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DOI

[10.1016/j.pbi.2023.102483](https://doi.org/10.1016/j.pbi.2023.102483)

Publication date

2023

Document Version

Final published version

Published in

Current Opinion in Plant Biology

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[Link to publication](#)

Citation for published version (APA):

Wippel, K. (2023). Plant and microbial features governing an endophytic lifestyle. *Current Opinion in Plant Biology*, 76, Article 102483. <https://doi.org/10.1016/j.pbi.2023.102483>

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Plant and microbial features governing an endophytic lifestyle

Kathrin Wippel



Abstract

Beneficial microorganisms colonizing internal plant tissues, the endophytes, support their host through plant growth promotion, pathogen protection, and abiotic stress alleviation. Their efficient application in agriculture requires the understanding of the molecular mechanisms and environmental conditions that facilitate in planta accommodation. Accumulating evidence reveals that commensal microorganisms employ similar colonization strategies as their pathogenic counterparts. Fine-tuning of immune response, motility, and metabolic crosstalk accounts for their differentiation. For a holistic perspective, in planta experiments with microbial collections and comprehensive genome data exploration are crucial. This review describes the most recent findings on factors involved in endophytic colonization processes, focusing on bacteria and fungi, and discusses required methodological approaches to unravel their relevance within a community context.

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Current Opinion in Plant Biology 2023, **76**:102483

This review comes from a themed issue on **Biotic interactions (2024)**

Edited by **Pierre-Marc Delaux** and **Jacqueline Monaghan**

For complete overview of the section, please refer the article collection - **Biotic interactions (2024)**

Available online 6 November 2023

<https://doi.org/10.1016/j.pbi.2023.102483>

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Introduction

Within the plant tissue, bacteria and fungi can colonize inter- or intra-cellularly, and some are able to migrate through the entire plant via the root and shoot vasculature [1,2]. This intimate relationship between host and microbes can be beneficial for both partners (mutualistic), beneficial for the microbial partner without harming the plant (commensal), or detrimental for the plant (pathogenic), although intermediate

associations certainly exist along the continuum from one to another [3]. Although microbial pathogens mostly proliferate within the host tissue, i.e., endophytically, the term endophyte is commonly used for commensal and beneficial organisms that colonize the plant without causing diseases [4]. The interaction with diverse microorganisms is crucial for plant health, development, and nutrition [5]. However, it is assumed that microbial load in host tissue should be limited, because high microbial titer is commonly associated with detrimental microbes [6]. Interestingly, commensal as well as pathogenic bacterial strains seem to be able to live within the *Arabidopsis* leaf apoplast in a multiplication-death equilibrium that avoids excessive proliferation for several weeks [7]. In addition, a harmless endophyte can turn into a pathogen depending on host developmental stage and genotype or changing environmental conditions [4,8,9]. Which type of plant-microbe association is established, is determined to a large extent by the genetic constitution and compatibility of both partners. This article aims to provide an overview of recently identified host and microbial factors and responses that are crucial for microbial endophytic lifestyles, focusing on commensal and beneficial interactions of bacteria and fungi with land plants, mainly in the root compartment, also covering more complex community interactions. Finally, an overview of current methodological approaches to uncover new mechanisms underlying endophytism is provided.

Benefits of hosting endophytic microorganisms

Among the best-described plant-microbe mutualistic interactions is the root nodule symbiosis between legumes and rhizobia under low nitrate conditions, where the bacteria live in specialized root compartments, the nodules, and fix atmospheric nitrogen for the host in return for carbon [10]. Similarly, plants engage with arbuscular mycorrhiza fungi (AMF) that allow the plant to explore a much larger volume of soil and transport phosphate through their hyphae to the intracellular arbuscules in the host root tissue [11]. These are very differentiated interactions for which the required, tightly regulated molecular crosstalk has been intensively studied and described.

Besides the microbial mutualists, commensal bacteria and fungi also exhibit plant growth-promoting (PGP) activities. These can be stimulation via hormones such as cytokinin [12], auxin [13], and gibberellic acid [14], or fertilization by mobilizing nutrients from the surroundings. For example, inaccessible soil organic and inorganic phosphate is mobilized by bacteria by acidifying the rhizosphere through the release of organic acids, or via the activity of phosphatase and phytase enzymes, the latter of which has also been shown for *Penicillium* and *Aspergillus* fungi [15]. It should be noted that the actual ecological relevance for direct plant nutrition by these mechanisms is still under debate [16]. Another example is iron uptake, since iron becomes available through chelating properties of microbial siderophores [17], or through cooperation of plant-secreted coumarins with commensal bacteria [18], although the mechanism for the latter remains to be identified. It was suggested that plants can take up microbe-derived nitrogen and carbon through microbiory by degrading the endophytes, possibly coupled to a rhizophagy cycle where bacteria enter, exit, and re-enter root cells [19]. However, the efficiency of this process seems very low and thus its significance may be weak [20].

Additionally, endophytes may grant protection against pathogens, which can be mediated through niche competition, direct antagonism, or priming of the host immune system. For example, besides the general competition for carbon sources and space [21], pathogens may face iron limitations due to siderophore activity of commensal bacteria. Secreted compounds like the bacterial 2,4-diacetylphloroglucinol (DAPG), pyoverdine, hydrogen cyanide, pyoluteorin, or pyrrolnitrin have direct antimicrobial activity in a contact-independent manner similar to antibiotics [22–24]. Moreover, colonization of the host by beneficial bacteria stimulates the host immune system which can lead to induced systemic resistance, priming the host for subsequent attack by detrimental strains [25].

In the context of abiotic stress, adverse environmental conditions such as drought and salt stress were shown to be alleviated by bacteria via ACC deaminase activity to decrease stress-induced ethylene levels [26,27]. The beneficial fungus *Trichoderma harzianum* conferred salt tolerance in two barley cultivars with different levels of salt sensitivity, by differentially modifying root lipid and polar metabolite profiles in either host [28].

These beneficial activities are often demonstrated for individual strains under laboratory conditions. In natural environments, the presence of other microorganisms, variable abiotic factors, and soil properties may affect these functions. On the other hand, microbial functions may only become evident in a community context and

may be more robust and stable as part of the entire microbiota [29,30].

The role of the host immune system

Both pathogens and endophytes possess microbe-associated molecular patterns (MAMPs), such as the bacterial flagellin peptide flg22 or the fungal cell wall component chitin and are thus recognized by the plant's pattern recognition receptors. However, non-pathogenic microbes are thought to trigger only a weak immune response [31], although beneficial interaction may indeed require a certain immune induction. This was shown for a *Bacillus velezensis* strain which only colonizes efficiently and produces auxin when the *Arabidopsis* reactive oxygen species (ROS) signaling is initiated [32]. Metabolomics analysis of *Brachypodium distachyon* roots revealed an initial defense activation by beneficial *Azospirillum brasilense* before triggering phosphorus deficiency stress alleviation [33]. In a more holistic context, the tryptophan metabolism as part of the immune response was shown to be responsible for controlling fungal load to maintain the beneficial effects of a bacteria-oomycetes-fungi root microbiome [34]. Strikingly, a comprehensive screen of flagellin peptides revealed that commensal bacteria display a large diversity of flg22 epitopes, many of which are immune-evasive and -modulating, linking to host colonization success [35]. To avoid *Arabidopsis* defense activation completely, *Bacillus subtilis* BSn5 produces subtilomycin that binds the bacterial flagellin, thus reducing the flg22-induced immune response and allowing colonization [36]. Similarly, the LysM domain-containing effector Tal6 of the endophytic fungus *Trichoderma atroviride* sequesters chitin-derived GlcNAc and binds to chitin for protection against plant-produced chitinases, thus evading host immune response [37]. A part of the immune system can be suppressed through acidification of the rhizosphere by bacteria secreting gluconic acid (*Pseudomonas capitata* WCS358), or through amino acid synthesis to avoid accumulation of alkaline precursors (*P. simiae* WCS417), although pH-independent mechanisms likely exist [38].

Co-cultivation of *Arabidopsis thaliana* and *Lotus japonicus* with host-adapted commensal communities also resulted in up-regulation of immunity-related transcripts in the roots [39]. Two other studies investigated bacterial commensal communities that exhibited either immune-suppressive or non-suppressive properties in *Arabidopsis*, where suppressive activity usually correlated with better root colonization [40,41]. On the other hand, a consortium of commensal *Pseudomonas* strains triggered a strong immune response in *Arabidopsis* leaves compared to marginal effects of a group of pathogenic strains [6]. With a new approach, comparison of both host and bacterial *in planta* transcriptome profiles of individual commensals inoculated in the *Arabidopsis* leaf apoplast

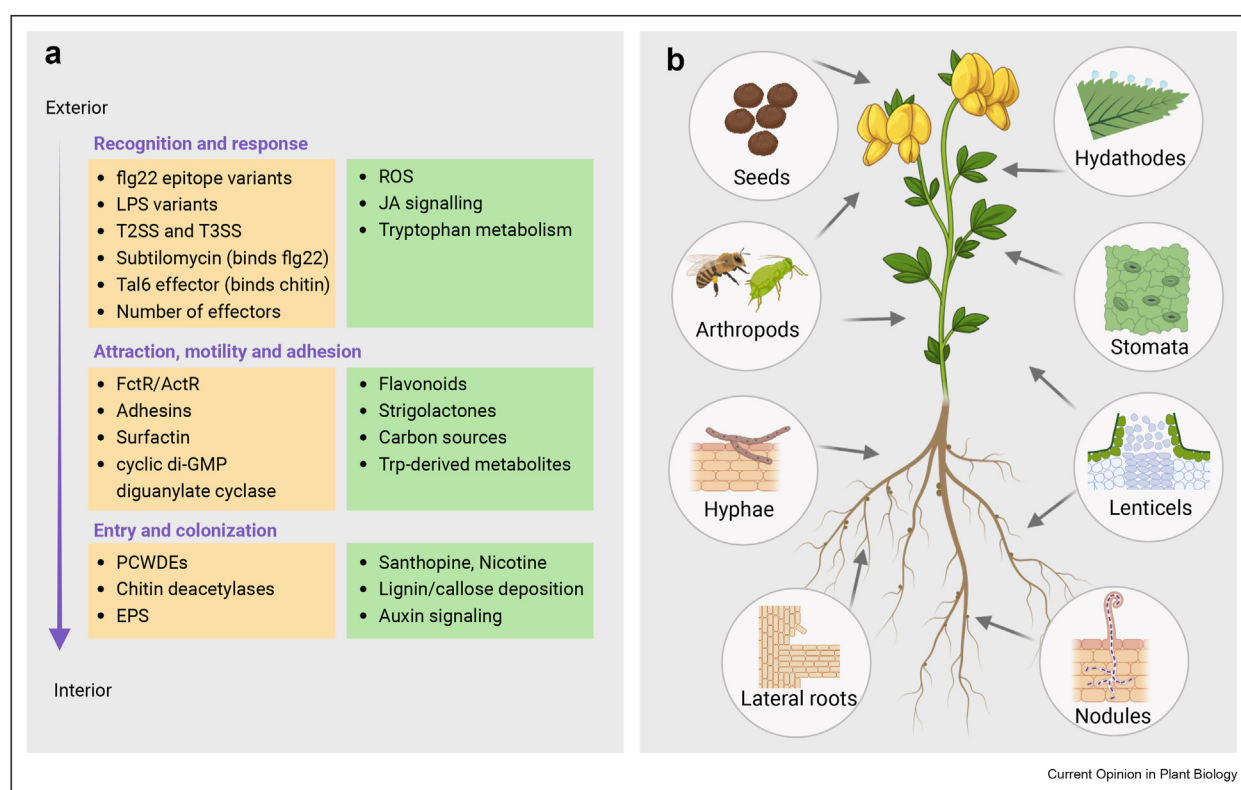
showed that the host responds in a strain-specific manner, for example with different levels of immune activation [42]. Similarly, endophytic accommodation triggered variable expression levels of genes related to energy production and nutrient uptake in these individual strains, which could be a means for the host to orchestrate colonization of diverse microbes by modifying metabolite availability [42].

Other MAMPs like lipopolysaccharides (LPS) - structural components of the bacterial membrane - can act as modulators of symbiosis efficiency through the plant immune system. For example, when the O-antigen component of the LPS was mutated in *Mesorhizobium huakuii*, the mutant strain displayed host accession-specific symbiotic phenotypes, ranging from enhanced to abolished and coupled to host immune defense gene expression [43]. To counteract defense, bacteria deliver effectors into host cells via the bacterial type III secretion system (T3SS). Accordingly, parts or all of the T3SS system is often missing in commensal bacteria,

apart from mutualistic rhizobia and some non-pathogenic *Pseudomonas* and *Xanthomonas* strains [9,44]. Interestingly, the T2SS has been shown to be responsible for the immunosuppressive activity and enhanced colonization of a commensal *Dyella* bacterium on *Arabidopsis* roots [41]. In fungi, endophytic *Fusarium* species tend to possess fewer putative effector genes than pathogenic species [45], making host immune evasion more likely. Finally, plant defense hormones distinguish different microbes. In rice, endophytic *Azoarcus olearius* induced host jasmonate signaling, thought to restrict excessive proliferation, whereas it responded with salicylic acid production to a bacterial pathogen [46].

These findings show that commensals and beneficial microbes can modulate host defense to invade plant tissue (Figure 1a). However, more research on microbial communities in the context of a plant holobiont is needed to decipher how plants integrate signals from welcomed and unwelcomed colonizers [47].

Figure 1



a. Microbial (orange boxes) and plant (green boxes) features discussed in this article as determinants for successful endophytic colonization of commensal or mutualistic organisms. T2SS/T3SS, type II/III secretion system; LPS, lipopolysaccharides; ROS, reactive oxygen species; JA, jasmonate; Fctr/ActR, transcriptional motility regulators; PCWDEs, plant cell wall-degrading enzymes; EPS, exopolysaccharide. **b.** Ports of microbial entry. Schematic depicting different ways and tissues by which microorganisms can enter plant tissue, including seeds (movement of microbes internally to the flower tissue, or externally from the aerial environment), arthropods (vector-mediated entry into flower, leaf, or stem tissue), hydathodes (passive entry), stomata (passive entry or active opening), lenticels (in stems or roots), lateral roots (through tissue cracks at the emergence sites), root nodules (via hitch-hiking along with symbionts), and fungal hyphae, shown representatively for the model plant *Lotus japonicus*.

The role of plant metabolites, microbial motility, and attachment

Plant-derived chemical compounds play an important role in the attraction and maintenance of associated microorganisms, including specialized metabolites that can not only repel pathogens but also control colonization of endophytes [48]. For example, balanced colonization and PGP activity of *Colletotrichum tofieldiae* on *Arabidopsis* were dependent on metabolites of the tryptophane pathway, as shown with corresponding plant mutants [49]. In tobacco, the specialized metabolites santhopine and nicotine are both responsible for the endophytic enrichment of the bacterial genus *Arthrobacter*, which catabolizes these compounds [50].

Indispensable microbial properties for endophytic colonization are motility and the ability for attachment. Flagellum activity, swarming, swimming, adhesion, and biofilm formation are essential for rhizobacteria to move towards the host plant root. For example, the production of *Bacillus velezensis* surfactin is stimulated by root exudates and this lipopeptide is essential for swarming and biofilm formation, which enhances early tomato root colonization and priming of host immunity [51]. In another *B. velezensis* strain, several mutants defective in individual adhesin genes were strongly affected in adhesion to cucumber roots, resulting in impaired colonization [52]. Deleting the gene for the bacterial motility regulator ActR in *Azorhizobium caulinodans* led to loss of swimming ability and impaired biofilm formation, which reduced colonization and nodulation capacities on the host plant *Sesbania rostrata* [53]. Another study showed that cyclic-di-GMP diguanylate cyclase, which is important for biofilm formation, is also required for internal colonization of *Azospirillum baldaniorum* in wheat roots [54].

To move more efficiently through the soil, especially through dry patches where flagellum and ciliary movement are impaired, bacteria may hitchhike on fungal hyphae, the so-called fungal highways [55]. This association requires a give-and-take strategy. For example, in return for transport along its mycelia, *Bacillus subtilis* provides the fungus *Aspergillus nidulans* with the vitamin thiamine, promoting mycelial growth and thus further dispersal of the bacteria [56]. Beneficial AM fungus *Rhizophagus irregularis* attracts and feeds *Rahnella aquatilis* with fungal exudates which allows the bacterium to proliferate and move along the water film around the hyphae [57]. On the other hand, after hitchhiking hyphae of the fungal pathogen *Fusarium oxysporum* to reach the host plant, *Rahnella* can even prevent fungal growth and infection of tomato seedlings through gluconic acid-induced acidification of the rhizosphere, which counteracts virulence-associated alkalization [58]. During peanut-rhizobium root nodule symbiosis, nodulation capacity was enhanced due to the modification of bacterial growth and

chemotaxis in response to root exudates from plants co-cultivated with the beneficial fungus *Phomopsis liquidambaris* [59].

These studies highlight the importance of extending binary host-microbe interaction assays to at least tripartite or more complex interactions that better mimic the natural soil microbiota.

Entry and migration within plant tissue

Natural entry routes for microorganisms can be openings such as stomata, lenticels, or hydathodes, or cracks in the tissue through wounding by soil particles, pathogen attack, lateral root emergence, or abiotic stress (Figure 1b). Non-mutualistic microbes can also enter root nodules as cheaters by hijacking the entry of mutualistic strains. Even though these commensals are not able to fix nitrogen, they can still be beneficial by promoting shoot growth and nodule development, as shown for two *Serratia* strains from lentil nodules [60]. A *Pseudomonas* strain was found to co-colonize healthy, effective nodules of *Lotus burtii*, and to decrease the number of ineffective nodules on *L. japonicus* [61]. Depending on tissue type and plant species, *Enterobacter* sp. SA187 can utilize different entry mechanisms such as passive entry, enzymatic cell wall degradation, or active stomata opening [62]. In general, to proceed into deeper-lying cell layers, plant cell wall-degrading enzymes (PCWDEs) such as cellulases, cutinases, pectinases, and lignin peroxidases are important for colonization by many fungal and bacterial species [8,63,64]. Accordingly, *Arabidopsis* can restrict beneficial *Fusarium* localization to xylem vessels of lateral roots and elongation zone of the primary root compared to pathogenic ones via induced lignin and callose deposition [65], likely preventing fungal overgrowth while keeping beneficial activities. In line with this, this strain possesses fewer PCWDEs than its pathogenic relatives tested in this study [65]. Another interesting example is the beneficial fungus *Epichloë festucae*, which converts its cell wall chitin to chitosan with the help of chitin deacetylases in order to switch from epiphytic to endophytic lifestyle in *Lolium perenne* leaf blades [66]. Furthermore, exopolysaccharides (EPS) are important membrane components of both bacterial and fungal species. EPS outside of fungal cell walls was shown to be crucial for colonization of host plant cells based on a dual function of fungal cell wall modification and plant ROS detoxification with the help of a beta-glucan, which is released through plant hydrolase activity on the EPS [67]. Recently, a plant receptor kinase has been identified that binds these glucans and facilitates AMF symbiosis in *L. japonicus* [68]. Pectinase, cellulase, and ligninolytic activities have also been described for bacterial endophytes of the medicinal plant *Alkanna tinctoria*, and may be linked to the induction of the pharmaceutically important antimicrobials alkannin and shikonin [69].

Microorganisms can be transmitted directly into the plant interior via arthropod vectors. Besides various examples for vector-derived pathogens [70], there has recently been stronger recognition for the significance of insect-mediated exchange of commensal or beneficial microbes between plants. For example, strawberry flowers were shown to function as hubs from which pollinator bees transfer *Streptomyces* strain SP6C4 from one plant to another, so that the beneficial bacterium can exert its antagonism towards fungal and bacterial pathogens on both the plant and the insect [71]. SP6C4 was also able to move from the rhizosphere or from flower tissue to the plant vasculature [71]. Another study found that beneficial sap-sucking mirids acquire and transfer two PGP strains, *Paraburkholderia* and *Enterobacter*, from and between tomato plants, and bacteria migrated from shoots to roots [72].

Once in the plant's interior compartments, microbes may migrate to different organs. The endophytes can subsequently be vertically transferred from the seeds to the next plant generation and influence plant health and microbiota [73]. In rice, seed endophytic communities were shown to be stable over two generations, and shape the root microbiome of rice plants [74]. In maize, protection against *Fusarium* was reported for bacterial seed isolates, alongside of seedling growth promotion [75]. *Sphingomonas melonis* was found to be stably transmitted over three generations of rice seeds, and to confer resistance to detrimental bacterium *Burkholderia planatarii* by interfering with the production of its virulence factor tropolone through anthranilic acid [76].

These studies highlight the versatility of microbial entry modes into host tissue, and that movement between plant organs and generations is possible. Especially regarding stable, field-applicable bioagents, it is important to investigate microbial colonization, heritability, and soil legacy with sophisticated methods.

Methodological approaches for the identification of colonization factors

Characterizing microbial growth, chemotaxis, carbon preferences, and plant beneficial traits of commensals isolated from the inner plant tissues is relevant to understand their individual functions. However, behavior and mechanisms may differ in planta. Culture-independent approaches offer the opportunity to identify host and microbial features involved in host-microbe associations in a more holistic, yet potentially more targeted, manner, better capturing the state of host interaction. With more economic high-throughput sequencing platforms and large public databases being available, comparative omics analyses can provide such comprehensive insights (Table 1). For example, a metagenome analysis of 74 *Micromonospora* strains derived from marine, soil, and plant-associated environments was

used to identify both potential plant growth-promoting and host colonization features [77], while comparative genomics between *Arthrobacter* strains from tobacco roots and soil revealed genes for plant-specialized metabolites that are crucial for plant root colonization [50]. Comparative analysis of 41 fungal genomes of natural *Arabidopsis* isolates showed that PCWDEs are key contributors to endophytic root colonization [63]. Moreover, specific enzyme families are linked to either endophytic (xylanases, lytic polysaccharide monooxygenases) or detrimental (pectate lyases) behavior. Accordingly, over-expression of a pectate lyase in an initially non-detrimental strain resulted in aggressive colonization and negative impact on host fitness [63]. Omics approaches may also be used to predict plant-microbial interactions, as in a recent study that compiled genomes from pathogenic and non-pathogenic *Xanthomonas* strains and employed a machine learning approach to predict plant-associated and pathogenicity factors for diverse strains [44]. Despite the big value of such genomics analyses, they do not consider if genetic elements are indeed expressed under relevant conditions. It is therefore important to validate the results obtained from in silico analyses by in planta experiments and orthogonal assays.

On the other hand, to identify differential host responses according to the colonization patterns and of microbes with different lifestyles, cell type-specific expression of host genes using TRAP (translating ribosome affinity purification) can be analyzed [78]. However, this method requires knowledge of cell type-specific markers and availability of corresponding transgenic plant lines.

The significance of combining experimental data with bioinformatic analyses was also demonstrated in a recent study where bacterial substrate preference determined in the laboratory was integrated with genome-based models of metabolic pathways [79]. This allowed the authors to predict with high confidence which strains compete or cooperate based on the niche overlap, which was transferable to community colonization dynamics on *Arabidopsis* leaves [79]. If an analogous approach proves successful in the root compartment, where additional organic matter for consumption is present in the rhizosphere, or with fungi, remains to be shown.

Another way to investigate colonization niches and dynamics and decipher the corresponding genetic features on the microbial side is to use forward genetic screens with barcoded insertion mutant libraries coupled to high-throughput sequencing, such as TnSeq, BarSeq, and INSeq [80,81]. To track colonization patterns of multiple commensal species of similar taxonomy or mutant strains of the same species in microbiota reconstitution experiments, new methods are being developed, where members of microbial collections are individually barcode-

Table 1**Methodologies for the identification of colonization features.**

| Feature | Method | Required resource/material | Challenges | Limitations | Representative reference | Examples for functional comparisons |
|-----------------|-----------------------------------------------------------------------|-------------------------------------------------------|-------------------------------------------------------------|-----------------------------------------------------------------|--------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Microbial | Comparative genomics of multiple microbes | Annotated microbial whole genome sequences | Available genomes only from cultured microorganisms unknown | Actual in vivo expression/activity unknown | [63] | <ul style="list-style-type: none"> • Beneficial vs. pathogenic • Rhizospheric vs. endophytic |
| Microbial | Forward genetic mutant screen | (Barcoded) mutant library of strain of interest | Genetic tractability of certain taxa | Only one or few strains | [81] | <ul style="list-style-type: none"> • Beneficial vs. pathogenic • Rhizospheric vs. endophytic |
| Microbial | Tracking and quantification of multiple individually barcoded strains | Tagged single (mutant) strains | Genetic tractability of certain taxa | Only one or few focal strains | [82] | <ul style="list-style-type: none"> • Competitiveness (within different communities) |
| Microbial, host | In planta (meta) transcriptomics | Annotated microbial whole genome sequences | Low ratio of microbial to plant RNA, low sensitivity | Transcript assignment to specific organisms not always possible | [42] | <ul style="list-style-type: none"> • Different host species • Host exposure to stress |
| Host | Comparative transcriptomics within one or across host species | Annotated plant genome sequence(s) | Qualitative differences in plant genome annotation | Direct comparison only possible for homologous genes | [39,42] | <ul style="list-style-type: none"> • Beneficial vs. pathogenic • Rhizospheric vs. endophytic |
| Host | Comparative metabolomics within one or across host species | Plant metabolite profile under conditions of interest | Identification of different metabolites of similar mass | Differentiation between plant and microbial metabolites | [33] | <ul style="list-style-type: none"> • Different microbial communities • Beneficial vs. pathogenic • Rhizospheric vs. endophytic • Different microbial communities |

tagged [82]. These technologies, however, are restricted to culturable bacterial and fungal strains.

In the next step, the integration of complex data sets from such studies has the potential to reveal novel networks between hosts and their inhabiting microorganisms, including molecular factors involved in their interaction.

Conclusions

The current state of the art underlines the idea that i) there are shared strategies between endophytes and pathogens for entering host tissue, ii) beneficial microbes also elicit a plant immune response, and iii) the outcome of this response is modulated by additional, endophyte-specific factors. Viewed from the host perspective, the plant is surveilling and managing the behavior of colonizing microorganisms, which sit along a continuum between beneficial and detrimental strains. Omics techniques are being employed to discover further host and microbial factors that are crucial for endophytic host plant colonization by comparing the genetic makeup either of pathogens vs. mutualists, or of microbes inhabiting different host compartments. Currently, there is a need for robust *in planta* microbial metatranscriptomic studies, which are still technologically challenging, and for tracking and quantifying individual microbial strains within communities to explore community dynamics. Host specificity and priority effects will play a significant role in this context. Subsequently, different analyses should be integrated to, e.g., link microbial occurrence or abundance with function. Beyond these approaches to identify genetic factors, spatiotemporal imaging of host tissue colonization and proliferation of microbes will help to understand endophytic microbial lifestyles. Finally, the knowledge gained from these studies needs to be translated to agricultural settings, where the effective usage of beneficial strains and consortia is challenged by application strategies and variations of stability in the field.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

No data was used for the research described in the article.

Acknowledgments

I would like to thank Ana Alexandre for her valuable comments on the manuscript, and two anonymous reviewers for their constructive feedback. Funding was received by German Research Foundation (DFG) Special Priority Programme 2125 DECRyPT; and the Research Priority Area Systems Biology program of the Faculty of Science, University of Amsterdam. Figures created with BioRender.com.

References

Papers of particular interest, published within the period of review, have been highlighted as:

- * of special interest
- ** of outstanding interest

1. Planas-Marquès M, Kressin JP, Kashyap A, Panthee DR, Louws FJ, Coll NS, Valls M: **Four bottlenecks restrict colonization and invasion by the pathogen *Ralstonia solanacearum* in resistant tomato.** *J Exp Bot* 2020, **71**: 2157–2171.
 2. Raiol-Junior LL, Cifuentes-Arenas JC, De Carvalho EV, Girardi EA, Lopes SA: **Evidence that '*Candidatus liberibacter asiaticus*' moves predominantly toward new tissue growth in Citrus plants.** *Plant Dis* 2021, **105**:34–42.
 3. Stengel A, Drijber RA, Carr E, Eggerja T, Hillman E, Krause T, Reese S, Herr JR: **Rethinking the roles of pathogens and mutualists: exploring the continuum of symbiosis in the context of microbial ecology and evolution.** *Phytobiomes J* 2022, **6**:108–117.
 4. Collinge DB, Jensen B, Jørgensen HJ: **Fungal endophytes in plants and their relationship to plant disease.** *Curr Opin Microbiol* 2022, **69**:102177.
 5. Chialva M, Lanfranco L, Bonfante P: **The plant microbiota: composition, functions, and engineering.** *Curr Opin Biotechnol* 2022, **73**:135–142.
 6. Shalev O, Karasov TL, Lundberg DS, Ashkenazy H, Pramod Na Ayutthaya P, Weigel D: **Commensal *Pseudomonas* strains facilitate protective response against pathogens in the host plant.** *Nat Ecol Evol* 2022, **6**:383–396.
 7. Velásquez AC, Huguet-Tapia JC, He SY: **Shared *in planta* population and transcriptomic features of nonpathogenic members of endophytic phyllosphere microbiota.** *Proc Natl Acad Sci USA* 2022, **119**, e2114460119.
- With a 4-week time-course transcriptional profiling and bacterial quantification of commensal and pathogenic bacterial strains colonizing the *Arabidopsis* phyllosphere, this work establishes that both types of strains remain in a stationary phase-like state that follows an equilibrium of multiplication and death. This is one of few studies examining microbial abundance dynamics and features during long-term cocultivation within the host.
8. Redkar A, Sabale M, Zuccaro A, Di Pietro A: **Determinants of endophytic and pathogenic lifestyle in root colonizing fungi.** *Curr Opin Plant Biol* 2022, **67**:102226.
 9. Brader G, Compani S, Vescio K, Mitter B, Trognitz F, Ma L-J, Sessitsch A: **Ecology and genomic insights into plant-pathogenic and plant-nonpathogenic endophytes.** *Annu Rev Phytopathol* 2017, **55**:61–83.
 10. Roy S, Liu W, Nandety RS, Crook A, Mysore KS, Pislaru CI, Frugoli J, Dickstein R, Uvdardi MK: **Celebrating 20 Years of genetic discoveries in legume nodulation and symbiotic nitrogen fixation.** *Plant Cell* 2020, **32**:15–41.
 11. Paries M, Gutjahr C: **The good, the bad, and the phosphate: regulation of beneficial and detrimental plant–microbe interactions by the plant phosphate status.** *New Phytol* 2023, **239**: 29–46.
 12. Zaheer MS, Ali HH, Iqbal MA, Erinle KO, Javed T, Iqbal J, Hashmi MIU, Mumtaz MZ, Salama EAA, Kalaji HM, et al.: **Cytokinins production by *Azospirillum brasiliense* contributes to increase in growth, yield, antioxidant, and physiological systems of wheat (*Triticum aestivum* L.).** *Front Microbiol* 2022, **13**:886041.
 13. Poveda J, González-Andrés F: **Bacillus as a source of phytohormones for use in agriculture.** *Appl Microbiol Biotechnol* 2021, **105**:8629–8645.
 14. Keswani C, Singh SP, García-Estrada C, Mezaache-Aichour S, Glare TR, Borris R, Rajput VD, Minkina TM, Ortiz A, Sansinenea E: **Biosynthesis and beneficial effects of microbial gibberellins on crops for sustainable agriculture.** *J Appl Microbiol* 2022, **132**:1597–1615.

15. Timofeeva A, Galyamova M, Sedykh S: **Prospects for using phosphate-solubilizing microorganisms as natural fertilizers in agriculture.** *Plants* 2022, **11**:2119.
16. Raymond NS, Gómez-Muñoz B, Van Der Bom FJT, Nybroe O, Jensen LS, Müller-Stöver DS, Oberson A, Richardson AE: **Phosphate-solubilising microorganisms for improved crop productivity: a critical assessment.** *New Phytol* 2021, **229**: 1268–1277.
17. Gao B, Chai X, Huang Y, Wang X, Han Z, Xu X, Wu T, Zhang X, Wang Y: **Siderophore production in *Pseudomonas* sp. strain SP3 enhances iron acquisition in apple rootstock.** *J Appl Microbiol* 2022, **133**:720–732.
18. Harbort CJ, Hashimoto M, Inoue H, Niu Y, Guan R, Rombolà AD, Kopriva S, Voges MJEEE, Sattely ES, Garrido-Oter R, et al.: **Root-secreted coumarins and the microbiota interact to improve iron nutrition in *Arabidopsis*.** *Cell Host Microbe* 2020, **28**:825–837.e6.
19. White J, Kingsley K, Verma S, Kowalski K: **Rhizophagy cycle: an oxidative process in plants for nutrient extraction from symbiotic microbes.** *Microorganisms* 2018, **6**:95.
20. Hill PW, Marsden KA, Jones DL: **How significant to plant N nutrition is the direct consumption of soil microbes by roots?** *New Phytol* 2013, **199**:948–955.
21. Macías-Rodríguez L, Guzmán-Gómez A, García-Juárez P, Contreras-Cornejo HA: ***Trichoderma atroviride* promotes tomato development and alters the root exudation of carbohydrates, which stimulates fungal growth and the biocontrol of the phytopathogen *Phytophthora cinnamomi* in a tripartite interaction system.** *FEMS Microbiol Ecol* 2018, <https://doi.org/10.1093/femsec/fiy137>.
22. Matuszewska M, Maciąg T, Rajewska M, Wierzbicka A, Jafra S: **The carbon source-dependent pattern of antimicrobial activity and gene expression in *Pseudomonas donghuensis* P482.** *Sci Rep* 2021, **11**, 10994.
23. Balthazar C, St-Onge R, Léger G, Lamarre SG, Joly DL, Filion M: **Pyoluteorin and 2,4-diacetylphloroglucinol are major contributors to *Pseudomonas protegens* Pf-5 biocontrol against *Botrytis cinerea* in cannabis.** *Front Microbiol* 2022, **13**: 945498.
24. Getzke F, Hassani MA, Crüsemann M, Malisic M, Zhang P, Ishigaki Y, Böhrringer N, Jiménez Fernández A, Wang L, Ordon J, et al.: **Cofunctioning of bacterial exometabolites drives root microbiota establishment.** *Proc Natl Acad Sci USA* 2023, **120**, e2221508120.
25. Teixeira PJPL, Colaianni NR, Fitzpatrick CR, Dangi JL: **Beyond pathogens: microbiota interactions with the plant immune system.** *Curr Opin Microbiol* 2019, **49**:7–17.
26. Zhang L, Zhang W, Li Q, Cui R, Wang Z, Wang Y, Zhang Y-Z, Ding W, Shen X: **Deciphering the root endosphere microbiome of the desert plant *alhagi sparsifolia* for drought resistance-promoting bacteria.** *Appl Environ Microbiol* 2020, **86**, e02863. 19.
27. Roy Choudhury A, Trivedi P, Choi J, Madhaiyan M, Park J, Choi W, Walitang DI, Sa T: **Inoculation of ACC deaminase-producing endophytic bacteria down-regulates ethylene-induced pathogenesis-related signaling in red pepper (*Capsicum annuum* L.) under salt stress.** *Physiol Plantarum* 2023, **175**, e13909.
28. Gupta S, Smith PMC, Boughton BA, Rupasinghe TWT, Natera SHA, Roessner U: **Inoculation of barley with *Trichoderma harzianum* T-22 modifies lipids and metabolites to improve salt tolerance.** *J Exp Bot* 2021, **72**:7229–7246.
29. Finkel OM, Castrillo G, Herrera Paredes S, Salas González I, Dangi JL: **Understanding and exploiting plant beneficial microbes.** *Curr Opin Plant Biol* 2017, **38**:155–163.
30. Durán P, Thiergart T, Garrido-Oter R, Agler M, Kemen E, Schulze-Lefert P, Hacquard S: **Microbial interkingdom interactions in roots promote *Arabidopsis* survival.** *Cell* 2018, **175**:973–983.e14.
31. Thoms D, Liang Y, Haney CH: **Maintaining symbiotic homeostasis: how do plants engage with beneficial microorganisms while at the same time restricting pathogens?** *Mol Plant-Microbe Interactions* 2021, **34**:462–469.
32. Tzipilevich E, Russ D, Dangi JL, Benfey PN: **Plant immune system activation is necessary for efficient root colonization by auxin-secreting beneficial bacteria.** *Cell Host Microbe* 2021, **29**:1507–1520.e4.
- Employing plant and microbial mutants, chemical complementation, and microscopy, the authors of this study dissect the molecular mechanism how a beneficial *Bacillus* strain induces host ROS production to activate microbial auxin secretion necessary for bacterial survival and root colonization, which in turn ensures its antifungal activity.
33. Schillaci M, Kehelpannala C, Martinez-Seidel F, Smith PMC, Arsova B, Watt M, Roessner U: **The metabolic response of *Brachypodium* roots to the interaction with beneficial bacteria is affected by the plant nutritional status.** *Metabolites* 2021, **11**:358.
34. Wolinska KW, Vannier N, Thiergart T, Pickel B, Gremmen S, Piasecka A, Piślewska-Bednarek M, Nakano RT, Belkhadir Y, Bednarek P, et al.: **Tryptophan metabolism and bacterial commensals prevent fungal dysbiosis in *Arabidopsis* roots.** *Proc Natl Acad Sci USA* 2021, **118**, e2111521118.
35. Colaianni NR, Parys K, Lee H-S, Conway JM, Kim NH, Edelbacher N, Mucyn TS, Madalinski M, Law TF, Jones CD, et al.: **A complex immune response to flagellin epitope variation in commensal communities.** *Cell Host Microbe* 2021, **29**: 635–649.e9.
- The authors screened flg22 peptide sequences of hundreds of *Arabidopsis* root and leaf commensal bacteria and found a huge sequence diversity. These flg22 variants are enriched in plant-associated microbiota and confer evasion the host immune activation by modulating the immune receptor signaling through various individual mechanisms.
36. Deng Y, Chen H, Li C, Xu J, Qi Q, Xu Y, Zhu Y, Zheng J, Peng D, Ruan L, et al.: **Endophyte *Bacillus subtilis* evade plant defense by producing lantibiotic subtilomycin to mask self-produced flagellin.** *Commun Biol* 2019, **2**:368.
37. Romero-Contreras YJ, Ramírez-Valdespino CA, Guzmán-Guzmán P, Macías-Segoviano JL, Villagómez-Castro JC, Olmedo-Monfil V: **Tai6 from *Trichoderma atroviride* is a LysM effector involved in mycoparasitism and plant association.** *Front Microbiol* 2019, **10**:2231.
38. Liu Y, Wilson AJ, Han J, Hui A, O'Sullivan L, Huan T, Haney CH: **Amino acid availability determines plant immune homeostasis in the rhizosphere microbiome.** *mBio* 2023, **14**, e03424. 22.
39. Wippel K, Tao K, Niu Y, Zgadzaj R, Kiel N, Guan R, Dahms E, Zhang P, Jensen DB, Logemann E, et al.: **Host preference and invasiveness of commensal bacteria in the *Lotus* and *Arabidopsis* root microbiota.** *Nat Microbiol* 2021, **6**:1150–1162.
- Using microbiota competition experiments with *Arabidopsis*, *Lotus*, and bacteria isolated from both plant species, this work demonstrates that root-colonizing commensals exhibit host preference. Moreover, adapted strains induce immune-responsive host transcripts upon co-cultivation display competitive advantage in an invasion scenario.
40. Ma K-W, Niu Y, Jia Y, Ordon J, Copeland C, Emonet A, Geldner N, Guan R, Stolze SC, Nakagami H, et al.: **Coordination of microbe–host homeostasis by crosstalk with plant innate immunity.** *Nat Plants* 2021, **7**:814–825.
41. Teixeira PJPL, Colaianni NR, Law TF, Conway JM, Gilbert S, Li H, Salas-González I, Panda D, Del Risco NM, Finkel OM, et al.: **Specific modulation of the root immune system by a community of commensal bacteria.** *Proc Natl Acad Sci USA* 2021, **118**, e2100678118.
42. Nobori T, Cao Y, Entila F, Dahms E, Tsuda Y, Garrido-Oter R, Tsuda K: **Dissecting the cotranscriptome landscape of plants and their microbiota.** *EMBO Rep* 2022, **23**, e55380.
- In this study, co-transcriptomics of host and commensal bacteria in the *Arabidopsis* leaf apoplast identify common and distinct responses between the plant-microbe pairs, suggesting that plants in nature may coordinate microbiota structure via specific regulations for each strain. In addition, varying transcript levels of conserved bacterial genes in planta indicate that relying on genomic data only may not always be sufficient to draw conclusions about microbial colonization abilities.
43. Tang Z, Cai S, Liu Y, Li D, Xie F, Lin H, Chen D-S, Li Y: **An LPS O-antigen synthesis gene in *Mesorhizobium huakuii* plays**

- differentiated roles in root nodule symbiotic compatibility with *Astragalus sinicus*. *Mol Plant-Microbe Interactions* 2023, <https://doi.org/10.1094/MPMI-05-23-0066-R>.**
44. Te Molder D, Poncheewin W, Schaap PJ, Koehorst JJ: **Machine learning approaches to predict the plant-associated phenotype of *Xanthomonas* strains.** *BMC Genom* 2021, **22**:848.
 45. Constantin ME, Fokkens L, De Sain M, Takken FLW, Rep M: **Number of Candidate effector genes in accessory genomes differentiates pathogenic from endophytic *Fusarium oxysporum* strains.** *Front Plant Sci* 2021, **12**:761740.
 46. Chen X, Marszałkowska M: **Reinhold-hurek B: jasmonic acid, not salicylic acid restricts endophytic root colonization of rice.** *Front Plant Sci* 2020, **10**:1758.
 47. Mesny F, Hacquard S, Thomma BP: **Co-evolution within the plant holobiont drives host performance.** *EMBO Rep* 2023, **24**, e57455.
 48. Singh G, Agrawal H, Bednarek P: **Specialized metabolites as versatile tools in shaping plant–microbe associations.** *Mol Plant* 2023, **16**:122–144.
 49. Frerigmann H, Piotrowski M, Lemke R, Bednarek P, Schulze-Lefert P: **A network of phosphate starvation and immune-related signaling and metabolic pathways controls the interaction between *Arabidopsis thaliana* and the beneficial fungus *Colletotrichum tofieldiae*.** *Mol Plant-Microbe Interactions* 2021, **34**:560–570.
 50. Shimasaki T, Masuda S, Garrido-Oter R, Kawasaki T, Aoki Y, Shibata A, Suda W, Shirasu K, Yazaki K, Nakano RT, et al.: **Tobacco root endophytic *arthrobacter* harbors genomic features enabling the catabolism of host-specific plant specialized metabolites.** *mBio* 2021, **12**, e00846. 21.
 51. Hoff G, Arguelles Arias A, Boubis F, Pršić J, Meyer T, Ibrahim HMM, Steels S, Luzuriaga P, Legras A, Franzl L, et al.: **Surfactin stimulated by pectin molecular patterns and root exudates acts as a key driver of the *bacillus*-plant mutualistic interaction.** *mBio* 2021, **12**, e01774. 21.
 52. Huang R, Feng H, Xu Z, Zhang N, Liu Y, Shao J, Shen Q, Zhang R: **Identification of adhesins in plant beneficial rhizobacteria *Bacillus velezensis* SQR9 and their effect on root colonization.** *Mol Plant-Microbe Interactions* 2022, **35**:64–72.
 53. Sun L, Wang D, Yin Z, Zhang C, Bible A, Xie Z: **The FtcR-like protein ActR in *Azorhizobium caulinodans* ORS571 is involved in bacterial motility and symbiosis with the host plant.** *Front Microbiol* 2021, **12**:744268.
 54. Sierra Cacho D, Zamorano Sánchez DS, Xiqui-Vázquez ML, Viruega Góngora VI, Ramírez-Mata A, Baca BE: **CdgC, a cyclic-di-GMP diguanylate cyclase of *Azospirillum balduiniorum* is involved in internalization to wheat roots.** *Front Plant Sci* 2021; 12.
 55. Kohlmeier S, Smits THM, Ford RM, Keel C, Harms H, Wick LY: **Taking the fungal highway: mobilization of pollutant-degrading bacteria by fungi.** *Environ Sci Technol* 2005, **39**: 4640–4646.
 56. Abeysinghe G, Kuchira M, Kudo G, Masuo S, Ninomiya A, Takahashi K, Utada AS, Hagiwara D, Nomura N, Takaya N, et al.: **Fungal mycelia and bacterial thiamine establish a mutualistic growth mechanism.** *Life Sci Alliance* 2020, **3**, e202000878.
 57. Jiang F, Zhang L, Zhou J, George TS, Feng G: **Arbuscular mycorrhizal fungi enhance mineralisation of organic phosphorus by carrying bacteria along their extraradical hyphae.** *New Phytol* 2021, **230**:304–315.
 58. Palmieri D, Vitale S, Lima G, Di Pietro A, Turrà D: **A bacterial endophyte exploits chemotropism of a fungal pathogen for plant colonization.** *Nat Commun* 2020, **11**:5264.
 59. Wang H-W, Ma C-Y, Xu F-J, Lu F, Zhang W, Dai C-C: **Root endophyte-enhanced peanut-rhizobia interaction is associated with regulation of root exudates.** *Microbiol Res* 2021, **250**: 126765.
 60. Debnath S, Chakraborty S, Langthasa M, Choure K, Agnihotri V, Srivastava A, Rai PK, Tilwari A, Maheshwari DK, Pandey P: **Non-rhizobial nodule endophytes improve nodulation, change root exudation pattern and promote the growth of lentil, for prospective application in fallow soil.** *Front Plant Sci* 2023, **14**: 1152875.
 61. Crosbie DB, Mahmoudi M, Radl V, Brachmann A, Schloter M, Kemen E, Marín M: **Microbiome profiling reveals that *Pseudomonas* antagonises parasitic nodule colonisation of cheater rhizobia in *Lotus*.** *New Phytol* 2022, **234**:242–255.
 62. Synek L, Rawat A, L'Haridon F, Weisskopf L, Saad MM, Hirt H: **Multiple strategies of plant colonization by beneficial endophytic *Enterobacter* sp. SA187.** *Environ Microbiol* 2021, **23**: 6223–6240.
 63. Mesny F, Miyauchi S, Thiergart T, Pickel B, Atanasova L, ** Karlsson M, Hüttel B, Barry KW, Haridas S, Chen C, et al.: **Genetic determinants of endophytism in the *Arabidopsis* root mycobiome.** *Nat Commun* 2021, **12**:7227.
- This study couples comparative genome analyses of 41 culturable fungal strains with plant performance data after co-cultivation with individual strains, and thereby identifies fungi-encoded plant cell wall degrading enzyme classes that are specific for either endophytic or pathogenic lifestyles. The authors verify their results with genetic manipulation of a commensal strain into an aggressive colonizer.
64. Gluck-Thaler E, Cerutti A, Perez-Quintero AL, Butchacis J, Roman-Reyna V, Madhavan VN, Shanharaj D, Merfa MV, Pesce C, Jauneau A, et al.: **Repeated gain and loss of a single gene modulates the evolution of vascular plant pathogen lifestyles.** *Sci Adv* 2020, **6**, eabc4516.
 65. Martínez-Soto D, Yu H, Allen KS, Ma L-J: **Differential colonization of the plant vasculature between endophytic versus pathogenic *Fusarium oxysporum* strains.** *Mol Plant-Microbe Interactions* 2023, **36**:4–13.
 66. Noorifar N, Savoian MS, Ram A, Lukito Y, Hassing B, Weikert TW, Moerschbacher BM, Scott B: **Chitin deacetylases are required for *Epichloë festucae* endophytic cell wall remodeling during establishment of a mutualistic symbiotic interaction with *Lolium perenne*.** *Mol Plant-Microbe Interactions* 2021, **34**:1181–1192.
 67. Chandrasekar B, Wanke A, Wawra S, Saake P, Mahdi L, Charura N, Neidert M, Poschmann G, Malisic M, Thiele M, et al.: **Fungi hijack a ubiquitous plant apoplastic endoglucanase to release a ROS scavenging β-glucan decasaccharide to subvert immune responses.** *Plant Cell* 2022, **34**:2765–2784.
 68. Kelly S, Hansen SB, Rübsam H, Saake P, Pedersen EB, Gysel K, Madland E, Wu S, Wawra S, Reid D, et al.: **A glycan receptor kinase facilitates intracellular accommodation of arbuscular mycorrhiza and symbiotic rhizobia in the legume *Lotus japonicus*.** *PLoS Biol* 2023, **21**, e3002127.
 69. Rat A, Naranjo HD, Krigas N, Grigoriadou K, Maloupa E, Alonso AV, Schneider C, Papageorgiou VP, Assimopoulou AN, Tsafantakis N, et al.: **Endophytic bacteria from the roots of the medicinal plant *Alkanet tinctoria* tausch (boraginaceae): exploration of plant growth promoting properties and potential role in the production of plant secondary metabolites.** *Front Microbiol* 2021, **12**:633488.
 70. Eigenbrode SD, Bosque-Pérez NA, Davis TS: **Insect-borne plant pathogens and their vectors: ecology, evolution, and complex interactions.** *Annu Rev Entomol* 2018, **63**:169–191.
 71. Kim D-R, Cho G, Jeon C-W, Weller DM, Thomashow LS, Paulitz TC, Kwak Y-S: **A mutualistic interaction between *Streptomyces* bacteria, strawberry plants and pollinating bees.** *Nat Commun* 2019, **10**:4802.
 72. Galambos N, Companat S, Wäckers F, Sessitsch A, Anfora G, Mazzoni V, Pertot I, Perazzoli M: **Beneficial insects deliver plant growth-promoting bacterial endophytes between tomato plants.** *Microorganisms* 2021, **9**:1294.
 73. Nelson EB: **The seed microbiome: origins, interactions, and impacts.** *Plant Soil* 2018, **422**:7–34.
 74. Zhang X, Ma Y-N, Wang X, Liao K, He S, Zhao X, Guo H, Zhao D, Wei H-L: **Dynamics of rice microbiomes reveal core vertically transmitted seed endophytes.** *Microbiome* 2022, **10**:216.
 75. Pal G, Kumar K, Verma A, Verma SK: **Seed inhabiting bacterial endophytes of maize promote seedling establishment and**

- provide protection against fungal disease. *Microbiol Res* 2022, **255**:126926.
76. Matsumoto H, Fan X, Wang Y, Kusstatscher P, Duan J, Wu S, Chen S, Qiao K, Wang Y, Ma B, *et al.*: **Bacterial seed endophyte shapes disease resistance in rice.** *Nat Plants* 2021, **7**:60–72.
 77. Riesco R, Ortúzar M, Fernández-Ábalos JM, Trujillo ME: **Deciphering genomes: genetic signatures of plant-associated *Micromonospora*.** *Front Plant Sci* 2022, **13**:872356.
 78. Fröschel C, Komorek J, Attard A, Marselli A, Lopez-Arboleda WA, Le Berre J, Wolf E, Geldner N, Waller F, Korte A, *et al.*: **Plant roots employ cell-layer-specific programs to respond to pathogenic and beneficial microbes.** *Cell Host Microbe* 2021, **29**:299–310.e7.
 79. Schäfer M, Pacheco AR, Künzler R, Bortfeld-Miller M, Field CM, Vayena E, Hatzimanikatis V, Vorholt JA: **Metabolic interaction models recapitulate leaf microbiota ecology.** *Science* 2023, **381**, eadf5121.
 80. Wetmore KM, Price MN, Waters RJ, Lamson JS, He J, Hoover CA, Blow MJ, Bristow J, Butland G, Arkin AP, *et al.*: **Rapid quantification of mutant fitness in diverse bacteria by sequencing randomly bar-coded transposons.** *mBio* 2015, **6**: e00306–e00315.
 81. Wheatley RM, Ford BL, Li L, Aroney STN, Knights HE, Ledermann R, East AK, Ramachandran VK, Poole PS: **Lifestyle adaptations of *Rhizobium* from rhizosphere to symbiosis.** *Proc Natl Acad Sci USA* 2020, **117**:23823–23834.
 82. Ordon J, Thouin J, Nakano RT, Ma K-W, Zhang P, Huettel B, Garrido-Oter R, Schulze-Lefert P: **Simultaneous tracking of near-isogenic bacterial strains in synthetic *Arabidopsis* microbiota by chromosomally-integrated barcodes.** *Preprint on bioRxiv* 2023, <https://doi.org/10.1101/2023.04.20.537712>.