interactions that dictate the microbiomes in the rhizosphere (roots) and phyllosphere (leaves) are expected to influence the microbiome in the anthosphere (flowers).

Floral traits can influence the floral microbiome via distinct mechanisms. For instance, flower size is hypothesized to affect the niche availability for microbiota [6]. Flower abundance, on the other hand, can influence floral microbiome assembly by governing the source pool of colonizing microbes [1, 7] and by enhancing pollinator-mediated microbial dispersal [1]. In particular, floral volatile organic compounds (VOCs) have been shown to mediate pollinator foraging [8], which can potentially influence microbial dispersal [3]. In addition to the pollinator-mediated effect of floral VOCs on the floral microbiome, VOCs are known for their antimicrobial properties (e.g. terpenes, benzenoids, aliphatics) [9–12] and their roles as carbon sources (e.g. methanol) [9], and thus can potentially reduce pathogenic microbes [10, 13], modulate microbe–microbe interactions [14] and thereby affect microbial communities [12]. Microbiota, in turn, can influence plant VOCs by directly contributing to VOC emissions [3] or indirectly by metabolizing plant VOCs and/or affecting plant physiology that can influence plant VOC emissions [9, 10, 13]. Thus, the effect of microbiota on floral VOCs is determined, in part, by the quantity and quality of plant metabolites [9] in the anthosphere. Such bidirectional relationships between floral VOCs and floral microbiome are likely to be influenced...
samples were collected from 93 plants flowering on August 16, 2019, which represented a subset of 144 total plants in the experiment [18].

Supplementary Table 1). Bacterial Shannon diversity of floral bacterial and fungal communities (Fig. 2 and floral traits including VOCs [13, 15], which can influence herbivores) [16, 17] may affect the source–sink dynamics (e.g. mycorrhizae) [15] and phyllosphere antagonists (e.g. instance, plant interactions with rhizosphere mutualists by biotic interactions experienced by the whole plant. For example, plant interactions with rhizosphere mutualists (e.g. mycorrhizae) differ among genotypes depending upon herbivory treatment on the floral microbiome. Stronger effects of plant genotype than herbivory (e.g. mycorrhizae) were grouped into and randomized within blocks (N = 1–12). The three strawberry genotypes (Seascape, Tribute, and Wasatch) were assigned to and completely randomized within each treatment. Floral microbiome samples were collected from 93 plants flowering on August 16, 2019, which represented a subset of 144 total plants in the experiment [18].

Here we leveraged a factorial field experiment [18] that manipulated biotic interactions belowground (mycorrhizae treatments) and aboveground (herbivory treatments) in three day-neutral strawberry cultivars (Fragaria ×ananassa “Seascape”, “Tribute” and “Wasatch”; Fig. 1). As clonal plants of each cultivar were used in this study, we referred to cultivars as genotypes. We first examined how epiphytic bacterial and fungal communities in flowers responded to plant genotype, mycorrhizae treatments and herbivory treatments. We then assessed how plant genotypes varied in floral traits (flower abundance, size and VOCs) that may shape the floral microbiome. Lastly, we used structural equation modeling to link genotypic variation in floral abundance and size and biotic treatments to the bidirectional relationships between floral VOCs and floral microbiome (bacterial and fungal α- and β-diversity).

**Results**

**Stronger effects of plant genotype than treatment on the floral microbiome**

Strawberry genotypes influenced the α- and β-diversity of floral bacterial and fungal communities (Fig. 2 and Supplementary Table 1). Bacterial Shannon diversity differed among genotypes depending upon herbivory treatments, as revealed by a general linear mixed model (LMM, genotype–herbivory interaction, $F = 7.06$, $df = 2$, $P = 0.002$). For instance, bacterial Shannon diversity was higher in Tribute than Seascape in herbivory-absent treatments (least-squares mean, LS-mean, contrast: control, $\chi^2 = 6.16$, $df = 1$, $P = 0.013$; mycorrhizae, $\chi^2 = 5.46$, $P = 0.019$; Fig. 2a), but was similar in herbivory-present treatments (herbivory, $\chi^2 = 2.47$, $P = 0.12$; herbivory + mycorrhizae, $\chi^2 = 0.90$, $P = 0.34$). In addition, Tribute had lower bacterial Shannon diversity than Wasatch but only in herbivory + mycorrhizae treatment ($\chi^2 = 4.74$, $df = 1$, $P = 0.030$; Fig. 2a). Within genotypes, bacterial Shannon diversity was influenced by treatments, especially for Tribute (control > herbivory, $\chi^2 = 7.22$, $df = 1$, $P = 0.007$; control > herbivory + mycorrhizae, $\chi^2 = 8.93$, $P = 0.003$) and Seascape (herbivory > mycorrhizae, $\chi^2 = 5.45$, $P = 0.020$), but not for Wasatch (Fig. 2a).

Similar to α-diversity, the β-diversity of floral bacterial communities was influenced by plant genotype (Fig. 2c and Supplementary Table 1). The permutational multivariate analysis of variance (PERMANOVA) of Bray–Curtis dissimilarity identified genotype as the primary source of variation in bacterial community composition (4% of variation, $F = 1.70$, $df = 2$, $P = 0.035$; Supplementary Table 1), which was driven by bacterial community differences between Tribute and Wasatch (pairwise PERMANOVA, $F = 2.53$, $df = 1$, $P = 0.016$) but not other pairs (Seascape vs. Tribute, $F = 1.08$, $P = 0.32$; Seascape vs. Wasatch, $F = 1.32$, $P = 0.22$). Supporting the PERMANOVA results, the constrained principal coordinates analysis (cPCoA) of bacterial communities revealed significant separation between Tribute and Wasatch (pairwise cPCoA, $F = 2.48$, $df = 1$, $P = 0.016$; Fig. 2c), with Seascape being intermediate between them but not different from either (Seascape vs. Tribute, $F = 1.07$, $P = 0.33$; Seascape vs. Wasatch, $F = 1.31$, $P = 0.23$).

For floral fungal communities, although Shannon diversity was similar among genotypes (LMM, $F = 0.34$, $df = 2$, $P = 0.71$; Fig. 2b), community composition varied...
significant with genotype (PERMANOVA, 3% of variation, \(F = 1.32, df = 2, P = 0.045\); cPCoA, 3%, \(F = 1.31, P = 0.050\); Fig. 2d and Supplementary Table 1).

### Plant genotypes differ in flower abundance, size and VOCs

Given the significant effect of genotype on the floral microbiome, we then examined genotypic variation in aspects of floral phenotype (e.g. flower abundance, size and VOCs) potentially important in governing floral microbiome assembly. The LMM revealed that flower abundance differed among genotypes (\(F = 6.60, df = 2, P = 0.004\)) but not among treatments (herbivory treatments, \(F = 1.36, df = 1, P = 0.25\); mycorrhizae treatments, \(F = 0.11, P = 0.74\); Supplementary Table 2). In particular, Wasatch produced more flowers than Tribute (LS-mean contrast, \(\chi^2 = 12.1, df = 1, P = 0.001\); Fig. 3a), with Seascape being intermediate (Wasatch vs. Seascape, \(\chi^2 = 3.3, P = 0.068\); Tribute vs. Seascape, \(\chi^2 = 2.5, P = 0.12\)). Similar to flower abundance, flower size (estimated by flower diameter) also differed among genotypes (LMM, \(F = 3.34, df = 2, P = 0.041\); Supplementary Table 2), with Seascape having the largest and Wasatch the smallest (Fig. 3b). Yet, there was no significant trade-off between flower abundance and flower size (Pearson’s correlation, \(t = -0.78, df = 78, P = 0.44\)) across genotypes.

The floral VOC profile consisted of 22 identified VOCs: 10 terpenes (\(\alpha\)-pinene, \(\beta\)-pinene, 6-methyl-5-hepten-2-one, \(\delta\)-limonene, ocimene isomers, geranyl acetate, dihydro-\(\beta\)-ionone, \(\beta\)-ionone, \(\alpha\)-murolene and \(\alpha\)-farnesene), seven benzenoids (anisole, benzaldehyde, benzyl alcohol, methyl salicylate, p-anisaldehyde, 3-hexen-1-ol benzoate and benzyl benzoate), three aliphatics (trans-2-hexen-1-al, (E)-3-hexen-1-ol and cis-3-hexenyl acetate), one (sulfur) S-containing compound (benzothiazole) and one C5 branched-chain compound (hexenyl tiglate). Plant genotype explained the largest source of variation in VOCs (PERMANOVA, 12%, \(F = 4.47, df = 2, P = 0.001\); cPCoA, 11%, \(F = 3.81, P = 0.001\); Fig. 3c and Supplementary Table 3), with the first two axes of VOC cPCoA (cPCoA1 and cPCoA2; Fig. 3c) driven primarily by variation in benzaldehyde and cis-3-hexenyl acetate, respectively (Supplementary Table 3). Pairwise PERMANOVAs and cPCoAs further revealed that all three genotypes produced significantly different VOC profiles from each other (all \(P < 0.05\)). The random forest classification identified the specific VOCs that differed between genotypes, that is, \(\alpha\)-pinene, \(\beta\)-pinene, geranyl acetate and benzaldehyde between Seascape and Tribute (out-of-bag error, OOB = 6.12%), \(\alpha\)-pinene and benzaldehyde between Seascape and Wasatch (OOB = 17.5%), and \(\alpha\)-pinene, ocimene isomers and benzyl alcohol between Tribute and Wasatch (OOB = 9.3%).

### SEMs linking floral phenotype and treatment to the floral microbiome

Given the significant genotypic differences in both floral phenotype and microbiome, we then used SEMs to link genotypic variation in flower abundance, size and VOCs, as well as treatments, to the \(\alpha\) - and \(\beta\) -diversity of floral bacterial and fungal communities (Figs. 4a and 5a). The SEMs focused on VOCs at three different levels: (1) individual VOCs that differed between genotypes as identified by the random forest classification (see above), and that predicted the \(\alpha\)- or \(\beta\)-diversity of bacterial and fungal communities based on backward predictor selection (see Materials and methods; Supplementary Tables 4 and 5); (2) the major classes of VOCs (terpenes, aliphatics, benzenoids, S-containing compounds and C5 branched-chain compounds); and (3) VOC profile (VOC cPCoA1 and cPCoA2; Fig. 3c). The SEMs were examined in parallel
considering overall genotypic (within- and between-genotype) variation using lavaan [19] (Figs. 4 and 5), and within-genotype variation alone using piecewiseSEM [20] (with genotype as a random effect; Supplementary Figs. 1 and 2). This approach was used to confirm whether the detected relationships based on overall genotypic variation were also supported within genotypes.

The SEMs based on overall genotypic variation revealed that α-pinene and β-pinene were negatively linked with bacterial Shannon diversity ($r = -0.21$, $P = 0.050$ and $r = -0.26$, $P = 0.017$, respectively; Fig. 4b, c and Supplementary Table 4). Bacterial ASVs associated with the emission of α-pinene and β-pinene (Supplementary Table 6) were common residents of the floral microbiome in cultivated strawberry [1] and wild strawberries [5], including, for instance, Hymenobacter, Massilia, Deinococcus and nectar taxa (Acinetobacter, Arthrobacter, Methylobacterium, Nocardioides, Sphingomonas and Streptomyces) [21]. While treatments and flower abundance showed no notable effects ($P > 0.10$), flower size influenced bacterial Shannon diversity indirectly via enhancing the emission of α-pinene ($r = 0.41$, $P < 0.001$; Fig. 4b) and β-pinene ($r = 0.41$, $P < 0.001$; Fig. 4c). Different from α-pinene and β-pinene, other VOCs, major VOC classes and VOC profile (VOC cPCoA1 and cPCoA2) were not linked with bacterial Shannon diversity (Supplementary Table 4). For fungal Shannon diversity, only S-containing compound (benzothiazole) showed a positive relationship ($r = 0.31$, $P = 0.039$; Fig. 5b). Eight fungal ASVs that were commonly found in strawberry flowers [1] were identified as associated with the emission of the S-containing compound (Kendall’s correlation coefficient, $P < 0.05$; Supplementary Table 6), including those positively associated with the S-containing compound such as Cystofilobasidium macerans (yeast found in both flowers [1] and pollinators [22]), Phaeosphaeria caricola (strains of this genus often pathogenic [23]) and Ramularia pratensis (some strains with biocontrol activities [24]), as well as those negatively associated with the S-containing compound such as pathogenic Blumeria graminis [25]. These detected relationships based on overall genotypic variation were also supported by the SEMs that examined within-genotype variation (Supplementary Fig. 1b, c; Supplementary Fig. 2b).

The β-diversity of floral bacterial communities was linked with a different set of VOCs relative to the α-diversity (Fig. 4), as revealed by the SEMs based on overall genotypic variation. Specifically, p-anisaldehyde was positively linked with the first axis of bacterial community composition (cPCoA1, $r = 0.24$, $P = 0.019$; Fig. 4d), which was dominated by Lactobacillus micheneri (found in both flowers [1] and pollinators [1, 26]) and Methylobacterium and Acinetobacter nectaris (nectar taxa [21]) on the positive axis and Delftia (common taxa in strawberry flowers [1]) on the negative axis (Fig. 2c; Supplementary Table 1). Moreover, ocimene isomers ($r = -0.32, P = 0.029; Fig. 4e$), aliphatics ($r = -0.18, P = 0.069; Fig. 4f) and the C5 branched-chain compound (hexenyl tiglate, $r = -0.28, P = 0.023; Fig. 4g$) were negatively linked with the cPCoA2 of bacterial communities, which was dominated by Delftia, Lactobacillus micheneri, Methylobacterium and Acinetobacter nectaris on the positive axis and Phaseolibacter (common taxa in strawberry flowers [1]) and Sphingomonas and Rosenerbergiella (nectar taxa [21]) on the negative axis (Fig. 2c; Supplementary Table 1). The cPCoA2 of bacterial communities was also influenced by flower size indirectly via the enhanced emission of VOCs (ocimene isomers, $r = 0.28, P = 0.021$, Fig. 4e; aliphatics, $r = 0.29, P = 0.008$, Fig. 4f), but was not influenced by flower abundance or treatments (Fig. 4). These relationships based on overall genotypic variation were also detected within genotypes (Supplementary Fig. 1).

The β-diversity of fungal communities, on the other hand, were influenced by treatments and flower abundance (Fig. 5c–f), as revealed by the SEMs based on overall genotypic variation. Relative to the control, mycorrhizae treatment positively affected the cPCoA1 of fungal communities (all $P < 0.10$; Fig. 5c, e, f and Supplementary Table 5), which was dominated by Pseudophytomyces chartarum (potentially pathogenic [25]) and Candelaria concolor (lichenized fungal taxon [27]) on the positive axis and pathogenic Alternaria alternata [28] and Cladosporium delicatulum [25] on the negative axis (Fig. 2d; Supplementary Table 1). In addition, herbivory
Figure 4. Structural equation models (SEMs) linking floral phenotype and treatment to floral bacterial microbiome. a, Hypothesized effects of flower abundance and size as well as treatments on floral volatile organic compounds (VOCs) and bacterial microbiome. The floral bacterial microbiome was characterized using $\alpha$-diversity (Shannon diversity) and $\beta$-diversity (the first two axes of constrained principal coordinates analysis, cPCoA1 and cPCoA2). VOCs were evaluated at three levels (see Materials and methods): individual VOCs, major classes of VOCs, and VOC profile (VOC cPCoA1 and cPCoA2). The bidirectional links between VOCs and bacterial microbiome reflected potential reciprocal influence on each other. Treatments were coded using the control treatment as the reference level. SEMs were fitted based on overall genotypic variation using lavaan [19]. b-g, SEMs that contained notable positive (solid black) and negative (dashed black) links between VOCs and bacterial microbiome, as well as other variables, were presented (see SEM details in Supplementary Table 4): †$P < 0.10$; ∗$P < 0.05$; ∗∗$P < 0.01$; ∗∗∗$P < 0.001$. Grey links indicate $P > 0.10$. Link width and the numbers adjacent to links indicate standardized path coefficients.

Discussion

Our results demonstrated that strawberry genotype played a stronger role, overall, in shaping the floral microbiome than whole-plant biotic interactions both belowground (with mycorrhizae) and aboveground (with herbivores). In particular, we found that genotypic variation in flower abundance and size influenced floral VOCs, especially terpenes and benzenoids, which in turn
affected the floral bacterial and fungal communities. While the effects of biotic interactions on floral phenotype including VOCs were weak (Figs. 3–5), mycorrhizae treatments (mycorrhizae and herbivory + mycorrhizae) directly affected the fungal communities in flowers relative to the control (Fig. 5). By linking genotypic variation in floral traits and biotic interactions to floral bacterial and fungal communities, our study revealed the mechanisms by which plant genotype influences floral microbiome assembly [1] and also provided the first evidence that plant-level biotic interactions at the rhizosphere and phyllosphere can influence the anthosphere microbiome.

In line with previous studies on the floral microbiome [1], genotypic variation in flower abundance influenced the floral microbiome. Flower abundance has been shown to govern floral microbiome assembly both directly by influencing the source pool of microbes that colonize individual flowers [1, 7] and indirectly by influencing pollinator visits [30] that can mediate microbial dispersal [1]. Our results showed that flower abundance affected fungal α- and β-diversity via the bidirectional links between VOCs and the fungal communities among genotypes but not within genotypes. In addition to flower abundance, flower size that is hypothesized to influence the floral microbiome has, nevertheless, rarely been tested [6]. In this study, we found that flower size affected bacterial and fungal communities via the bidirectional links between VOCs and the floral microbiome, which was supported by SEMs based on overall genotypic variation and within-genotype variation as well. While the causation of these relationships awaits experimental validation, it provides a new insight into the mechanisms by which flower abundance and size drive microbiome assembly.

Our results provided evidence for the hypothesized bidirectional relationships [9] between plant VOCs and microbiome in the anthosphere. Specifically,
monoterpenes, namely \( \alpha \) - and \( \beta \)-pinene and ocimene isomers contributed most to the variation in floral VOC profile and floral bacterial communities. Bacterial Shannon diversity was negatively associated with \( \alpha \) - and \( \beta \)-pinene, whereas bacterial community composition was associated with p-anisaldehyde, ocimene isomers, aliphatics and hexenyl tiglate (the only C5 branched-chain compound in this study). These bidirectional relationships were supported by overall genotypic variation and within-genotype variation. On the other hand, fungal Shannon diversity was positively linked with benzothiazole, the only S-containing compound detected in strawberry volatile collections here, whereas fungal community composition was associated with terpenes (especially ocimene isomers) and benzenoids (especially benzaldehyde). Yet, the bidirectional relationship between benzaldehyde and fungal community composition was only supported among genotypes. To further verify that the relationship is not caused by unmeasured genotypic differences, experimental investigation that directly tests the reciprocal effects is needed. Overall, despite the known antimicrobial effects of floral VOCs such as terpenes, benzenoids, aliphatics and S-containing compounds [9–12], our results detected microbial taxa both negatively and those positively associated with the emission of these VOCs. This finding suggests the potential dual roles of floral VOCs as antimicrobial agents and carbon sources for some microbial taxa [9, 10] and/or their role in mediating complex microbe–microbe interactions [14] where the suppression of some taxa can potentially increase other taxa via direct and indirect interactions through the microbial network [1]. While how dynamic the relationships are between floral VOCs and the floral microbiome remains an open question, our results are expected to hold despite the time interval between VOC and microbiome collections, due to stable VOC composition (i.e. relative abundances of individual VOCs) among these genotypes detected over a longer time period (4 weeks) in this field experiment [18]. Taken together, our findings highlight the needs for future research on unraveling the specific mechanisms that underlie the relationships between floral VOCs and the floral microbiome as well as their temporal dynamics.

While plant interactions with herbivores at the phyllosphere have been found to influence the leaf microbiome [16, 17], our results showed, for the first time, that plant interactions at the rhizosphere and phyllosphere can influence the floral microbiome. Such effects were detected on shaping the floral fungal communities. Specifically, the mycorrhizae treatment caused the deviation of fungal community composition from the control (Fig.5) by reducing pathogenic taxa (e.g. A. alternata, Cladosporium delicatum) in flowers, whereas the herbivory + mycorrhizae treatment influenced fungal community composition by reducing floral yeasts (e.g. R. graminis, Naganishia randhawae) and increasing pathogenic taxa (e.g. A. alternata). While the effect of mycorrhizae treatment occurred among genotypes, the effect of herbivory + mycorrhizae treatment was supported by SEMs based on overall genotypic variation and within-genotype variation as well. These effects of biotic interactions on the floral microbiome were, nevertheless, not via floral VOCs, although plant mutualists and antagonists have been shown previously to alter floral VOCs [15, 31, 32]. In this study, the mycorrhizae treatments and herbivory treatments exhibited weak effects on the VOC profile. Yet, we could not rule out the possibility that the effects of the biotic treatments on floral VOCs can be dynamic and thus attenuate over time, and plant genotypes and their floral VOCs might respond differently to natural mycorrhizal inocula compared to a single mycorrhizal inoculum used in this study. Whether the effects of mycorrhizae and herbivory + mycorrhizae treatments on floral fungal microbiome are caused by altered source pools of microbes from the rhizosphere and phyllosphere that can affect the source–sink dynamics or altered plant traits not measured here that can affect floral fungal taxa merits further investigation.

To conclude, plant genotype governs floral microbiome assembly via both direct and indirect mechanisms mediated by diverse floral traits, especially the bidirectional relationships between floral VOCs and bacterial and fungal communities. In addition to the intrinsic characteristics of plant genotypes, interactions with non-focal mutualists and antagonists at the rhizosphere and phyllosphere influence the anthosphere microbiome that can profoundly affect plant reproduction [1, 2, 4]. These findings should not be unique to strawberries but may be generalizable to other crops and wild plants, because the VOCs of important relevance to the floral microbiome are common among plant species [9–13], and mycorrhizal partners and herbivore pests are ubiquitous in natural and agricultural ecosystems [33, 34]. We emphasize that our findings are based on plants with pre-existing endophytic microbiomes; as such, the extent to which endophytic microbiomes influence the detected relationships remain to be determined. Nevertheless, our findings based on strawberries improve the understanding of microbiome assembly in flowers and offer important insights into agricultural microbiomes that can influence plant reproduction and crop productivity.

Materials and methods
Factorial field experiment
The factorial field experiment [18] was conducted during March–September, 2019, at the Oakland University Student Farm, Rochester, Michigan (42.66073°N, 83.19558°W). Three strawberry genotypes (“Seascape”, “Tribute” and “Wasatch”) with 48 bare-root nursery stock plants per genotype were included in the experiment. The fresh biomass was recorded for each bare-root plant (i.e. initial plant weight) to account for potential
influence of variation in initial plant size on the experiment. These bare-root plants (N = 144 total) were grown in 1-gallon pots filled with 2:1:1 mixture of sphagnum peat moss (Lambert, Rivière-Ouelle, Québec, Canada), washed play sand (Kolorscape, Atlanta, GA) and Turface MVP (Turface, Buffalo Grove, IL) during March 10–16, 2019, and re-potted into 2-gallon white fabric pots (247Garden, Montebello, CA) filled with the same media during May 24–29, 2019. These plants were subjected to four treatments (control, herbivory, mycorrhizae, and herbivory + mycorrhizae). Within each treatment, a single plant from each genotype was randomly selected (Fig. 1). The four treatments were then grouped into and replicated across 12 blocks (Fig. 1).

For mycorrhizae-present treatments (mycorrhizae and herbivory + mycorrhizae), plants were inoculated with *Rhizophagus irregularis* spores (Elite 91 Myco Jordan, Murietta, CA) upon initial potting by dusting roots with inocula and also filling the 1-gallon pots half full of media and adding a thin layer of 5 mL inocula at the point of contact with roots. For herbivory-present treatments (herbivory and herbivory + mycorrhizae), plants experienced both natural herbivory from *Vanessa cardui* caterpillars (Carolina Biological Supply, Charlotte, NC; with three to five 2nd and 3rd instar larvae feeding on a single leaf per plant for six days) and simulated herbivory (by clipping the distal half each leaf from half of the leaves per plant and then applying 1 mM jasmonic acid solution in water [35] to cover all upper leaf surface) during June 17–July 6, 2019. The combination of natural and simulated herbivory is commonly used [36] to ensure that plants received the stimulus of a live herbivore and associated cues (e.g. enzymes in caterpillar saliva) while standardizing for the leaf area damaged per unit time.

### Floral traits and VOC collection

In this study, we recorded the number of open flowers produced over a two-week period (August 4–17, 2019) prior to and during the floral microbiome collection. We sampled floral volatile emissions eight weeks after herbivory treatments, and 5–15 days prior to floral microbiome collection (July 30–August 10, 2019) to minimize the potential influence of the activities of VOC measurements that can cause artificial microbial arrival and destructive flower collection on floral microbiome. Floral volatiles were collected from new fully expanded flowers between 11 AM and 3 PM on sunny to partly sunny days, with no to little chance of rain. During the 4-h sampling period each day, 16 floral and two blank samples were collected using a dynamic headspace sampling method [31], by enclosing flowers in 12 oz. polyethylene terephthalate cups with dome lids (Comfy Package, Rikkel Corp, NY) and pulling air through a semi-open system at a flow rate of 200 mL/min through volatile traps with 30 mg of HayeSep Q adsorbent (VAS, Rensselaer, NY). We used either a PVAS22 portable volatile assay system (VAS, Rensselaer, NY) or an Air Lite low-flow air sample pump (SKC, Eighty Four, PA), which were shown functionally equivalent in volatile collection during initial trials. Volatiles were analyzed on an Agilent 7890A gas chromatograph (GC) with an Agilent 5977B mass spectrometer (MS) and separated on a HP-5 column (30 m × 250 μm × 0.25 μm) with helium as a carrier gas. We measured flower diameter and dry weight for all flowers sampled. Floral VOC emissions (per hour) were standardized by flower dry weight.

### Floral microbiome collection and sequencing

The floral microbiome was collected from individual strawberry plants that flowered on August 16, 2019 (N = 93 out of 144 plants). A single flower (without pedicel) per plant was collected into a sterile 15 mL centrifuge tube using ethanol-rinsed forceps. These flower samples were transported on dry ice to the University of Pittsburgh and stored at −20°C within 12 h. Following the previous protocol [1], epiphytic microbes were collected by sonicating flowers in 3 mL phosphate-buffered saline at 40 kHz for 10 min and vortexing for 3 min. The microbes were pelleted by centrifuging at 10000 x g for 5 min, and then subject to DNA extraction using Quick-DNA Fecal/Soil Microbe Kits (Zymo Research, Irvine, CA). Samples and one negative control (without flower but in the process of microbial isolation and DNA extraction) were sent to Argonne National Laboratory for bacterial (16S rRNA V5–V6 region, 799F–1115r primer pair) and fungal (ITS1f–ITS2) library preparation [1]. Because the negative control failed in library preparation, the 93 samples were sequenced on a 1/2 lane of Illumina MiSeq (paired-end 250 bp).

### Microbial sequence analysis

Demultiplexed paired-end (PE) reads were used for detecting bacterial and fungal amplicon sequence variants (ASVs) using package DADA2 v1.14.0 [37] in R v3.6.2 [38] and QIIME 2 v2019.4 [39]. Following the previous pipeline [1] for bacterial ASV detection in DADA2, PE reads were trimmed and filtered [truncLen = c(245, 245), trimLeft = c(0, 10), maxN = 0, truncQ = 2], prior to specific variant identification that took into account sequence errors. The PE reads were then end joined (minOverlap = 20, maxMismatch = 4) for ASV detection. After chimera removal, ASVs were assigned with taxonomic identification based on the SILVA reference database (132 release NR 99) implemented in DADA2. For fungal ASV detection in DADA2, the PE reads were first screened to remove potential primer contaminations. Due to the low-quality end (~50 bp), the fungal reads were truncated at 200 bp during quality filtering [truncLen = c(200, 200), maxN = 0, truncQ = 2] to ensure the accuracy in ASV detection. This method provided a conservative estimate of fungal ASVs [1], due to the potential loss of information based on ITS length variation [37]. After chimera removal, fungal ASVs were assigned with taxonomic identification based on the UNITE reference database (v8.0 dynamic release) using QIIME 2.
Bacterial and fungal ASV tables were further filtered separately before conversion into microbial community matrices using package phyloseq [40]. First, we removed non-focal ASVs (Archaea, chloroplasts and mitochondria). Second, we filtered out samples of low reads (<200 reads). Third, we normalized per-sample reads to the same number (i.e. the median reads, 4114 and 10377, bacterial and fungal data set, respectively) [1,5]. Lastly, we removed low-frequency ASVs (<0.001% of total observations). The final community matrices consisted of 1500 and 976 ASVs for bacteria (N = 86 samples) and fungi (N = 87), respectively.

Floral microbial α- and β-diversity
To evaluate how floral microbial α-diversity was influenced by plant genotype, herbivory treatments (present: herbivory, herbivory + mycorrhizae; absent: control, mycorrhiza), and mycorrhizae treatments (present: mycorrhizae, herbivory + mycorrhizae; absent: control, herbivory), we analyzed the bacterial and fungal communities separately using general linear mixed models (LMMs) in package lme4 [41]. The response variable (Shannon diversity) was calculated using package vegan [42], and power transformed to improve normality with the optimal power parameter determined using the Box–Cox method in package car [43] (i.e. power parameter = 1 and 2 for bacterial and fungal Shannon diversity, respectively). The predictors included genotype, herbivory treatments and mycorrhizae treatments as well as their two-way interactions and three-way interaction, with initial plant weight as a covariate. The split plot with randomized complete block design in this experiment (Fig. 1) required nested random effects (i.e. treatment plots nested within blocks). However, due to only a subset of the 144 samples (N = 86 and 87 for bacterial and fungal data sets, respectively) for the LMMs, nested random effects led to difficulties in model convergence and thus were not used. Instead, the LMMs considered the random effect of blocks alone. Statistical significance (type III sums of squares) and least-squares means (LS-means) of predictors were assessed using packages lmerTest [44] and emmeans [45].

Microbial β-diversity (Bray–Curtis dissimilarity) was evaluated using PERMANOVA and constrained principal coordinates analysis (cPcoA) in vegan for bacterial and fungal communities separately. To assess the significance of the main effects, PERMANOVA and cPcoA included genotype, herbivory treatments and mycorrhizae treatments, with initial plant weight as a covariate and block as the random effect. To assess the significance of two-way interactions, PERMANOVA and cPcoA included both the main effects and their two-way interaction terms. Similarly, to assess the significance of the three-way interaction, the main effects, two-way interactions and three-way interaction were included. Once significant genotypic effect was identified, we further conducted the same PERMANOVAs and cPcoAs between pairwise genotypes to detect which specific genotypes were divergent from each other in microbial community composition.

Floral phenotype analysis
To evaluate whether strawberry genotypes differed in flower abundance (sample size N = 93) and flower size (N = 80), we conducted LMMs with the response variables power transformed to improve normality (i.e. power parameter = 0.5, square-root transformation, for flower abundance, and 2 for flower diameter). The predictors included genotype, herbivory treatments and mycorrhizae treatments as well as their two-way interactions and three-way interaction, with initial plant weight as a covariate and block as the random effect.

To evaluate whether strawberry genotypes differed in VOC profile (sample size N = 66), we conducted PERMANOVA and cPcoA. The predictors included genotype, herbivory treatments and mycorrhizae treatments as well as their two-way interactions and three-way interaction, with covariates (i.e. initial plant weight, pump ID and average daily temperature during VOC collection) and random effect (block). The PERMANOVA and cPcoA were conducted as in the above-mentioned microbial analysis to assess the significance of the main effects and interaction terms. Once significant genotypic effect was detected, we further identified the specific VOCs that differed between genotypes using random forest (RF) classification in packages caret [46] and randomForest [47]. The RF classification models were run for the full data with 1000 trees, and the number of randomly selected variables (i.e. individual VOCs) at each split of a decision tree was optimized using 10-fold cross validation in caret. RF model performance was assessed using out-of-bag (OOB) error. The set of important, non-redundant VOCs (in the presence of potentially correlated VOCs) were selected using backward variable elimination with package varSelRF [48].

SEMs linking floral phenotype to microbial α- and β-diversity
SEMs were conducted to link genotypic variation in flower abundance, size and VOCs, as well as treatments, to the α-diversity and β-diversity of floral bacterial and fungal communities (Figs. 4a and 5a). We focused on VOCs at three different levels: (1) individual VOCs that differed between genotypes as identified by the random forest classification (see above), and that predicted the α-diversity (Shannon diversity) or β-diversity (the first and second axis of cPcoA, cPcoA1 and cPcoA2) of bacterial and fungal communities based on backward predictor selection using packages caret and leaps [49]; (2) the major classes of VOCs (terpenes, aliphatics, benzenoids, S-containing compounds and CS branched-chain compounds); and (3) VOC profile (VOC cPcoA1 and cPcoA2; Fig. 3c). For the SEMs, flower abundance and flower diameter were power transformed (i.e. power parameter = 0.5 and 2, respectively). For the categorical variables of treatments, we coded...
Conflict of interest

The authors declare no conflict of interest.

Supplementary data

Supplementary data is available at Horticulture Research Journal online.

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