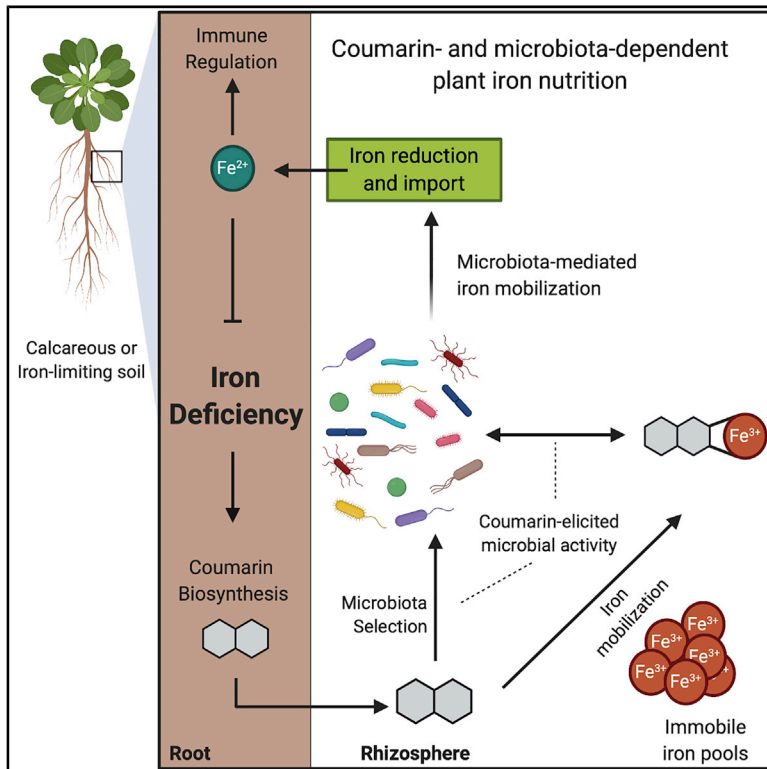


Cell Host & Microbe

Root-Secreted Coumarins and the Microbiota Interact to Improve Iron Nutrition in *Arabidopsis*

Graphical Abstract



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In Brief

Iron-limiting soils are widespread, causing significant losses in plant growth and productivity. Harbort et al. show that under iron limitation, plant-secreted coumarin compounds are mediators of a beneficial plant-microbiota interaction. These specialized metabolites alter root microbiota composition and are required for microbiota-mediated plant iron uptake and immune regulation.

Highlights

- Coumarins alter the root microbiota and improve plant growth in iron-limiting soil
- The microbiota improves plant iron nutrition via a coumarin-dependent mechanism
- The iron-beneficial commensal trait is taxonomically widespread but strain specific
- Coumarin-microbiota interaction resolves iron starvation and regulates immune response



Article

Root-Secreted Coumarins and the Microbiota Interact to Improve Iron Nutrition in *Arabidopsis*

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SUMMARY

Plants benefit from associations with a diverse community of root-colonizing microbes. Deciphering the mechanisms underpinning these beneficial services are of interest for improving plant productivity. We report a plant-beneficial interaction between *Arabidopsis thaliana* and the root microbiota under iron deprivation that is dependent on the secretion of plant-derived coumarins. Disrupting this pathway alters the microbiota and impairs plant growth in iron-limiting soil. Furthermore, the microbiota improves iron-limiting plant performance via a mechanism dependent on plant iron import and secretion of the coumarin fraxetin. This beneficial trait is strain specific yet functionally redundant across phylogenetic lineages of the microbiota. Transcriptomic and elemental analyses revealed that this interaction between commensals and coumarins promotes growth by relieving iron starvation. These results show that coumarins improve plant performance by eliciting microbe-assisted iron nutrition. We propose that the bacterial root microbiota, stimulated by secreted coumarins, is an integral mediator of plant adaptation to iron-limiting soils.

INTRODUCTION

Plant roots are colonized by a diverse community of microbes, collectively termed the root microbiota, originating from the surrounding soil biome (Bai et al., 2015; Bulgarelli et al., 2012; Lundberg et al., 2012). The structure of these communities is shaped by soil edaphic factors and root-secreted photosynthates and secondary metabolites (Berendsen et al., 2012; Bulgarelli et al., 2013). The root microbiota provides indirect protection against soil-borne fungal pathogens (Carrión et al., 2019; Durán et al., 2018) and is thought to improve host nutrition by improving the bioavailability of nutrients (Hacquard et al., 2015). However, the extent to which plants can selectively alter their microbiota and harness these beneficial traits in response to nutritional stress is unknown.

Iron is an essential mineral nutrient of plants, acting as a catalyst in many biological processes including photosynthesis and respiration. Although it is an abundant element in most soils, its bioavailability is often limiting due to its extremely low solubility at neutral and alkaline pH, as in calcareous soils containing a high proportion of calcium carbonate. Iron deficiency results in

stunted plant growth and leaf chlorosis and is responsible for decreased crop yields and nutrient content in ~30% of arable land (Morrissey and Guerinot, 2009). As such, there is great economic interest in improving plant iron nutrition. In response to iron-limiting conditions, non-graminaceous plants, such as *A. thaliana*, mount an iron starvation response that is coordinated by FER-LIKE IRON DEFICIENCY INDUCED TRANSCRIPTION FACTOR (FIT) (Colangelo and Guerinot, 2004; Jakoby et al., 2004) and a suite of associated basic helix-loop-helix (bHLH)-type transcription factors (Ivanov et al., 2012). This response serves to improve the solubility of iron through rhizosphere acidification by H⁺-ATPase AHA2 (Santi and Schmidt, 2009) and reduction of iron(III) to more-soluble iron(II) by plasma membrane protein FERRIC REDUCTION OXIDASE 2 (FRO2) (Robinson et al., 1999). Iron(II) is imported into the root epidermis by IRON-REGULATED TRANSPORTER1 (IRT1) (Vert et al., 2002).

The secretion of coumarins, phenolic secondary metabolites deriving from the general phenylpropanoid pathway, is also induced by iron starvation and is thought to contribute to iron nutrition through direct mobilization of recalcitrant iron pools (Fourcroy et al., 2014, 2016; Rajniak et al., 2018; Rodríguez-



Celma and Schmidt, 2013; Schmid et al., 2014; Schmid et al., 2014; Sisó-Terraza et al., 2016; Siwinska et al., 2018; Tsai et al., 2018). Three main coumarin compounds are produced in *A. thaliana* via a linear biosynthetic pathway (Figure 1A) (Rajniak et al., 2018; Tsai et al., 2018). FERULOYL-COA 6-HYDROXY-LASE1 (F6'H1) synthesizes scopoletin (Kai et al., 2008), which can be converted to fraxetin by SCOPOLETIN 8-HYDROXY-LASE (S8H) (Rajniak et al., 2018; Tsai et al., 2018), and further converted to sideretin by CYTOCHROME P450, FAMILY 82C4 (CYP82C4) (Rajniak et al., 2018). Each of these coumarins can be exported by the ABC transporter PLEIOTROPIC DRUG RESISTANCE 9 (PDR9) (Fourcroy et al., 2014), though other efflux pumps may also contribute to the export of some coumarins (Ziegler et al., 2017). Coumarin secretion was recently shown to influence the structure of root microbial communities in artificially limed soil (Stringlis et al., 2018) and synthetic media (Voges et al., 2019). However, the impact of coumarin secretion on the root microbiota in soils with different mineral nutrient availabilities and the consequences for plant productivity remain undefined.

RESULTS

Coumarin Biosynthesis Is Important for Plant Growth and Root Microbiota Composition in Iron-Limiting Soil

To assess potential links between coumarin secretion, plant growth, and the root-associated microbiota, we grew *A. thaliana* Col-0 wild-type (WT) plants and mutants with defects in coumarin biosynthesis or export (Figure 1A) on two soils with contrasting iron availability. These mutants were previously shown to have abolished coumarin biosynthesis at their respective steps in the pathway (Rajniak et al., 2018; Schmid et al., 2014; Tsai et al., 2018; Ziegler et al., 2017). Cologne agricultural soil (CAS), obtained from a local site, is slightly acidic with pH 6.5, at which iron is sufficiently available. We also obtained soil from a vineyard in Italy, termed here Italian soil (IS), which is alkaline and calcareous (pH 7.5, 9.7% of active CaCO₃). These conditions significantly reduce the availability of iron, despite this soil having a higher total iron content than CAS (Figure S1). We observed a decrease in shoot fresh weight (SFW) and leaf chlorophyll content in *f6'h1* and *s8h* plants grown on IS, whereas the measured performance parameters of all genotypes were indistinguishable on CAS (Figures 1B and 1C). A similar growth deficit was observed in *f6'h1* plants grown on other alkaline soils isolated from geographically diverse sites and could be improved by supplementation with solubilized iron (Figure S1). These results show that coumarin biosynthesis is important for growth in naturally iron-limiting soils. The performance of *pdr9* plants, however, was indistinguishable from WT on both CAS and IS; thus, coumarin export via the ABC transporter PDR9 was not crucial for growth in iron-limiting soils (Figures 1 and S1). This is in contrast to reported germ-free growth on synthetic media (Fourcroy et al., 2014; Rodríguez-Celma et al., 2013) (and Figure 5 below). These data suggest that in soil, sufficient coumarin export may occur via additional members of the diversified and promiscuous ABC transporter family (Borghi et al., 2019; Ziegler et al., 2017).

The root-associated bacterial microbiota of plants grown on CAS and IS were assessed by culture-independent 16S rRNA

gene amplicon sequencing and analyzed at the amplicon sequence variant (ASV) level. Alpha diversity was greater in IS than CAS for both unplanted soil and root samples but did not vary by genotype (Figure S2). Unconstrained principal coordinate analysis (PCoA) of Bray-Curtis distances between samples showed that the soil type and batch were the largest drivers of divergence between samples (Figure S2). PCoA analysis of beta diversity constrained (CPCoA) for the interaction between genotype and soil type revealed a significant separation of *f6'h1* and *s8h* plants from other genotypes when grown on IS, but not on CAS (Figure 1D). Analysis of bacterial community profiles from three batches of each soil type confirmed that *f6'h1* plants separate significantly from other genotypes when grown in IS, but not in CAS (Figure S2). Together, these results indicate that coumarin biosynthesis, especially of scopoletin and fraxetin, is important for plant growth and determining root microbiota composition in a naturally iron-limiting calcareous soil but is dispensable in an iron-replete soil. Furthermore, this illustrates that the interaction between soil type and plant genotype can serve as a major determinant of root microbiota structure, explaining 27.7% of community variation (Figure 1D).

Coumarin Biosynthesis Restructures the Root Microbiota at the ASV Level

In order to explore the nature of the observed changes in community structure, we determined which ASVs were differentially enriched (deASVs) in each mutant genotype compared with WT in each soil. For this analysis, we pooled data from three batches of each soil (except *s8h*, which was included in only one batch). The greatest number of deASVs was observed in *f6'h1* plants, with significantly more detected when grown in IS than CAS (260 deASVs in IS, 50 in CAS; Figure 2A). The impact of deASVs on the microbiota structure in terms of relative abundance was also greatest in *f6'h1* plants grown in IS (Figure S3).

Taxonomic analysis revealed that multiple bacterial families were significantly over- or under-represented within the deASV subset for *f6'h1* plants in IS compared with the full list of detected ASVs (Figures 2B and S3). *Burkholderiaceae* was the most prevalent family detected within the deASVs (56 deASVs; 2.65-fold enriched in deASVs subset compared with all ASVs detected). However, a correlation between coumarin production and deASV relative abundance could not be generalized at family-level resolution; most families contained deASVs which were more abundant in *f6'h1* plants than WT, as well as deASVs which were less abundant. Some patterns were observed at the genus level (Figure S3), but due to the overall low number of deASVs in each genus, their statistical significance remains unclear. This indicates that coumarin production restructures the root microbiota at the ASV level within multiple bacterial families. Despite being the family containing the most deASVs, the relative abundance of *Burkholderiaceae* was not significantly altered in coumarin-deficient plants (Figure 2C). The relative abundances of the next two most-impacted families, *Rhizobiaceae* and *Streptomyetaceae*, however, were slightly but significantly altered in *f6'h1* plants on IS. This indicates that, at the ASV level, disruption of coumarin biosynthesis has a quantitative impact within multiple root-associated commensal families, but with relatively minor effects on the microbiota structure at higher taxonomic ranks. This suggests the existence of ASV-level

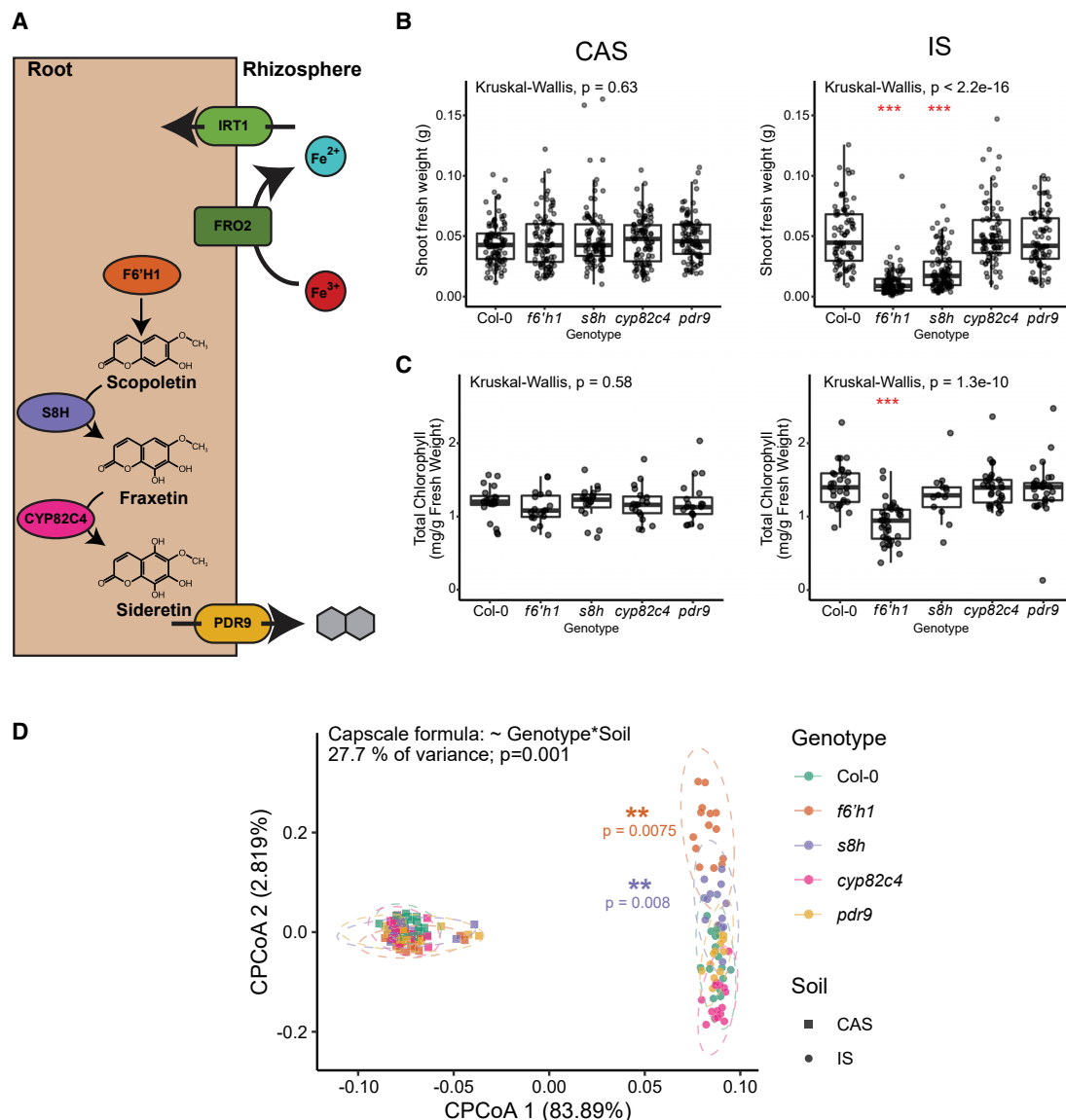


Figure 1. Coumarin Biosynthesis Is Important for Plant Growth and Root Microbiota Composition in a Naturally Calcareous Soil

(A) Diagram of pathways for coumarin biosynthesis and export, and reductive uptake of iron in *Arabidopsis*.

(B and C) (B) SFW and (C) total chlorophyll content of coumarin pathway mutants grown in a non-calcareous (CAS) and a calcareous (IS) soil. Statistical significance was determined by Kruskal-Wallis; each mutant was compared with Col-0 by Wilcoxon Ranked Sum post-hoc. Significance is indicated by red asterisks (*, **, ***, indicate $p < 0.05$, 0.01, and 0.001, respectively). For shoot fresh weight measurements, Col-0 $n = 171$, 204; *f6'h1* $n = 168$, 272; *s8h* $n = 93$, 113; *cyp82c4* $n = 164$, 209; and *pdr9* $n = 172$, 169 in CAS and IS, respectively. Chlorophyll content was measured from pooled leaf samples, (Col-0 $n = 35$, 29; *f6'h1* $n = 34$, 36; *s8h* $n = 19$, 14; *cyp82c4* $n = 34$, 30; and *pdr9* $n = 35$, 30 in CAS and IS, respectively).

(D) Constrained ordination of root bacterial community composition of coumarin pathway mutants, constrained for the interaction between soil and genotype. Ellipses delineate multivariate normal distribution at 95% confidence. Data are from one representative experiment of three (Col-0 $n = 17$, 15; *f6'h1* $n = 18$, 14; *s8h* $n = 15$, 14; *cyp82c4* $n = 18$, 15; and *pdr9* $n = 17$, 14 in CAS and IS, respectively). p values represent significance of separations between genotypes within each soil determined by pairwise PERMANOVA. Only *f6'h1* (orange) and *s8h* (purple) were significantly separated from Col-0 in IS. See also: Figures S1 and S2.

compensatory mechanism(s) within bacterial families which, during root microbiota establishment, maintain higher taxonomic structure in coumarin-deficient plants on iron-limiting soil. An increase in the number of deASVs was also observed for *s8h* plants grown in IS compared with CAS (Figure 2A), though fewer deASVs overall were detected compared with other mutants as this genotype was only included in one experimental replicate.

Consistently, the taxonomic profile of *s8h* deASVs was also enriched for *Burkholderiaceae*, and had considerable, but not complete, overlap with the deASVs detected on *f6'h1* plants in the same experiment (Figure S3). This indicates that production of both scopoletin and fraxetin impact the root microbiota.

As various coumarin compounds have been shown to exert antimicrobial activity, we examined the coumarin sensitivity of

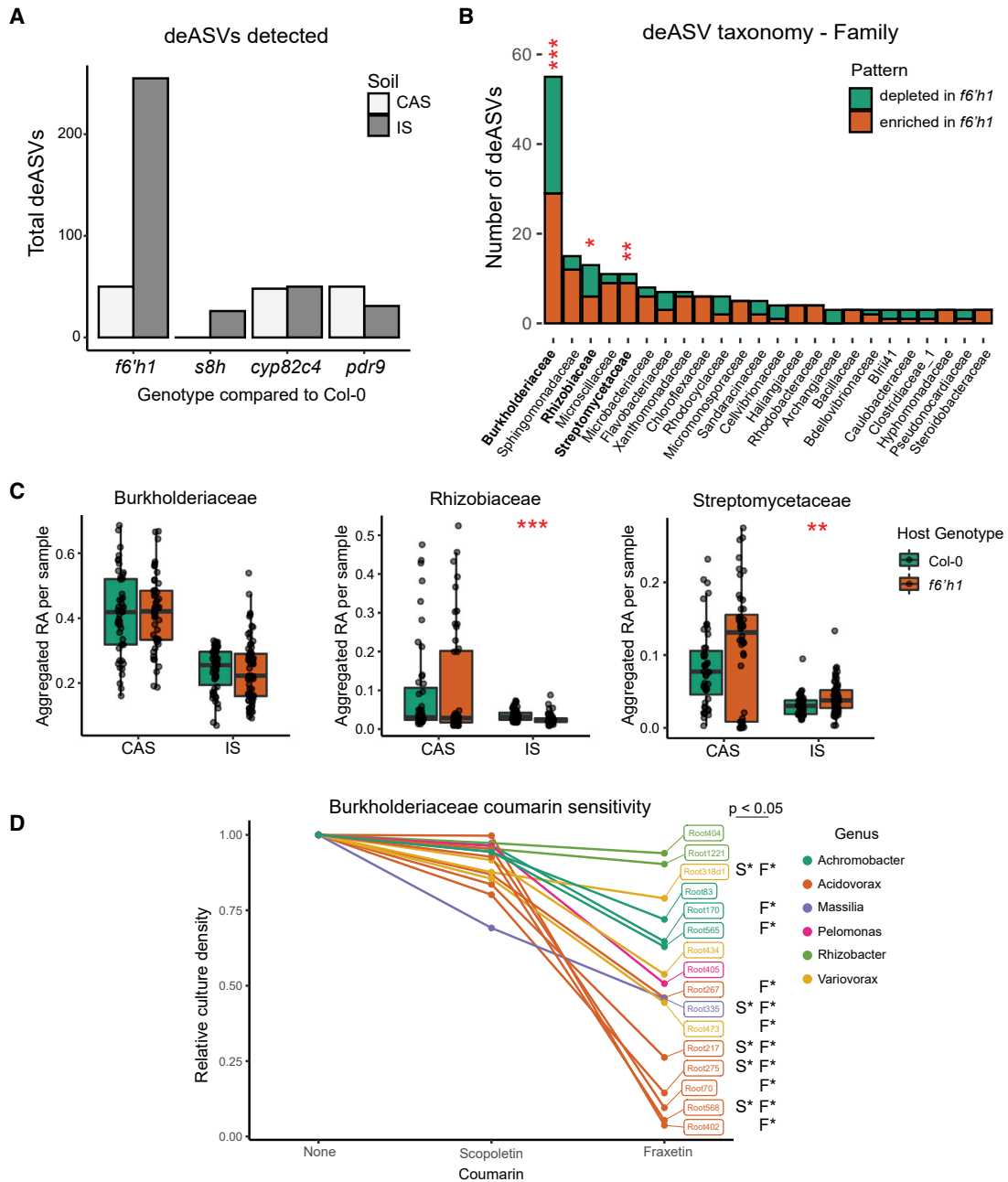


Figure 2. Coumarin Biosynthesis Restructures the Root Microbiota at the ASV Level

(A) Number of deASVs detected in indicated mutants compared with Col-0 in each soil. Data are pooled from three experiments (except *s8h*, which was included in only one), and filtered for ASVs found in at least three samples with RA > 0.05%. Differential enrichment was calculated using a negative binomial generalized log-linear model at an FDR-adjusted p value of 0.05.

(B) Family-level taxonomic classification of deASVs in *f6'h1* plants growing on IS. Colors indicate if deASVs were enriched or depleted in *f6'h1* compared with Col-0. Hypergeometric enrichment test was performed to determine if each family was over- or under-represented in deASV list compared with all detected ASVs. Red asterisks indicate significance with FDR-adjusted p values.

(C) Sample-wise aggregated relative abundance of the top three families most significantly over-represented in deASVs: *Burkholderiaceae*, *Rhizobiaceae*, and *Streptomycetaceae*. Each data point represents the average RA aggregated at the family level in a single sample. Significance between genotypes in each soil was determined by Wilcoxon ranked sum test.

(D) Overnight growth of *Burkholderiaceae* bacterial strains in the presence of scopoletin or fraxetin. Optical density (OD) of cultures was normalized to the OD of each strain in the absence of coumarins. Significant differences ($p \leq 0.05$ by Tukey's HSD) in growth compared with the control are indicated for scopoletin (S*) and fraxetin (F*) to the right of each strain. Data are averages of 2–4 experiments, each with 2–3 technical replicates, per strain. *, **, and *** in (B) and (C) indicate $p \leq 0.05$, 0.01, and 0.001, respectively. See also Figures S2 and S3.

a panel of root commensal *Burkholderiaceae* strains, the most prevalent family within the deASVs on *f6'h1* and *s8h* plants. These strains were previously isolated from roots of *Arabidopsis* growing in CAS soil (Bai et al., 2015). Bacterial growth was quantified in the presence of 50 μ M scopoletin or fraxetin, a concentration within the physiological range of coumarins observed within roots (Siwinska et al., 2014, 2018; Stringlis et al., 2018). We observed prevalent growth inhibition to a range of degrees in the presence of fraxetin, and to a lesser extent in response to scopoletin (Figure 2D). This indicates that fraxetin exerts variable antimicrobial activity on *Burkholderiaceae* strains. This strain-specific variation potentially explains part of the ASV-level shifts within *Burkholderiaceae* observed between WT and coumarin-deficient plants in iron-limiting soil.

Taxonomically Diverse Root Commensals Improve Iron-Limiting Plant Performance

To assess the impact of root commensals on plant performance under iron-limiting conditions, we employed an agar medium-based gnotobiotic system that allows control over both iron mobility and the presence of bacterial commensals. In this system, nutrient medium (1/2 MS) is strongly buffered at pH 7.4, similar to the pH of iron-limiting IS soil. Iron is provided at 100 μ M in one of two forms: available iron (avFe) FeEDTA, a complex that retains solubility even at alkaline pH, or an unavailable form (unavFe) FeCl₃, which forms Fe(OH)₃ and is highly insoluble at alkaline pH. Providing unavFe mimics iron-limiting conditions in calcareous and alkaline soils such as IS; iron is present but recalcitrant due to extremely low solubility and must be actively mobilized for utilization. Using this gnotobiotic system, we reconstituted plants with a synthetic community (SynCom) of bacterial commensals to assess the impact of the microbiota on iron-limiting plant performance. To achieve this, we took advantage of a diverse culture collection of bacterial commensals isolated from *A. thaliana* roots grown in CAS (Bai et al., 2015). We designed a taxonomically diverse SynCom of 115 members, which reflects the root bacterial diversity observed by culture-independent methods at high taxonomic ranks (Figure 3A; Table S2). SFW and chlorophyll content were measured as readouts of plant performance and as a proxy measurement for iron nutritional status.

A growth deficit and leaf chlorosis (Figures 3B and 3C), symptoms of iron starvation, were observed in plants grown on unavFe. Furthermore, elemental and transcriptomic analyses revealed decreased leaf iron content and induction of iron import components FRO2 and IRT1 and repression of ferritins, iron storage proteins (Figures 7 and S7). Together, these data confirm a robust induction of iron deficiency in this experimental system. We found that reconstitution of the microbiota with a bacterial SynCom greatly improved both SFW and chlorophyll content of plants grown on unavFe. This beneficial interaction was unexpected, given that iron starvation-induced coumarins exert antimicrobial activity against some commensals (Stringlis et al., 2018; Voges et al., 2019; Wang et al., 2014; Yang et al., 2016) (Figure 2D). Iron starvation was also induced by reducing the supply of soluble iron to an insufficient amount (1 μ M FeEDTA). However, commensal-mediated improvement of plant performance was not observed in plants grown on media containing only insufficient soluble iron (Figure 3C). Plant growth limitation

on unavailable iron and growth rescue by a live SynCom were maintained when plant roots were shielded from light (Figure S4), supporting the robustness of this experimental system. These results suggest that bacterial commensals can improve plant performance by improving access to immobile sources of iron.

To survey the ability of various taxa to improve iron-limiting plant growth, we grew plants in mono-associations with bacterial strains on unavFe. Fifty-three taxonomically diverse SynCom strains were tested for their ability to rescue iron-limiting plant growth (Figure 3A, red arrows; Table S2). Within each broader taxonomic lineage, we observed growth-rescuing strains as well as strains lacking this ability (Figure 3D), demonstrating the ubiquity of this beneficial activity as well as the strain-specific variation within all core taxonomic lineages of the root microbiota. Thus, in a community context, the capacity of bacterial commensals to rescue plant growth under iron-starved conditions is functionally redundant. Furthermore, the ability of these strains to grow in the presence of scopoletin and fraxetin was found not to correlate with plant growth rescue capacity, indicating that this plant-beneficial trait does not require selection via coumarin antimicrobial activity (Figure S4).

Microbiota-Mediated Plant Growth Rescue Occurs via Reductive Import of Iron and Requires Fraxetin Secretion

We utilized *A. thaliana* mutant lines deficient in genes involved in iron uptake and homeostasis to determine their importance for bacteria-mediated growth rescue under iron limitation. Mutants in components of the reduction-based iron uptake system (*fro2* and *irt1*), rhizosphere acidification (*aha2*), and a negative regulator of the iron starvation response (*bts*) were grown on unavFe in the presence of a live SynCom, or a heat-killed SynCom as a negative control. Genotypes *fro2* and *irt1* displayed an exaggerated growth deficit and leaf chlorosis (Figure 4), consistent with their reported hypersensitivity to iron starvation (Robinson et al., 1999; Vert et al., 2002). In contrast to WT plants, addition of the bacterial SynCom was unable to improve the phenotype of these iron import mutants. Performance of *irt1* plants was further reduced in the presence of the SynCom. This indicates that without the iron import channel, plants may be unable to compete with bacterial commensals for access to the already limited pool of available iron. Moreover, *bts* plants, which are tolerant to iron deficiency (Hindt et al., 2017; Selote et al., 2015), were larger than WT plants on unavFe, not chlorotic, and still displayed slightly improved performance when inoculated with the SynCom. No difference was observed between *aha2* and WT controls, however, indicating that plant-mediated rhizosphere acidification is not rate limiting for commensal-mediated plant growth rescue in strongly buffered alkaline conditions. When grown on avFe, the SynCom did not improve plant performance in any of these genotypes (Figure S5). These results validate our gnotobiotic system for microbiota reconstitution under iron-limiting conditions, confirm that the growth limitation and chlorosis on unavFe is due to iron starvation, and suggest that improved plant performance in the presence of commensals depends on the plant's endogenous system for iron reduction and import.

We next investigated the role of coumarins in commensal-mediated plant growth rescue under iron limitation. The addition of the SynCom provided no benefit to plant growth

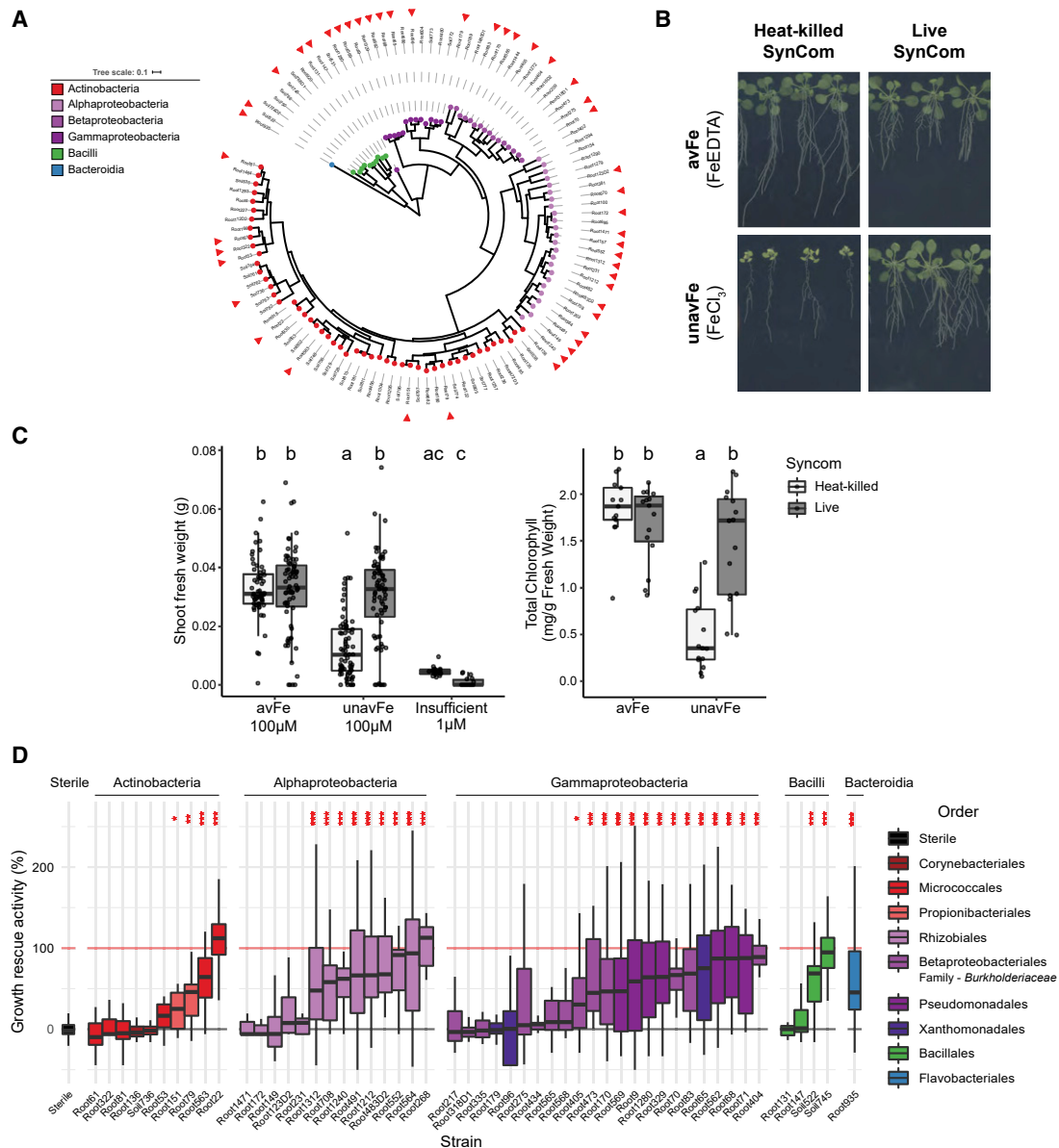


Figure 3. Taxonomically Diverse Root Commensals Improve Iron-Limiting Plant Performance

(A) Phylogenetic tree of 115-strain SynCom derived from At-RSPHERE culture collection (Bai et al., 2015) used for microbiota reconstitution. Red arrows indicate strains used in (D).

(B) Representative images of plants grown for 2 weeks on media containing available (avFe) and unavailable (unavFe) forms of iron inoculated with live SynCom or heat-killed control.

(C) SFW and total chlorophyll quantification of Col-0 plants after 2 weeks of growth on indicated iron conditions. Data are pooled from three experiments with avFe and unavFe: n = 42–54 plants per condition, and chlorophyll measured in pooled samples, n = 13–15 per group. Insufficient iron data are from one experiment, n = 18 plants. Letters indicate significant pairwise differences between groups ($p\text{-adj} \leq 0.05$ by Dunn's pairwise comparison with Bonferroni correction for SFW, and Tukey's HSD corrected for multiple comparisons for chlorophyll content).

(D) Iron-limiting growth rescue activity of SynCom strains in mono-association. SFW was measured and plotted as percent growth rescue of bacteria-inoculated plants on unavFe compared with the growth deficit between sterile plants on avFe versus unavFe. Black and red lines indicate 0% (axenic plants on unavFe) and 100% growth rescue (axenic plants on avFe), respectively. Data are pooled from 1–4 experiments per strain and normalized to respective sterile controls (n = 18 plants per experiment). Asterisks indicate significance from sterile plants by Wilcoxon ranked sum test with FDR adjustment (*, **, and *** indicate $p\text{-adj} \leq 0.05$, 0.01, and 0.001, respectively). See also Figure S4.

or chlorophyll content of *f6'h1*, *s8h*, or *pdr9* plants grown on unavFe (Figure 5). In contrast, the SynCom improved performance of *cyp82c4* plants similar to WT. These data suggest that plant biosynthesis of scopoletin and/or fraxetin (cata-

lyzed by F6'H1 and S8H, respectively) and their secretion (through PDR9) are required for bacteria-mediated plant growth rescue under iron limitation, while sideretin (synthesized by CYP82C4) is dispensable. No growth promotion by the

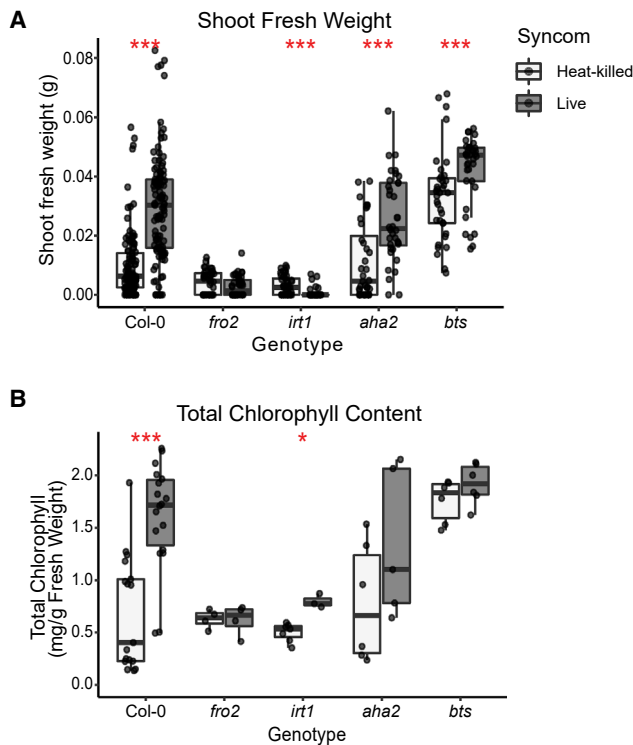


Figure 4. Microbiota-Mediated Plant Growth Rescue Occurs via the Reductive Import of Iron

(A and B) (A) SFW and (B) leaf chlorophyll content of indicated mutants in the reductive import of iron pathway grown on unavFe media inoculated with heat-killed or live bacterial SynCom. Total chlorophyll content was measured in pooled leaf samples from six plants. Data are from two independent experiments per genotype ($n = 36$ plants, 6 chlorophyll samples). Each experiment included Col-0 control ($n = 90$ plants, 18 chlorophyll samples). Asterisks indicate significance between heat-killed- and live SynCom-inoculated groups by Wilcoxon ranked sum test for SFW and Student's *t* test for chlorophyll content (*, **, and *** indicate $p \leq 0.05$, 0.01, and 0.001, respectively). See also Figure S5.

SynCom was observed in these genotypes grown on avFe (Figure S5).

To further assess the roles of scopoletin and fraxetin in commensal-mediated plant growth rescue, we chemically complemented *f6'h1* plants by supplementing the growth medium with each compound (Figures 6A and 6B). Addition of scopoletin did not improve plant performance, while fraxetin fully restored the ability of the SynCom to improve both plant growth and leaf chlorophyll content in coumarin-deficient *f6'h1* plants. That scopoletin was unable to complement *f6'h1* plants suggests that external scopoletin is not sufficiently taken up by roots and converted to fraxetin after secretion. Furthermore, the ability of the SynCom to rescue growth of *s8h* plants was also fully restored by supplementation with scopoletin and fraxetin together (Figure 6C). This confirms that fraxetin is the necessary coumarin structure type for commensal-mediated growth rescue. Notably, supplementation with coumarins failed to rescue growth or chlorophyll content of germ-free *f6'h1* or *s8h* plants at 50 μM , a concentration lower than what has been used to rescue iron starvation by directly mobilizing iron. This indicates that commensal-mediated improvement in iron-limiting

plant growth is induced by fraxetin concentrations lower than those required for sufficient mobilization of iron in axenic conditions. Together, these results confirm that secreted fraxetin is both necessary and sufficient to elicit growth rescue activity from bacterial commensals under iron limitation. These findings argue for an indirect activity of fraxetin in recruiting commensal-mediated mobilization of recalcitrant iron pools in addition to its direct iron-mobilizing activity.

Coumarins and the Microbiota Interact to Alleviate Iron Starvation

To determine if the observed iron starvation symptoms and their improvement by root commensals correlated with plant iron status, we measured leaf elemental content by inductively coupled plasma mass spectrometry (ICP-MS) (Figures 7A and S6). Leaf iron concentration was significantly reduced on unavFe, confirming iron deficiency. Furthermore, addition of a live SynCom to WT, but not *f6'h1* plants, restored plant iron content to replete avFe levels. However, addition of the SynCom had no impact on plant iron content on avFe. These results confirm that the plant growth rescue activity by commensals during iron starvation is due to improved iron nutrition. We next sought to determine if this nutritional benefit was a result of microbial stimulation of the plant iron deficiency response or increased iron availability in the presence of the SynCom. To identify plant pathways responding to the presence of coumarins and bacterial commensals under different iron regimes, we performed analysis of the whole-root transcriptome of WT and *f6'h1* seedlings in our gnotobiotic system. An earlier time point (1 week) was chosen to observe potential stimulation of iron deficiency response genes by the SynCom (Verbon et al., 2019) that may lead to plant growth rescue. At this time point, SynCom-mediated plant growth rescue was observed in WT, but growth was still comparable between WT and *f6'h1* plants (Figure S7).

Plotting transcriptome sample distances by PCA (Figure 7B) revealed that the supplied iron form was the largest determinant of dissimilarity (PC1, 18% of variance), followed by SynCom status (PC2, 9% of variance). When grown on avFe, both Col-0 and *f6'h1* plants clustered together, but separation was observed based on SynCom status. This indicates that a live SynCom impacts host transcriptional responses when iron is available, independently of plant coumarin status. The transcriptomes of plants grown on unavFe, however, were distinct from those of plants grown on avFe, and further separated based on both genotype and SynCom status. When inoculated with a heat-killed SynCom, both WT and *f6'h1* on unavFe separated from the avFe cluster (upper-left quadrant). A larger genotype-driven separation was observed between Col-0 and *f6'h1* plants when inoculated with a live SynCom on unavFe. Remarkably, WT plants inoculated with a live SynCom on unavFe clustered closely with SynCom-inoculated WT plants on avFe (lower-right quadrant), while *f6'h1* plants remained in the lower-left quadrant, clearly separated from the avFe cluster. This pattern indicates that the transcriptional responses to growth on unavFe are more pronounced in *f6'h1* plants than WT, consistent with their hypersensitivity to iron starvation. Indeed, more iron-responsive differentially expressed genes (DEGs) were detected in *f6'h1* than WT plants (Figure S7). Furthermore, the number of iron-responsive DEGs indicates that the iron starvation-induced

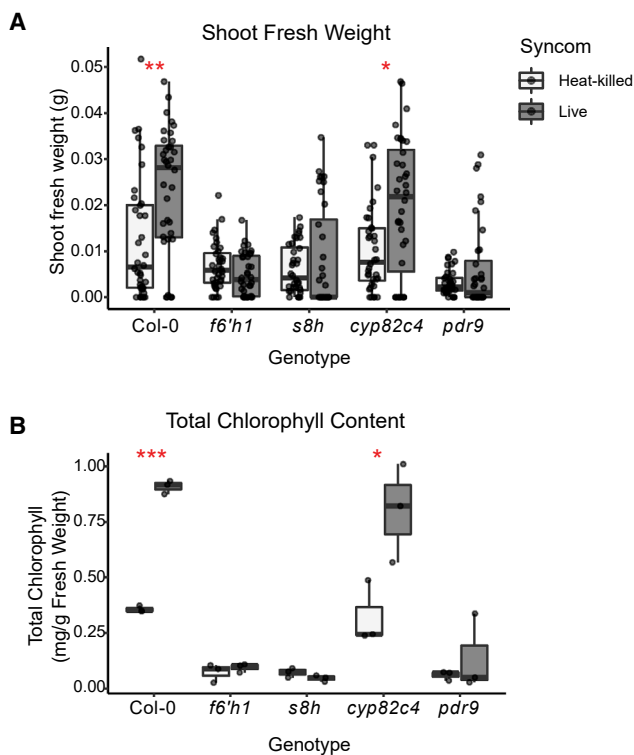


Figure 5. Plant Biosynthesis and Secretion of Fraxetin Is Necessary for Microbiota-Mediated Growth Rescue

(A and B) (A) SFW and (B) leaf chlorophyll content of indicated coumarin biosynthesis and export mutants grown on unavFe media inoculated with heat-killed or live bacterial SynCom. SFW data are from two experiments ($n = 36$ plants). Chlorophyll content is from one experiment ($n = 3$ pooled leaf samples). Asterisks indicate significance between heat-killed- and live SynCom-inoculated groups by Wilcoxon ranked sum test for SFW and Student's t test for chlorophyll content (*, **, and *** indicate $p \leq 0.05$, 0.01 , and 0.001 , respectively). See also Figure S5.

response was mitigated by the addition of the SynCom in Col-0 but not *f6'h1* plants.

We performed k -means clustering of all transcripts based on expression pattern and identified DEGs to investigate the interaction between iron availability, SynCom, and genotype (Figure 7C). The identified clusters showed iron- and SynCom-responsive gene sets (plots left of heatmap clusters). Cluster 4 revealed a set of genes activated in axenically grown Col-0 and *f6'h1* plants in response to unavFe, which were more strongly induced in *f6'h1* plants. Furthermore, their expression is reduced to homeostatic levels in Col-0 plants upon addition of the SynCom but remained elevated in *f6'h1* plants. This gene set corresponds to iron-responsive genes that were also responsive to SynCom in a coumarin-dependent manner. Gene ontology analysis revealed that this cluster is enriched for genes belonging to the iron starvation response, iron homeostasis, and metal ion transport (Figure 7C, annotations right of heatmap). Genes in cluster 8 displayed the inverse pattern: downregulated on unavFe and restored in the presence of the SynCom in Col-0, but not *f6'h1*. We compared our DEGs to a list of 25 previously identified core iron starvation marker genes (Mai et al., 2016) (Figure S7). Of the 12 genes reported to be induced under iron starvation, 11 were found in cluster 4, while 7

out of 13 genes reported to be downregulated under iron starvation were present in cluster 8. The cluster assignment and expression patterns (Figures 7D and S7) of selected iron homeostasis regulators and coumarin biosynthesis genes revealed that these genes are iron- and SynCom responsive in a genotype-dependent manner. Importantly, the addition of live SynCom did not stimulate expression of iron deficiency genes. Iron starvation-upregulated genes (including bHLH39, FRO2, IRT1, and MYB72), as well as iron starvation-downregulated genes (including FER1) displayed expression patterns consistent with alleviation of the iron deficiency response by the SynCom in WT plants. This indicates that, rather than biostimulation of plant iron uptake, iron nutrition is improved by a commensal mechanism. In addition, the inability of the SynCom to alleviate the iron-starvation response in *f6'h1* roots (cluster 4 and 8 genes; Figures 7C and 7D) supports a mechanism by which secreted coumarins are required to elicit microbiota-mediated iron nutrition, rather than two parallel mechanisms exerted by coumarins and the microbiota independently. Together, these data reveal a robust induction of iron starvation in plants grown on unavFe, which is alleviated by commensals in WT, but not in *f6'h1* plants. This implies the existence of a coumarin- and microbiota-dependent mechanism that improves plant performance via iron nutrition.

Analysis of SynCom-responsive genes also revealed a core set of DEGs common to both genotypes independently of iron availability (clusters 3 and 10, Figures 7C and S7). These clusters were enriched for genes related to defense responses and response to bacteria. Interestingly, genes associated with these terms were also significantly enriched in clusters 2 and 8, the expression patterns of which are dependent on host genotype, iron status, and SynCom (Figure 7C). The presence of immune-related genes in clusters 2 and 8 suggests that, in addition to the core SynCom-responsive genes, a subset of defense genes is regulated by the presence of commensals in a coumarin-dependent manner. Collectively, these results show that both coumarin secretion and the root microbiota have profound impacts on plant transcriptional responses to iron deprivation. Furthermore, this emergent interaction between coumarins and the microbiota improves plant performance through bolstered iron nutrition, resolving the iron starvation response and regulating a subset of defense-related genes.

DISCUSSION

Our results reveal unexpected impacts of root-secreted coumarins governing plant-bacteria interactions, including soil type-dependent alteration of root microbiota composition, elicitation of a commensal-mediated mechanism of plant iron nutrition, and regulation of a subset of defense genes. We show that *f6'h1* and *s8h* plants, which are deficient in the biosynthesis of scopoletin and fraxetin, assemble an altered root bacterial community. Individual members of the *Burkholderiaceae*, core members of the plant root microbiota that often exert plant-beneficial activities (Eberl and Vandamme, 2016; Thiergart et al., 2020), are impacted by fraxetin in a strain-specific manner in both roots and growth in culture. This strain-specific sensitivity may in part underlie ASV-level changes in abundance on iron-limiting IS. The greater impact on the microbiota observed in *f6'h1* compared with *s8h* plants, in terms of deASVs detected and their relative

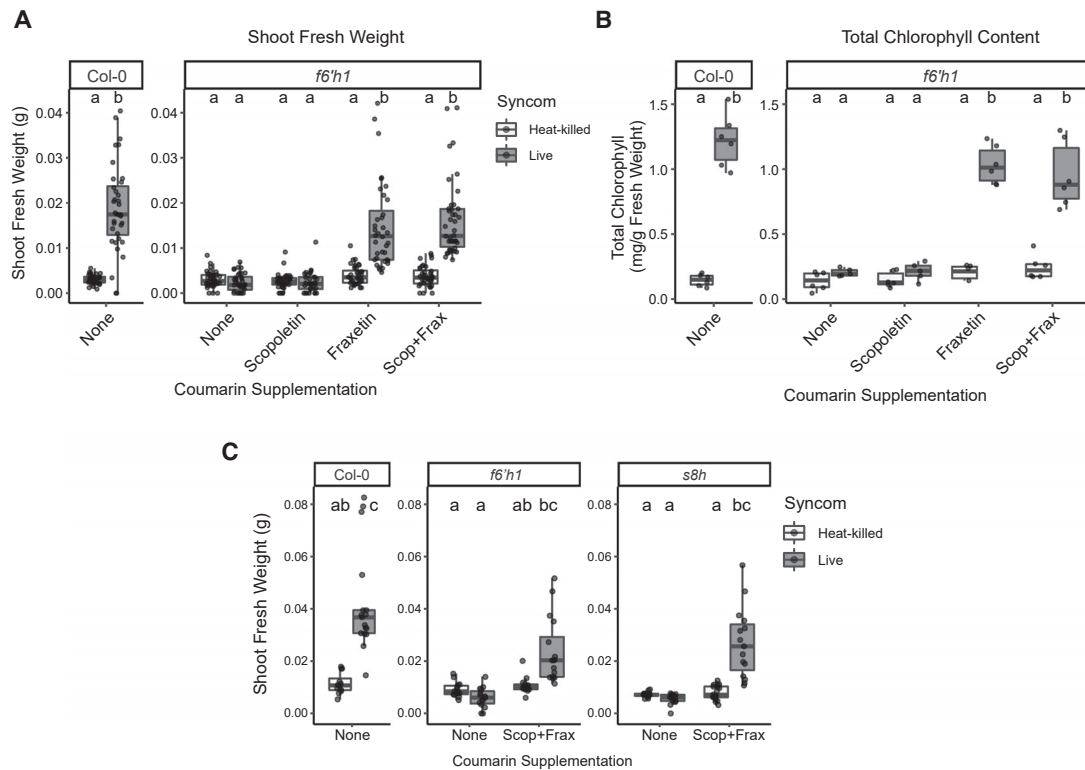


Figure 6. Supplementation with Fraxetin Restores Microbiota-Mediated Growth Rescue of *f6'h1* and *s8h* Plants

(A and B) (A) SFW and (B) leaf chlorophyll content of Col-0 plants, and *f6'h1* plants grown on unavFe supplemented with 50 μ M scopoletin and/or fraxetin and inoculated with heat-killed or live SynCom.

(C) SFW of Col-0, and *f6'h1* and *s8h* plants grown on unavFe supplemented with 50 μ M scopoletin and fraxetin and inoculated with heat-killed or live SynCom. Data in (A) and (B) are from two experiments ($n = 30$ – 42 plants, 6 pooled leaf chlorophyll samples). Data in (C) are from a single experiment ($n = 18$ plants). Letters indicate significant pairwise differences between groups (p -adj ≤ 0.05 by Dunn's pairwise comparison with Bonferroni correction for SFW, and Tukey's HSD corrected for multiple comparisons for chlorophyll content). See also Figure S5.

abundance (RA), suggests that both scopoletin and fraxetin impact bacterial microbiota assembly. Indeed, a metagenome analysis indicated an altered microbial multi-kingdom assemblage and provided evidence for selective scopoletin anti-fungal activity against soil-borne fungal pathogens *in vitro* (Ba et al., 2017; Carpinella et al., 2005; Kai et al., 2006; Stringlis et al., 2018). However, we cannot exclude that iron malnutrition in *f6'h1* plants has additional consequences on the microbiota.

Plant performance data, coupled with elemental content and transcriptomic analysis, confirm that benefits conferred by commensals under iron limitation occur via improved iron nutrition. In contrast to this beneficial interaction, under low phosphate concentrations *A. thaliana* must compete with a bacterial SynCom for access to the macronutrient, requiring integration of phosphate starvation and defense responses (Castrillo et al., 2017). We similarly observe emergent effects of coumarins and the microbiota on a subset of defense-related genes, indicating potential trade-offs between growth and defense. Of note, the impact of commensal communities on plant performance when phosphate is present in unavailable forms, as is characteristic of many soils, has not been tested. Importantly, in our system, growth promotion by microbes is observed only when iron is present but immobile, conditions characterizing most iron-limiting soils. Thus, our results highlight the importance of studying plant

nutritional phenotypes in systems closely mimicking natural conditions, including the presence of commensals and defined forms of soil minerals that are unavailable to the plant host. To the best of our knowledge, experimental evidence for clear plant nutritional benefits from commensals in a community context has not been reported before. This is a significant step forward in understanding how plant nutrition and productivity can be bolstered by harnessing endogenous soil microbes.

The presence of this beneficial activity across all core taxonomic lineages of the *A. thaliana* bacterial microbiota suggests that the underlying molecular mechanism(s) evolved independently rather than by common descent. As growth rescue depends on plant expression of the iron reductive import machinery, but does not involve microbial stimulation of its expression, the mechanism(s) must function upstream of reduction and import at the root surface. Multiple bacterial molecules can mobilize insoluble iron and are potentially utilized by plants, including siderophores and other metabolites, though the ability of plants to utilize microbially mobilized iron in the context of an intact microbiota has yet to be shown. The nutritional benefits provided by commensals requires plant-secreted fraxetin but is independent of its antimicrobial activity. This suggests that the impact of coumarins on root microbiota composition and on commensal-mediated iron nutrition may be independent

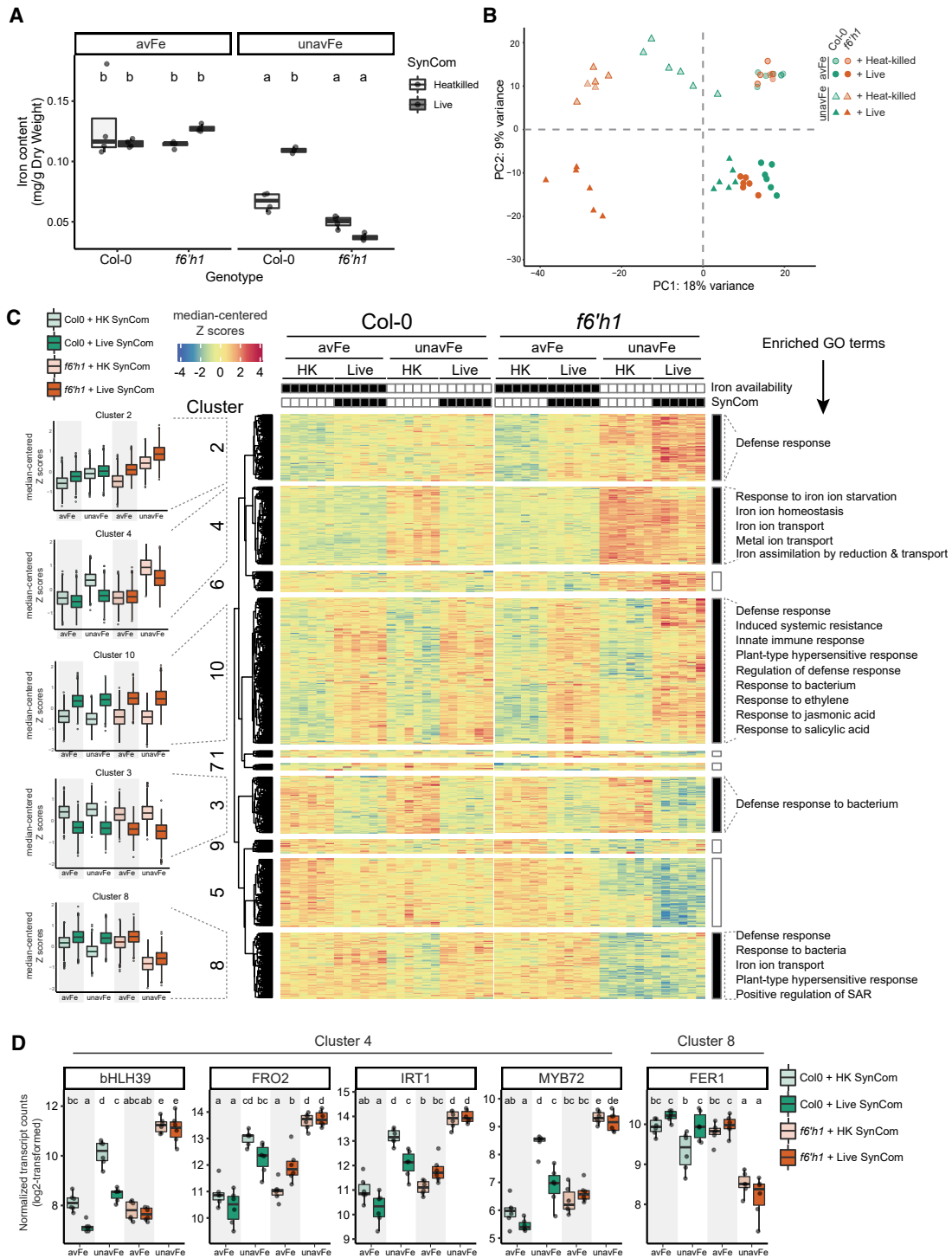


Figure 7. A Bacterial SynCom Improves Plant Iron Nutrition, Relieves the Iron Deficiency Response, and Modulates a Subset of Defense Genes in a Coumarin-Dependent Manner

(A) Shoot iron content of Col-0 and *f6'h1* plants grown on avFe and unavFe media with a live SynCom or heat-killed control measured by ICP-MS (n = 3–4 pooled plant samples per group).

(B) PCA ordination of sample distances between root transcriptional profiles of Col-0 and *f6'h1* plants grown for 1 week on avFe or unavFe media inoculated with a live SynCom or heat-killed control. Data are from two pooled experiments (n = 6 samples pooled from 6 plant roots each).

(legend continued on next page)

mechanisms. This would indicate that coumarins have multiple roles in mediating host-microbe interactions. Importantly, our results were obtained using a bacterial culture collection derived from *A. thaliana* roots grown in CAS, in which coumarin status did not affect plant performance or microbiota structure. The observation that taxonomically diverse commensals isolated from an iron-replete soil are capable of rescuing iron-limiting plant growth further suggests the involvement of microbial functions that are ubiquitous across soil types and can be elicited by fraxetin. Future work with commensal culture collections derived from soils with contrasting iron availability are required to directly link plant phenotypes in natural calcareous soils and gnotobiotic systems.

Root-secreted coumarins are inducible under iron starvation and mediate an interaction between the host and commensals that improves host iron nutrition. This genotype-environment interaction strongly suggests that the root microbiota is an integral component of plant edaphic adaptation to growth in iron-limiting soil. Quantitative variation in coumarin production has been demonstrated among *A. thaliana* accessions (Siwinski et al., 2014; Tsai et al., 2018) and was shown to correlate with performance under iron limitation (Tsai et al., 2018). As coumarins are both ubiquitously present and chemically diverse among flowering plants (Bourgand et al., 2014; Krieger et al., 2018; Rajniak et al., 2018), our findings provide an ecological framework for examining the consequences of their evolutionary diversification on microbiota-mediated mineral nutrition of plant hosts.

STAR★METHODS

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at <https://doi.org/10.1016/j.chom.2020.09.006>.

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AUTHOR CONTRIBUTIONS

P.S.-L., H.I., and M.H. conceived the project. C.J.H., M.H., H.I., and P.S.-L. designed the experiments. A.D.R. provided IS. M.H. and C.J.H. performed the root microbiota profiling experiments in natural soils. C.J.H., R.G., and R.G.-O. analyzed the microbiota profiling data. C.J.H. performed SynCom reconstitution experiments, bacterial growth assays, and RNA-seq experiments. RNA-seq data analysis was performed by Y.N., R.G.-O., and C.J.H.; R.G.-O., R.G., and Y.N. provided bioinformatic tools and support. S.K. advised and performed ICP-MS analyses. E.S.S. and M.J.E.E.V. provided scientific advice at an early stage of the project and feedback on the manuscript. P.S.-L. and R.G.-O. supervised the project. C.J.H. and P.S.-L. wrote the paper.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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(C) Heatmap of median-centered Z scores for 2,440 DEGs identified across samples, arranged by *k*-means clustering. Significantly enriched iron homeostasis-related and defense-related GO terms of pertinent clusters are indicated on right of heatmap. GO analysis was performed by comparing the indicated DEG cluster to the whole transcriptome ($p\text{-adj} \leq 0.05$).

(D) Expression of select iron deficiency response marker genes. Data are \log_2 -transformed, normalized counts. Letters in (A) and (D) indicate significant pairwise differences between groups ($p\text{-adj} \leq 0.05$ by Tukey's HSD corrected for multiple comparisons). See also Figures S6 and S7.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Bacterial and Virus Strains		
AtSphere <i>Arabidopsis thaliana</i> -derived culture collection. Full list of strains in this study are in Table S2	Bai et al., 2015	http://www.at-sphere.com/
Biological Samples		
Cologne agricultural soil	Bai et al., 2015	CAS
Italian soil	This study	IS
Chemicals, Peptides, and Recombinant Proteins		
DFS Taq Polymerase	Bioron	CAT#101005
Agar Bacteriological	Difco	CAT#214530
FeEDTA	Sigma-Aldrich	CAT#E6760
FeCl ₃ hexahydrate	Merck	CAT#103943
HEPES buffer	Roth	CAT#6763.1
Scopoletin	Sigma-Aldrich	CAT#S2500
Fraxetin	Sigma-Aldrich	CAT#18224
Tryptic Soy Broth	Sigma-Aldrich	CAT#T8907
Critical Commercial Assays		
FastDNA SPIN kit for Soil	MP Biomedicals	CAT# SKU 116560200
Quant-IT Pico Green dsDNA Assay	ThermoFischer	CAT#P7589
AMPure XP DNA purification beads	Beckman Coulter	CAT#A63881
RNEasy Plant Mini Kit	Qiagen	CAT#74904
Deposited Data		
16S rRNA root profiling data	This paper https://www.ebi.ac.uk/ena	ENA: accession PRJEB38663
Raw and processed data, scripts for analysis and figure generation	This paper	Mendeley Data DOI: 10.17632/tkdn6zbw7k.1
Root RNA-Seq data	This paper https://www.ebi.ac.uk/ena	ENA: accession PRJEB38663
Silva 132 rRNA database, Released Dec. 2017	Quast et al., 2013	RRID: SCR_006423
Experimental Models: Organisms/Strains		
<i>A. thaliana</i> : Col-0 wildtype	NASC	N60000
<i>A. thaliana</i> : <i>f6'h1-1</i> TDNA insertion mutant	NASC	SALK_132418C
<i>A. thaliana</i> : <i>s8h-2</i> TDNA insertion mutant	NASC	SM_3_23443
<i>A. thaliana</i> : <i>cyp82c4-1</i> TDNA insertion mutant	NASC	SALK_001585
<i>A. thaliana</i> : <i>pdr9-2</i> TDNA insertion mutant	NASC	SALK_050885
<i>A. thaliana</i> : <i>irt1-1</i> TDNA insertion mutant	Vert et al., 2002	<i>irt1-1</i>
<i>A. thaliana</i> : <i>fro2 (frd1-1)</i> TDNA insertion mutant	NASC	N3777
<i>A. thaliana</i> : <i>aha2-4</i> TDNA insertion mutant	NASC	SALK_082786
<i>A. thaliana</i> : <i>bts-1</i> TDNA insertion mutant	NASC	SALK_016526
Oligonucleotides		
All primers are found in Table S1		N/A
Software and Algorithms		
R statistical environment	https://www.r-project.org/	V 4.0.1
DADA2	Callahan et al., 2016	V1.12.1

(Continued on next page)

Continued

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Interactive Tree of Life (iTOL)	Letunic and Bork, 2007	https://itol.embl.de/
Vegan, R package	Dixon, 2003	V 2.5-6
RVAideMemoire, R package	Hervé, 2020	V 0.9-77
edgeR, R package	Robinson et al., 2010	V 3.24.3
Fastp	Chen et al., 2018	RRID: SCR_016962
Kallisto	Bray et al., 2016	RRID: SCR_016582
Tximport, R package	Soneson et al., 2015	V 1.16.1
SVA, R package	Leek et al., 2012	V 3.36.0
DESeq2, R package	Love et al., 2014	V 1.28.1
ComplexHeatmap, R package	Gu et al. 2016	V 2.4.2
Goseq, R package	Young et al., 2010	V 1.40.0
Gene Ontology Consortium	Ashburner et al., 2000	RRID: SCR_002811
Ggpubr, R package	Kassambara, 2020	V 0.3.0
FSA, R package	Ogle et al., 2020	V 0.8.30
Rcompanion, R package	Mangiafico, 2020	V 2.3.25
Emmeans, R package	Lenth, 2020	V 1.4.8
BioRender (Graphical Abstract)	BioRender.com	RRID: SCR_018361

RESOURCE AVAILABILITY

Lead Contact

Further information and requests for resources and reagents should be addressed to the Lead Contact, Paul Schulze-Lefert (schlef@mpipz.mpg.de)

Materials Availability

This study did not generate new unique reagents. All bacterial strains and *Arabidopsis* lines used in this study were previously described and are publically available.

Data and Code Availability

Raw sequencing data of 16S rRNA profiling experiments and RNA-Seq transcriptomic data have been deposited in the European Nucleotide Archive ENA: PRJEB38663. Datasets and scripts necessary to reproduce analyses and generate figures have been deposited to Mendeley Data: <https://doi.org/10.17632/tkdn6zbw7k.1>. Scripts for RNASeq analysis and heatmap generation are available at https://github.com/YulongNiu/MPIPZ_CJ_RNASeq.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Soils

Cologne agricultural soil (CAS) was obtained from a local site (GPS code : 50.958 N, 6.856 E) that has not been exposed to agriculture for >15 years. Italian soil (IS) was obtained from an organic vineyard in Tebano, Italy (GPS code : 44.292 N, 11.784 E) which has been maintained since 2007 without irrigation or fertilization. Soil was homogenized, sieved, and stored at 4°C until used in experiments.

Plant Model

All *A. thaliana* genotypes used in this study were in the Columbia wild-type (Col-0, N60000) background. See [Key Resources Table](#) for full information on all genotypes. Mutants related to coumarin biosynthesis (*f6'h1-1*, SALK_132418C; *s8h-2*, SM_3_23443; *cyp82c4-1*, SALK_001585) and export (*pdh9-2*, SALK_050885), regulation of the iron starvation response (*bts-1*, SALK_016526), and iron reductive import (*aha2-4*, SALK_082786; *fro2*, also known as *frd1-1* (Robinson et al., 1999);, and *irt1-1* (Vert et al., 2002)) were employed in this study and are available from the Arabidopsis Biological Resource Center (ABRC). Each of these genes are expressed in roots.

Bacterial Strains

All bacterial strains used in this study were previously described (Bai et al., 2015) and are summarized in [Table S2](#). Species phylogenetic trees were generated with iTOL version 5.5 (Letunic and Bork, 2007) from previous whole genome taxonomic classification

(Bai et al., 2015). Each of these strains was originally isolated from *A. thaliana* roots grown in CAS soil. Strains were stored in 20% glycerol stocks and cultured on 50% tryptic soy agar (TSA) plates and 50% tryptic soy broth (TSB) at 25°C.

Plant Growth Conditions

Seeds were surface-sterilized with 70% ethanol for 15 min under agitation, followed by two washes with 70% ethanol, one with 100% ethanol, and three with sterile distilled water. Sterilized seeds were stratified at 4°C in the dark for 2–3 days either imbibed in water (for soil experiments) or on agar media plates (for agar-media experiments) before transfer to growth conditions.

Soil

Surface-sterilized, stratified seeds were germinated in 7x7 cm square pots filled with CAS or IS. Pots were watered from the top with non-sterile distilled water every 2 days. Plants were grown in the greenhouse under long-day conditions (16hrs light, 8 h dark). Pots were distributed at random within trays. and shuffled periodically to minimize edge and location effects.

Agar-media

Surface-sterilized seeds were sown on plates containing 1% agar (Agar, granulated, Difco) in 50% Murashige and Skoog (MS) medium with vitamins (2.2 g/L, Duchefa Biochemie) supplemented with 0.5% sucrose. After two days of stratification at 4°C, plates were positioned vertically in a climate chamber (Panasonic, MLR-352) and grown for six days (10 h light, 21 °C; 14 h dark, 19 °C). Uniform seedlings were then transferred to experimental condition plates prepared fresh on the day of seedling transfer.

METHOD DETAILS

Soil Nutrient Content

Soil nutrient analysis was performed by Labor für Boden- und Umweltanalytik (Laboratory for Soil and Environmental Analyses, Switzerland). Plant-available iron was measured as iron mobilized by chelator diethylenetriaminepentaacetic acid (DTPA) extraction as in (Lindsay and Norvell, 1978). Briefly, 10g of air-dried soil was shaken for 2 h with 20 ml of extracting solution (0.005M DTPA, 0.1M triethanolamine, and 0.01M CaCl₂, with a pH of 7.3). The leachate was centrifuged and filtered through a 0.22 micron membrane. Iron content in the leachate was measured by ICP-MS at 1:10 and 1:50 dilutions.

Leaf Chlorophyll Measurement

Chlorophyll extraction and quantification was adapted from (Hiscox and Israelstam, 1979). Samples were prepared from 20–40mg of leaf tissue pooled from ~6 plants per sample and weighed. Samples were either processed immediately or frozen at -80°C until processing. Chlorophyll was extracted by adding 1 ml DMSO per ~30mg tissue and incubating samples at 65°C with shaking for 45–60 min until plant tissue was transparent and chlorophyll completely extracted. Absorbance of tissue-free chlorophyll extract was measured at 652 nm on a spectrophotometer (NanoDrop One, Thermo Scientific). Alternatively, absorbance of 100 µl of samples was measured in 96-well microtiter plate in a microplate reader (Infinite M200 PRO, Tecan). Absorbance_{652nm} cm⁻¹ was converted to total chlorophyll per ml and normalized to input sample tissue mass with the following formula:

$$\text{Chlorophyll}_{\text{Total}} (\text{mg} \cdot \text{g}_{\text{FW}}^{-1}) = \frac{\text{Abs}_{652\text{nm}}}{34.5} * \frac{\text{Volume}_{\text{DMSO}}}{\text{Input}_{\text{FW}}}$$

16S Profiling of the Root Microbiota

For 16S profiling, root samples were harvested and libraries were processed as in (Thiergart et al., 2020). Plant roots were harvested at the early flowering stage, average 37 days after sowing. The whole root system was carefully collected using tweezers, and large soil aggregates and debris were removed by gently washing with sterile water. Roots were transferred to a 15-ml conical with 5 ml PBS buffer (130 mM NaCl, 7 mM Na₂HPO₄, 3 mM NaH₂PO₄, pH 7.0) and washed twice for 20 min on a tube rotator at 120 rpm. Washed roots were briefly dried on filter paper and stored in Lysing Matrix E tubes (MP Biomedicals) at -80°C until processing.

Total root DNA was extracted using the FastDNA SPIN Kit for Soil (MP Biomedicals). Samples were homogenized using the Pre-cellys 24 homogenizer (Bertin Technologies) at 6,200 r.p.m. DNA was eluted in 80 µl of nuclease-free water and quantified using Quant-iT PicoGreen dsDNA Assay (ThermoFisher). Samples were diluted to 3.5 ng µl⁻¹ and used in a two-step PCR amplification protocol. In the first step, the V5–V7 region of the bacterial 16S rRNA gene was amplified in triplicate reactions for each sample with primers 799F and 1192R (Table S1). Amplification was performed in a 25-µl reaction volume containing 2 U DFS-Taq DNA polymerase, 1× incomplete buffer (both Bioron), 2 mM MgCl₂, 0.3% bovine serum albumin, 0.2 mM dNTPs (Life Technologies) and 0.3 µM forward and reverse primers. The same PCR parameters were used for each primer pair (94 °C for 2 min, 94 °C for 30 s, 55 °C for 30 s, 72 °C for 30 s and 72 °C for 10 min for 25 cycles). Single-stranded DNA and proteins were digested by adding 1 µl of Antarctic phosphatase, 1 µl Exonuclease I and 2.44 µl Antarctic Phosphatase buffer (New England Biolabs) to 20 µl of the pooled replicate reactions. Digestion was performed at 37 °C for 30 min, followed by enzyme deactivation at 85 °C for 15 min. Samples were centrifuged for 10 min at 4,000 r.p.m., and 3 µl of supernatant were used for the second PCR step in triplicate reactions to add barcodes and Illumina adapters (primer sequences in Table S1). Reactions were prepared with barcoded primer pairs and performed as above with the number of cycles reduced to ten. PCR quality was controlled by loading 5 µl of each reaction on a 1% agarose gel. The remaining reaction volume was loaded on a 1.5% agarose gel and run at 80 V for 2 h; bands with the correct size of ~500 base pairs were cut out and purified using the QIAquick Gel Extraction Kit (Qiagen). Concentration of the purified DNA was determined, and

30 ng DNA from each of the barcoded amplicons were pooled into one library per experiment, then purified and re-concentrated two-fold with AMPure XP beads (Beckman Coulter). Paired-end Illumina sequencing was performed in-house using the MiSeq sequencer and custom sequencing primers.

Analysis of 16S Profiling Data

ASV Table Generation

Forward and reverse sequencing reads were demultiplexed separately according to the barcode sequence and output in individual fastq files per sample. A denoising pipeline, DADA2 (v1.12.1) (Callahan et al., 2016) was used to obtain the final ASV table. Raw sequencing reads were truncated to 260 bp for the forward reads and 240 bp for the reverse reads and filtered with the following parameters: $maxN=0$, $maxEE=c(2,2)$, $truncQ=2$, $rm.phix=TRUE$. Subsequently, error rates were inferred from filtered reads until convergence or exceeding a maximum consistent number of 20. Sequence variants were then inferred from the trimmed, filtered, and error-corrected sequences and ASVs were obtained by merging the forward and reverse sequences together. Finally, chimeras were identified and removed when an ASV could be mapped to the left- and right-segments from two other, more abundant ASVs. Finally, the ASV table was generated by aggregating the number of reads mapped to each variant.

Alpha and Beta Diversity

Analyses and visualization were performed in the R statistical environment (Version 4.0.1). Analysis was performed on samples with a sequencing depth of at least 2,000 high-quality reads. Alpha and beta diversity were calculated on ASV count tables that were rarefied to 2,000 reads. Alpha diversity (Shannon index) was calculated with using the “diversity” function in *vegan* (Dixon, 2003) (R package version 2.5-6) and differences were tested with ANOVA.

Bray-Curtis dissimilarity index was calculated using the “vegdist” function in *vegan* and used for unconstrained ordination by Principal Coordinate Analysis (PCoA). Constrained PCoA (CPCoA) was performed with the “capscale” function in *vegan*, using the square-root distances of Bray-Curtis dissimilarity index. For ordination constrained on the interaction between genotype and soil type (Figure 1C), the formula used was “distance.matrix ~Host.Genotype*Soil”. Statistical significance of genotype separations was determined using adonis function and pairwise PERMANOVA with 999 permutations using *vegan* and the RVAideMemoire package (Hervé, 2020).

Analysis of deASVs

Analyses and visualization were all performed in the R statistical environment (R Development Core Team, 2010). Analysis was performed on samples with a sequencing depth of at least 2,000 high-quality reads. Relative abundance (RA) was calculated using non-rarefied ASV count data. To calculate deASVs between coumarin pathway mutants and Col-0 WT, data were pooled from three experiments (except *s8h*, which was included in only one experiment), and filtered for ASVs found in at least three samples at a RA > 0.05%. Statistically significant differential enrichment was determined with the *edgeR* package (Robinson et al., 2010) (version 3.24.3) using pair-wise genotype comparisons in a negative binomial generalized log-linear model at an FDR-adjusted p value of 0.05.

Taxonomic classification of ASVs and culture collection strains was performed using the Silva 132 database (released Dec. 2017) (Quast et al., 2013). Hypergeometric enrichment test was performed using the *stats* package in R. Each family was tested for over- or under-representation in the deASVs set by comparing to the list of all detected ASVs. Red asterisks indicate significance at an FDR-adjusted p value of 0.05.

Coumarin Antimicrobial Activity

The antimicrobial activity of scopoletin and fraxetin (Sigma Aldrich) against single bacterial strains was assayed in liquid culture in 50% tryptic soy broth (TSB, 15g/L; Sigma Aldrich). Scopoletin and fraxetin stocks were prepared in sterile DMSO (Sigma Aldrich) and stored at -80°C. Bacterial colonies were picked from TSA plates into liquid TSB and grown for 2–3 days at 25°C with 180 rpm agitation. Liquid cultures were subcultured by diluting 1:100 into fresh TSB and incubated for 1–2 h. In a clear flat-bottom 96-well microtiter plate, 100 μ l of subculture were added to 100 μ l of fresh TSB media supplemented with scopoletin or fraxetin for a final 50 μ M concentration, or equivalent DMSO negative control. The microtiter plate was sealed with a clear adhesive film to prevent evaporation. Growth was monitored kinetically in a microplate reader (Infinite M200 PRO, Tecan) with 30 seconds of shaking followed by measurement of optical density (OD) at 600 nm in four locations per well every 30–60 min for 18–20 h. The OD in each experiment was expressed as the average of triplicate wells per condition. Relative growth (Figure 2D) was calculated by dividing the average final OD measurement of each strain and indicated condition by the average OD in the coumarin-free control.

Gnotobiotic System for Iron Limitation

Iron limitation was achieved with a modified MS medium prepared from stock solutions. Stock solutions were prepared of ethylenediaminetetraacetic acid ferric sodium salt (Fe(III)EDTA, Sigma) in distilled water, and 100mM Fe(III)Cl₃ (Merck) in 10 mM HCl (to prevent precipitation), sterile-filtered and stored at 4°C protected from light. A 2M stock solution of 4-(2-Hydroxyethyl)piperazine-1-ethanesulfonic acid, N-(2-Hydroxyethyl)piperazine-N'-(2-ethanesulfonic acid) (HEPES) buffer (Roth) was prepared, and the pH was adjusted with KOH until a dilution to 10 mM in 50% MS resulted in a pH of 7.4 (approximately pH 8.2 for stock solution) and stored at 4°C.

As a base medium, modified 50% MS media without iron or pH buffer (750 μ M MgSO₄, 625 μ M KH₂PO₄, 10.3 mM NH₄NO₃, 9.4 mM KNO₃, 1.5 mM CaCl₂, 55 nM CoCl₂, 53 nM CuCl₂, 50 μ M H₃BO₃, 2.5 μ M KI, 50 μ M MnCl₂, 520 nM Na₂MoO₄, 15 μ M ZnCl₂, and 9.4mM KCl) was prepared from individual stock solutions. Base media with 1% agar was autoclaved and cooled to 50°C before adding iron

source (final 100 μ M) and HEPES (final 10mM, pH 7.4) with constant stirring. Media were allowed to cool to \sim 45°C, and 45 ml were measured into a conical tube. Live or heat-killed bacteria or SynCom (preparation see below) were added to a final $OD_{600nm} = 0.0001$, corresponding to approximately 10^5 cells/ml. For coumarin complementation experiments, scopoletin and fraxetin were added to a final concentration of 50 μ M, or equivalent DMSO-only control. Media was mixed thoroughly by inverting, poured into 12*12cm square petri dishes, dried with an open lid for 30 min, then allowed to solidify. Seedlings were transferred to experimental plates (six plants per plate, three replicate plates per experiment). For root light-shielding experiments, light-protected plates were placed in completely opaque black boxes with only shoots exposed to light as described in [Silva-Navas et al., 2015](#). Plates were returned to the growth chamber and grown vertically with random shuffling and re-distribution every 2–3 days for uniformity. After two weeks, SFW was measured and chlorophyll and root samples were collected. Plant growth rescue in mono-association assays ([Figure 3D](#)) was expressed as percent growth rescue of the differential between axenic plant growth on avFe and unavFe using the formula:

$$\% \text{ Growth Rescue} = \frac{SFW_{\text{inoculated on unavFe}} - SFW_{\text{axenic on unavFe}}}{SFW_{\text{axenic on avFe}} - SFW_{\text{axenic on unavFe}}} \times 100\%$$

Bacteria Preparation

SynCom

Cultures of 115 SynCom member strains were picked directly from glycerol stocks into 1 ml TSB in 96-well deep-well plates using a 96-well format microplate tip replicator and sealed with breathable plate sealer (AeraSeal, Sigma Aldrich). Cultures were grown at 25°C with 180 RPM agitation for five days and controlled to ensure that a majority of strains grew successfully. Fresh TSB (500 μ l per well) was added and cultures were grown overnight, for 12–18 h to harvest metabolically active cells. Cultures were then centrifuged (4000 g, 20 min), washed once with $MgCl_2$, and resuspended in 300 μ l/well. Cultures were combined, washed, resuspended in $MgCl_2$, and adjusted to $OD_{600} = 0.1$. Heat-killed SynCom was prepared by incubating an aliquot of SynCom suspension at 99°C for 30 min. Heat-killed or Live suspensions were used 1000x to inoculate media (final $OD_{600} = 0.0001$).

Mono-associations

Individual colonies were picked from TSA plates and grown in liquid TSB at 25°C with 150RPM agitation for five days. Strains were subcultured 1:10 in fresh TSB for 2 h, washed twice with sterile 10 mM $MgCl_2$ (Merck), resuspended, and adjusted to $OD_{600} = 0.1$. Strains forming clumps that could not be homogenously resuspended were diluted to a similar degree as other strains in the experiment.

Mineral Composition Analysis

The mineral composition of leaf tissue was determined by inductively coupled plasma mass spectrometry (ICP-MS) from 15–20 mg lyophilized shoot tissue by the CEPLAS Plant Metabolism and Metabolomics Laboratory, University of Cologne, using an Agilent 7700 ICP-MS (Agilent Technologies, Santa Clara, CA, USA) ([Almario et al., 2017](#)).

RNA Extraction and RNA-Seq Analysis

For transcriptomic analysis, 6-day-old *A. thaliana* seedlings were transferred to avFe or unavFe media with live or heat-killed SynCom as above and grown for eight days. Roots from six plants (one plate) were pooled for one replicate, with a total of three replicates per condition in each of two experiments (final n=6). Roots were homogenized with Lysing Matrix E and TissueLyser II (30 beats per second for 2x30 s; Qiagen) and RNA was extracted with the miRNeasy Mini Kit (Qiagen) according to the manufacturer's instructions. RNA quality was determined using a 2100 Bioanalyzer (Agilent Technologies). Preparation of Illumina sequencing libraries was conducted by the Max Planck Genome Center. Sequences were generated using the Illumina HiSeq2500 platform. Approximately 20M paired-end reads with a length of 150 bp were obtained per sample in one experiment, and 8M per sample in the second.

Initial paired-end RNA-Seq reads were pre-processed using *fastp* ([Chen et al., 2018](#)). High quality reads were aligned to *A. thaliana* reference transcriptome (TAIR 10) using *kallisto* ([Bray et al., 2016](#)) with default settings. After removal of low abundant transcripts, 35,886 transcripts were imported using the *tximport* R package ([Soneson et al., 2015](#)). Batch effects were detected and removed using the *SVA* R package ([Leek et al., 2012](#)). Differential gene expression analysis was performed using *DESeq2* ([Love et al., 2014](#)) by comparing the avFe and unavFe treatment with live or heat-killed SynCom in WT and *f6'h1* plants, respectively. DEGs were selected with the threshold $\log_2\text{FoldChange} > \log_2(1.5)$ and an adjusted p-value < 0.05 .

Scaled counts normalized to library size were generated using *DESeq2* ('*rlog*' function) and transformed as median-centered z-score (by transcripts, 'scale' function). Then, z-scores were used to conduct k-means clustering for all transcripts. The cluster number (k=10) was determined by sum of squared error and Akaike information criterion. Transcripts with similar expression patterns were grouped in the same cluster. Differentially expressed transcripts and cluster results were visualized using heatmaps generated with the *ComplexHeatmap* package in R ([Gu et al., 2016](#)). Gene expression in individual plots ([Figures 7D and S7](#)) was plotted using scaled counts data. Gene-set enrichment analyses were performed with the *goseq* package ([Young et al., 2010](#)) with gene ontology annotations from the Gene Ontology Consortium ([Ashburner et al., 2000](#); [The Gene Ontology Consortium, 2019](#)) (September 2019).

QUANTIFICATION AND STATISTICAL ANALYSIS

Details of data visualization, sample size and statistical analysis performed for each dataset can be found in the corresponding figure legend. All boxplots display individual data points, median values, interquartile range, and minimum and maximum values. Statistical analyses were performed in R using the packages and functions indicated above and in figure legends. For statistical analysis of SFW data, nonparametric tests were performed as data were not normally distributed as determined by Shapiro-Wilk test. Kruskal-Wallis one-way analysis of variance between groups was performed with the *stats* package, Wilcoxon signed-rank test to compare SynCom inoculation conditions was performed with *ggpubr* package (Kassambara, 2020), and Dunn's pairwise comparison test was performed with packages *FSA* (Ogle et al., 2020) and *rcompanion* (Mangiafico, 2020). Leaf chlorophyll and elemental content were approximately normally distributed according to Shapiro-Wilk test, and were analysed by ANOVA and Tukey's HSD with correction for multiple comparisons (*emmeans* package (Lenth, 2020)), or by Student's T-test to compare SynCom inoculation conditions with small samples sizes. p values from analyses with multiple comparisons were adjusted using methods indicated in figure legends. Significance was indicated by asterisks (*, **, and ***, indicate $p(\text{adj}) \leq 0.05$, 0.01, and 0.001, respectively) or by significance group ($p \leq 0.05$) by indicated tests. Figures were assembled in Adobe Illustrator. The graphical abstract was created with BioRender.com.