Modes of Action of Microbial Biocontrol in the Phyllosphere

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A fast-growing field of research focuses on microbial biocontrol in the phyllosphere. Phyllosphere microorganisms possess a wide range of adaptation and biocontrol factors, which allow them to adapt to the phyllosphere environment and inhibit the growth of microbial pathogens, thus sustaining plant health. These biocontrol factors can be categorized in direct, microbe–microbe, and indirect, host–microbe, interactions. This review gives an overview of the modes of action of microbial adaptation and biocontrol in the phyllosphere, the genetic basis of the mechanisms, and examples of experiments that can detect these mechanisms in laboratory and field experiments. Detailed insights in such mechanisms are key for the rational design of novel microbial biocontrol strategies and increase crop protection and production. Such novel biocontrol strategies are much needed, as ensuring sufficient and consistent food production for a growing world population, while protecting our environment, is one of the biggest challenges of our time.

Keywords: biocontrol, phyllosphere, plant immune system, induced systemic resistance, antipathogenic mechanisms, plant pathogens, beneficial microbes, probiotics

INTRODUCTION

Pathogens and pests cause between 20% and 30% of global crop yield losses (Savary et al., 2019). To ensure a sufficient and consistent yield we depend on chemical crop protection and growth-promoting products such as pesticides, herbicides, and fertilizers. Many of these chemical products pose a threat to human health and the environment, which fuels a demand for safer products (Nishimoto, 2019). A promising alternative is the use of microbial based products that can protect crops against diseases. Such microbial products are classified under biological control agents, defined as “a natural enemy, antagonist, or other organism, used for pest control” (ISPM 05, International Standards for Phytosanitary Measures). Although biocontrol is a broad term, including eukaryotic biocontrol agents such as yeasts, fungi, beneficial insects, and other non-microbial pests, in this review we focus on bacterial biocontrol agents. We will use the term biocontrol agent defined similarly as probiotics, “live microorganisms which when administered in adequate amounts confer a health benefit on the host” (Hill et al., 2014). We use this interpretation of a biocontrol agent because it does not only focus on antagonizing the pathogen, but also on improving plant health. Moreover, this definition allows to draw parallels between probiotic and biocontrol research. We will focus on the mechanisms of bacterial biocontrol agents targeting microbial pathogens.
The phyllosphere, the above-ground surface of plants, is a complex ecosystem where microorganisms and the host plant interact extensively to create specific, yet dynamic, communities. Microbial communities inhabit both the external surfaces (epiphytes) as the internal spaces (endophytes) and these communities play an important role in protecting the plant against diseases. Pathogens often have an epiphytic phase before entering the plant cell or the apoplast (intercellular space) (Pfeilmeier et al., 2016). In this review, we focus on external leaf applications of biocontrol agents, unless otherwise specified.

In analogy to a successful probiotic micro-organism, a successful biocontrol agent needs both specific adaptations that allow survival in the phyllosphere habitat (adaptation factors), as well as factors that contribute to the health of the host plant, by inhibiting the pathogen (probiotic or biocontrol factors) (Lebeer et al., 2008). To exert their beneficial properties, biocontrol agents need to be adapted to abiotic environmental factors as well as biotic host-specific factors. A general overview of environmental adaptation factors for the phyllosphere can be found in a review by Vorholt (2012). Adaptation factors are often overlooked in biocontrol research. However, low efficacy of biocontrol agents in field studies is often due to a lack of adaptation rather than a lack of biocontrol factors (Zeriouh et al., 2014; Salvatierra-Martinez et al., 2018). Moreover, a successful biocontrol agent needs a variety of adaptation and biocontrol factors to inhibit a pathogen and improve plant health (Köhl et al., 2019). Biocontrol factors can be related to direct or indirect microbial interactions (Figure 1). Direct interactions occur between the pathogen and the biocontrol agent. Indirect interactions are the interactions between the biocontrol agent and the host plant which improve the plant's fitness, like its resistance to the disease. In this review, we will give an overview of direct and indirect biocontrol and adaptation mechanisms relevant for biocontrol in the phyllosphere. Furthermore, we will describe these mechanisms and the genetic basis in detail, and indicate whether these mechanisms have been validated in the field, in vitro or in greenhouse experiments. An overview of biocontrol and adaptation factors discussed in the text is given in Table 1.

THE PHYLLOSPHERE MICROBIAL HABITAT

The phyllosphere is inhabited by a complex and dynamic community. The composition of this community depends on which microbes reach the phyllosphere in addition to abiotic factors such as climate, season and surrounding land use, and biotic factors such as leaf characteristics and host plant species (Maignien et al., 2014; Agler et al., 2016; Laforest-Lapointe et al., 2016; Smets et al., 2016). Microbes arrive on the phyllosphere rather stochastically via the air, soil, rain or insects. However, only selected taxa successfully colonize the phyllosphere (Maignien et al., 2014). Frequently occurring genera in phyllosphere communities are *Methyllobacterium*, *Sphingomonas*, and *Pseudomonas* (Delmote et al., 2009; Vorholt, 2012). These common phyllosphere bacteria possess specific adaptation factors to the phyllosphere. For example, *Methyllobacterium* spp. have adapted to the low-nutrient environment by metabolizing single-carbon compounds such as methanol (Kutschera, 2007). *Sphingomonas* spp. cope with the scarcity of nutrients by being able to metabolize a wide range of carbon sources (Delmote et al., 2009). *Pseudomonas* spp. use flagellar motility to reach more favorable sites (Haefele and Lindow, 1987), synthesize the biosurfactant syringafactin to increase the water availability on leaf surfaces (Hernandez and Lindow, 2019), and use effectors to leak water from the cells into the apoplast (Xin et al., 2016).

DIRECT INTERACTIONS

**Antibiotic Metabolites and Binary Inhibitory Interactions**

A key first step in the identification of novel biocontrol agents is the screening of antagonistic activities. Such screenings are increasingly applied at a larger scale. For example, Helfrich et al. (2018) recently screened more than 200 leaf isolates from *Arabidopsis thaliana* for binary inhibitory interactions, novel antagonistic strains and interesting metabolites. Most of these strains (88%) engaged in such inhibitory interactions. The orders *Bacillales* and *Pseudomonadales* were especially strong inhibitors, making up only 8% of the tested isolates but engaging in over 60% of the observed inhibitions. Most of the inhibitions also took place between distinct phylogenetic groups rather than within the same family or genus. Genome analysis using the antiSMASH tool (Blin et al., 2019) revealed that many of the inhibitory strains contained more biosynthetic gene clusters than average. These clusters can encode for metabolites with inhibitory effects. The top inhibitor of the collection, *Brevibacillus* sp. Leaf182, was shown to produce several non-ribosomal peptides with antimicrobial activity, such as marthiampetide A (an anti-infective and cytotoxic polythiazole cyclopeptide previously isolated from deep-sea *Marinactinospora thermotolerans*), streptocidin D (a cyclic decapetide antibiotic from the tyrocidine family, named after tyrothricin, the first commercially available antibiotic containing tyrocidine and gramicidin), and an unusual lysophospholipid (a bioactive molecule that possesses a large polar or charged head and a single hydrophobic carbon chain), which was active against Gram-negative bacteria. Previously, biocontrol activity by a *Brevibacillus brevis* strain against *Botrytis cinerea* had been observed in the phyllosphere of Chinese cabbage (Edwards and Seddon, 2001). This strain produces the antibiotic gramicidin S, another cyclic antibiotic non-ribosomal decapetide and major constituent of tyrothricin. Comparison of biocontrol activity with an antibiotic-negative mutant and pure gramicidin S showed that gramicidin S was the mechanism behind the observed biocontrol.

The *Pseudomonas* genus is frequently found in the phyllosphere in relatively high abundances (Delmote et al., 2009; Maignien et al., 2014). The *Pseudomonas* genus includes several commercialized biocontrol strains, such as *Pseudomonas chlororaphis* MA342 and *Pseudomonas* sp. DSMZ 13134. However, also several plant pathogens belong to this genus, such as the model phyllosphere pathogen *Pseudomonas syringae*.
TABLE 1 | Overview of several known mechanisms by which phyllosphere microbes can inhibit pathogen growth.

| In vitro screening | In silico screening | Compound | Mechanism/specific activity | Identified in | References | BC | A |
|--------------------|--------------------|----------|-----------------------------|---------------|------------|----|----|
| **1.1 Antibiotic metabolites** | | | | | | | |
| Binary inhibitory interactions, purification and identification of compounds in supernatant | Screening for biosynthetic gene clusters using the antiSMASH tool | Antimicrobial secondary metabolites | Various | Various species | Helfrich et al., 2018 | X |
| | srfAC, srfAD | | Lipopeptide, surfactin | Triggers biofilm formation | Bacillus spp. | Chen et al., 2007; Ongena and Jacques, 2008; Dunlap et al., 2013; Kim et al., 2015 | X |
| | fenF, mycABC | | Lipopeptide, iturin | Interferes with lipid layers | Bacillus spp. | Chen et al., 2007; Ongena and Jacques, 2008; Dunlap et al., 2013; Kim et al., 2015 | X |
| | ppsABCDE | | Lipopeptide, fengycin | Interferes with lipid layers | Bacillus spp. | Chen et al., 2007; Ongena and Jacques, 2008; Dunlap et al., 2013; Kim et al., 2015 | X |
| | phz gene cluster, ehp gene cluster | | Phenazine | Interferes with histone acetylation and biofilm formation | Pseudomonas spp., Pantoea spp. | Giddens et al., 2002; Chn-A-Woeng et al., 2003 | X | X |
| | ddaA-I | | Herbicolin I | | Pantoea vagans C9-1 | Kamber et al., 2012 | X |
| **1.2 Hydrolytic enzymes** | | | | | | | |
| Zymogram, or specific colorimetric assays | chiA, chiB, chiC or other genes encoding for glycosyl hydrolases from family 18 or 19 in the CAZy database | Hydrolytic enzymes: e.g., chitinase | Hydrolyses fungal cell wall | Bacillus subtilis | Essghaier et al., 2012 | X |
| | msp1 (p75) | | Bifunctional peptidoglycan hydrolase/chitinase | Inhibits hyphae formation | Lactobacillus casei group species | Allonsius et al., 2019 | X |
| **1.3 Quorum quenching and sensing** | | | | | | | |
| | nis gene cluster (nisin), spa gene cluster (subtilin), luxI and luxR (AHLs) | | Signalling molecules. Some gram-bacteria use bacteriocins (nisin and subtilin) that also have a signalling function | Quorum sensing | Nisin in lactococcus lactis and subtilin in Bacillus subtilis | Kleerebezem, 2004 | X |
| | carAB (degradation signalling molecule of Xylella fastidiosa), aiiA (AHL lactonase) | | Enzymes involved in degradation signalling factors | Quorum quenching | Bacillus, Paenibacillus, Microbacterium, Staphylococcus, and Pseudomonas | Newman et al., 2008; Morohoshi et al., 2009; Alymanesh et al., 2016 | X |
| **1.4 Competition for nutrients and space** | | | | | | | |
| Carbon source profiling and calculation of NOI | | | | | | | |

(Continued)
| In vitro screening | In silico screening | Compound | Mechanism/specific activity | Identified in | References | BC | A |
|--------------------|--------------------|----------|-----------------------------|---------------|------------|----|----|
| Selective media with methanol as sole carbon source | mxaF | Conserved enzyme responsible for methanol dehydrogenase | Methylotrophy, increased adaptability in a carbon limited environment | Methylobacterium spp. | Mcdonald and Murrell, 1997 | X | |
| 1.5 Siderophores Plate assay with indicator for detection of siderophores (Chrome azurol S assay) | Screening for siderophore gene clusters, using antiSMASH | Siderophores | Primary function is iron chelation. Siderophores can also have antibacterial properties through the production of ROS and play a role in motility on the phyllosphere | Pseudomonas protegens CS1 | Burbank et al., 2015; Santos Kron et al., 2020 | X | X |
| 2.1 Modulation plant hormone levels | Colorimetric assays | iac gene cluster | Enzymes responsible for the degradation of indole-3-acetic acid (IAA) | Pseudomonas putida 1290 | Leveau and Gerards, 2008 | X? | X |
| | HPLC analysis of extracts of the supernatant | ipdC/iaH or dcc/iaH or iaM/iaA | Enzymes involved in the production of IAA, several pathways possible, described in text | P. agglomerans | Brandl et al., 2001; Duca et al., 2014; de Souza et al., 2019 | X? | X |
| | HPLC analysis of extracts of the supernatant | acoS or accD | Enzymes responsible for lowering ethylene levels | Methylbacterium spp., R. fascians | Chinnadurai et al., 2009; Francis et al., 2016 | X? | X |
| | HPLC analysis of extracts of the supernatant | fas4 or IPT | Enzymes responsible for production cytokinins | Methylbacterium spp., R. fascians | Madhaiyan et al., 2006; Francis et al., 2016; Jorge et al., 2019 | X | X |
| 2.2 Induced systemic response | Transcriptomics of the host plant | Creation of a MAMP database, compare between beneficial and pathogenic microbes | MAMPs that trigger an immune response, that increases protection against pathogens | Sphingomonas melonis fr1 | Ryffel et al., 2016; Vogel et al., 2016 | X | X |
| | Transcriptomics of the host plant | Creation of an effector database, screening for type III secretion system gene clusters | Detectors that trigger an immune response, that increases protection against pathogens | Pseudomonas spp., Parabulholderia sp. | Stringlis et al., 2019; Herpell et al., 2020 | X | X |

The table includes (i) information on in vitro assays to test for the presence of these mechanisms, (ii) known genes involved in these mechanisms (in silico screening), (iii) the compound and (iv) the mechanism resulting in antipathogenic activity, (v) microbes in which the mechanism has been identified, (vi) references and the last two columns indicate whether the mechanism is (vii) a biocontrol factor (BC) and/or (viii) an adaptation factor (A). The screening methods, strains and references are not exhaustive but rather examples, which are also discussed in the text. The table follows the same structure as the review.
Biocontrol Pseudomonas strains have been observed to directly inhibit the growth of a wide variety of pathogens (such as P. syringae and B. cinerea) in lab and in field experiments (Völksch and May, 2001; Romero et al., 2016; Simionato et al., 2017). Biocontrol activity of Pseudomonas spp. is often attributed to the production of phenazines, a group of heterocyclic nitrogen-containing secondary metabolites (Chin-A-Woeng et al., 2003). Biosynthesis in Pseudomonas spp. is encoded by the phz gene cluster. Phenazines inhibit the growth of a variety of fungal pathogens, such as B. cinerea and Fusarium oxysporum [a more detailed overview is given in Chin-A-Woeng et al. (2003)]. The antifungal mode of action of phenazines is multifaceted. For example, Chen et al. (2018) demonstrated that phenazines inhibit mycelial growth of Fusarium graminearum by interference with fungal histone acetylation, and are involved in the formation of a bacterial biofilm on the hyphae, further decreasing pathogenicity. Biofilm formation on fungal hyphae is a widespread trait in soil bacteria (Guennoc et al., 2018). More studies are needed to determine how frequent this occurs in the phyllosphere. Next to phenazines, many other potential biocontrol metabolites have been identified in Pseudomonas spp. such as siderophores (see section “Siderophores”), 4-hydroxy-2-alkylquinolines (Yasmin et al., 2017), volatile compounds such as cyanide and other volatile organic compounds (Bailly and Weisskopf, 2017), and cyclic lipopeptides (non-ribosomal peptides) and rhamnolipids (glycolipids synthesized in a three-step biosynthetic pathway including rhlABC enzymes) (Nielsen et al., 2006; Strano et al., 2017; Yasmin et al., 2017). Rhamnolipids are effective against zoosporic root-infecting pathogens such as Phytophthora and Phytophthora spp. Furthermore, spraying purified rhamnolipids on leaves of Arabidopsis triggers an immune response in the host characterized by the accumulation of signaling molecules and defense genes (Sanchez et al., 2012) (this is an indirect biocontrol mechanism and is further discussed in the section “Plant Hormones”). Interestingly, cyclic lipopeptides and rhamnolipids are also biosurfactants. Biosurfactants generally improve surface motility, biofilm formation and colonization of plant surfaces. Therefore, these adaptation factors could play an important role in the effectiveness of Pseudomonas biocontrol agents. Although, to our knowledge, the importance of rhamnolipids in adaptation, has not yet been investigated in the phyllosphere. Recently, Santos Kron et al. (2020) investigated the role of three antibiotic compounds in the antagonism by Pseudomonas orientalis F9 via experiments with mutants deficient in the production of the siderophore pyoverdine (also see section “Siderophores”), safracin (a tetahydroisoquinoline alkaloid) and phenazine. In vitro double-layer assays showed antagonism against Erwinia amylovora and three P. syringae pathovars by the parental strain P. orientalis F9 and surprisingly also by the pyoverdine and phenazine deficient mutants. Only the mutant deficient in safracin production did not inhibit the pathogens. This
indicates that safracin rather than pyoverdine and phenazine was causing the in vitro antagonism. In contrast, all mutants were able to inhibit the pathogen Pythium ultimum, in vivo in a soil microcosm and E. amylovora, in vitro in a detached blossom assay. These unexpected results indicate that the biocontrol mechanism of P. orientalis is versatile and that other mechanisms could play a role in the observed biocontrol. Recently, Bernal et al. (2017) described the use of a type VI secretion system for the secretion of the RhS effector Tke2 in Pseudomonas putida. The secretion of this effector was shown to be responsible for inhibiting P. syringae, Xanthomonas campestris, Pectobacterium carotovorum, and Agrobacterium tumefaciens in vitro, as well as reducing colonization of X. campestris and reducing disease symptoms on Nicotiana benthamiana leaves. Furthermore, Chen et al. (2016) described the secretion of the antimicrobial siderophore pyoverdine by a type VI secretion system, which inhibits Xanthomonas oryzae pv. oryzae (see also section “Siderophores”). Many Pseudomonas spp., both pathogenic as non-pathogenic, as well as other Gram-negative phyllosphere bacteria, harbor type VI secretion systems, suggesting that these are an adaptation factor (Bernal et al., 2018).

Less frequent inhabitants of the phyllosphere, but often used in commercial biocontrol products, are Bacillus spp. (Ongena and Jacques, 2008). Bacilli isolated from the phyllosphere often engage in inhibitory interactions with other microbial competitors (Helfrich et al., 2018) and their ability to form resistant endospores facilitates their formulation and shelf life (Ongena and Jacques, 2008). Bacillus subtilis and Bacillus amyloliquefaciens are the two most described biocontrol agents in this genus thus far. B. subtilis strains inhibit a wide range of pathogens, both fungal and bacterial, such as F. graminearum (Wang et al., 2007), B. cinerea (On et al., 2015), Alternaria spp. (Ali et al., 2016), X. campestris, and P. carotovorum (Zeriouh et al., 2011). The antipathogenic activity of bacilli has mainly been attributed to the synthesis of non-ribosomal peptides and polyketides (Ongena and Jacques, 2008; Chen et al., 2009). The three classes of non-ribosomal lipopeptides, surfactin, iturin, and fengycin, often act in a synergistic manner. Interestingly, surfactins produced by B. subtilis do not appear to play a role in the antipathogenic activity in vitro, whereas they are necessary for biocontrol in planta (Zeriouh et al., 2014). Surfactins trigger biofilm formation, allowing B. subtilis to successfully colonize the phyllosphere in sufficient numbers and to manage the release of antimicrobial compounds. Therefore, surfactins are rather adaptation factors than biocontrol factors. Wei et al. (2016) confirmed that B. subtilis QST 713, which is used in commercial products, was able to colonize the leaf surface in sufficient numbers. However, despite successful colonization of the phyllosphere, difficulty to colonize new leaves (i.e., dispersal), limited the biocontrol potential of this product. Fengycins and iturins are mostly active against filamentous fungi, by interfering with the lipid layers and altering cell membrane structures (Ongena et al., 2007), but also against the Gram-negative pathogens X. campestris pv. cucurbitae and P. carotovorum subsp. carotovorum (Zeriouh et al., 2011). B. amyloliquefaciens strains have been proven to be successful biocontrol agents in the field for a wide range of pathogens, such as Sclerotinia sclerotiorum causing canola stem rot (Fernando et al., 2007), and fusarium head blight on durum wheat (Schisler et al., 2002). Salvatierra-Martinez et al. (2018) described biocontrol activity of two B. amyloliquefaciens strains against B. cinerea on tomato plants. These two strains had similar antagonistic effect in vitro, while strain BBC047 showed better results in planta. BBC047 was also able to produce a robust biofilm and maintain higher population density over time on the plants. Therefore, it is assumed that adaptation factors explain why strain BBC047 is a more effective biocontrol agent. The genomes of biocontrol B. amyloliquefaciens strains contain several gene clusters encoding for the three lipopeptides surfactin, iturin and fengycin, and polyketide compounds, such as bacillaene, macrolactin and difficidin (Chen et al., 2007; Dunlap et al., 2013; Kim et al., 2015). A clear overview of the secondary metabolite synthetase gene clusters in the genome of B. amyloliquefaciens AS 43.3 is given in Dunlap et al. (2013). Chen et al. (2009) demonstrated that in the mix of these antimicrobial metabolites, the polyketide difficidin and the dipeptide bacilysin, are most important for biocontrol against E. amylovora on apple blossoms. This was proven in planta by applying three mutants of the commercial strain FZB42 on detached apple blossoms, one mutant deficient in production of difficidin, the second unable to synthesize non-ribosomal lipopeptides and polyketides, and a third double mutant deficient in polyketide and bacilysin synthesis. Similarly, Wu et al. (2015) also show the role of difficidin and bacilysin from strain FZB42 in the antagonistic mechanism against X. oryzae pv. oryzae and X. oryzae pv. oryzicola. Moreover, microscopic techniques revealed that difficidin and bacilysin cause changes in the cell wall of Xanthomonas spp.

The genus Pantoea contains several plant pathogens, as well as biocontrol agents effective against a range of pathogens such as B. cinerea, X. campestris, and, the most extensively studied, E. amylovora [as reviewed by Walterson and Stavrinides (2015)]. Several antibiotics, such as pantocins (Smits et al., 2019), herbicolins (Kamber et al., 2012), and phenazines (Giddens et al., 2003), have been identified to play a role in the inhibition of E. amylovora. Stockwell et al. (2002) compared biological control of E. amylovora in field conditions by Pantoea agglomerans (syn. Erwinia herbicola) Eh252, known to produce only one antibiotic, and by its near-isogenic derivative, strain 10:12. Strain 10:12 is deficient in the production of mccEh252, involved in the synthesis of microcin C7. Strain Eh252 reduced the incidence of fire blight more effectively than 10:12. However, the mutant strain still protected the plants more effectively than a mock treatment, indicating that other mechanisms also contribute to biocontrol. The antibiotic herbicolin I was identified and characterized in Pantoea vagans C9-1 via the construction of a mutant library (Kamber et al., 2012). The herbicolin I biosynthetic gene cluster responsible dda-A1 is located on the plasmid pPag2. Remarkably, this cluster was not detected in many other biocontrol strains. Using a similar approach, another antibiotic gene cluster, PNP-1 was identified in Pantoea ananatis BFT175, also effective against E. amylovora (Walterson et al., 2014). The PNP-1 cluster shows similarities to a gene cluster encoding for phenazine in Pseudomonas chloraphis. Previously, the ehp gene cluster encoding for phenazine synthesis had been identified in the
Hydrolytic Enzymes

Production of chitinases, as well as other cell wall degrading enzymes, such as β-1,3-glucanase, is a common defense mechanism of plants (Boller, 1993). Microbes can also produce chitinases, which are an important biocontrol mechanism in the rhizosphere (reviewed by Veliz et al., 2017). Their importance in the rhizosphere indicates the potential of further studying the microbial chitinase activity on the phyllosphere. It has been demonstrated that B. subtilis J9 strain protects strawberry plants against B. cinerea in field conditions and that this strain produces extracellular chitinase and protease (Essghaier et al., 2012).

Recently, we observed that certain lactobacilli can inhibit hyphae formation of fungi in vitro by producing bifunctional enzymes with chitinase/peptidoglycan hydrolyase activity (Allonsius et al., 2019). Lactobacilli are not typical phytophphere inhabitants, and often have a low survival rate (Miller et al., 2019). However, they have been shown to dominate the endosphere of Origanum vulgare plants (Pontonio et al., 2018) and have been correlated negatively with disease symptoms of leaf spot on cucumber plants, presumably caused by P. syringae pv. lachrymans (Luo et al., 2019). Next to the production of hydrolytic enzymes by the biocontrol agents themselves, microbes can induce the production of chitinases in the host plant, a common defense reaction in plants. Inhibition of a pathogen by triggering a defense reaction in the host is further discussed in section “Indirect Interactions.”

Quorum Sensing and Quenching

Quorum sensing systems are systems by which bacteria change their behavior once a certain concentration threshold of signaling molecules is passed. In the phyllosphere, signaling molecules mediate behavior that enables bacteria to survive on the leaf surface, such as biofilm development, adhesion, motility, and production of cell-wall-degrading enzymes. Pathogenic bacteria use quorum sensing to measure their population size and regulate the moment to enter the apoplast or plant cell (Pfeilmeier et al., 2016; Leach et al., 2017). Gram-negative bacteria often use N-acyl-homoserine lactones (AHLs) as signaling molecules, which are synthesized by AHL synthase (luxI) and detected by a transcriptional regulator (luxR). Interestingly, AHL molecules can also trigger a response in the host plant (Delalande et al., 2005; Sieper et al., 2014), which is further discussed in the section “Indirect Interactions.” Gram-positive bacteria do not make use of AHL systems, but typically use small post-translationally processed peptides as signal molecules or diffusible signal factors (see further in this section). A wide variety of small communication peptides exist, and these peptides sometimes have other functions as well. For example, Lactococcus lactis and B. subtilis produce the antibiotic lantipeptides nisin and subtilin, respectively, which are also involved in quorum sensing (Kleerebezem, 2004). Both B. subtilis (Wei et al., 2016) and L. lactis (Trias et al., 2008) can survive in the phyllosphere and even have biocontrol characteristics. However, involvement of the bifunctional peptides nisin and subtilin in the biocontrol activity on the phyllosphere has not been described. Therefore, it would be interesting to investigate their role in the biocontrol mechanism of these bacteria.

Interestingly, non-pathogenic bacteria use the same signaling molecules as pathogens and can thereby contribute to disease development or inhibition, depending on the way of interfering. A shared quorum sensing system using AHL-signal molecules was observed between the symbiotic bacteria P. agglomerans, Erwinia toletana and Pseudomonas savastanoi pv. savastanoi, the causative agent for knot disease in olive trees (Hosni et al., 2011). The symbionts, or in this case opportunistic pathogens, benefit from the niche created by disease development by the pathogen and thus participate in its communication system. By cooperating with the pathogen, E. toletana and P. agglomerans aggravated the infection in the olive trees in this study (Hosni et al., 2011). In contrast, other P. agglomerans strains showed biocontrol activity against the pathogen P. syringae pv. tomato in tomato plants (Morella et al., 2019), but it is at present not known whether quorum sensing could be involved. It remains to be determined whether actual biocontrol agents can have this effect on target and non-target pathogens.

Next to cross-communication by producing the same signaling molecules, bacteria can degrade each other’s signals, also known as quorum quenching. Strains belonging to the genera Bacillus, Paenibacillus, Microbacterium, Staphylococcus, and Pseudomonas are able to rapidly degrade the diffusible signal factor, cis-11-methyl-2-dodecenoic acid. This signal is involved in the regulation of virulence of Xanthomonas spp. and Xylella fastidiosa in a quorum-sensing AHL-independent way (Newman et al., 2008). In the quorum-quenching strains, the genes carAB, involved in synthesis of carbamoylphosphate, a precursor for pyrimidines and arginine, were identified to be required for the rapid degradation of this diffusible signal factor. Bacteria containing the carAB genes could reduce disease incidence and severity of X. campestris pv. campestris in a detached leaf assay with mustard, cabbage and turnip plants.
and of *X. fastidiosa* when co-inoculated into the xylem of grape stems. Furthermore, Wu et al. (2015) showed that difficidin and bacilysin produced by *B. amyloliquefaciens* FZB42 (see section "Antimicrobial Metabolites") can downregulate the expression of several virulence genes in *X. oryzae*, including *rpfF*, involved in the production of a diffusible signal factor.

Morohoshi et al. (2009) screened 109 isolates from the potato phyllosphere for the ability to degrade several short-chain and long-chain AHLs, as Gram-negative pathogens use AHLs as a signaling molecule to regulate their virulence. They screened the isolates *in vitro* by using AHL biosensors, i.e., bacteria that respond to the presence of AHLs by producing a reporter protein. One of the enzymes involved in AHL degradation is AHL-lactonase, encoded by the *aaiA* gene, initially identified in *Bacillus* spp. *Microbacterium testaceum* strains StLB018 and StLB037 tested positive for AHL degradation and decreased disease symptoms in potato tissue caused by *P. carotovorum* subsp. *carotovorum*. In contrast, *M. testaceum* ATCC 15829, lacking AHL-degrading activity, did not decrease disease symptoms, indicating that quorum quenching was the mode of action of biocontrol. Alymanesh et al. (2016) used a similar method to screen isolates from the phyllosphere and rhizosphere from saffron, fig, and pomegranate, for the degradation of the AHL 3-oxo-C6-HSL. They concluded that quorum quenching is a common trait among the isolates tested and is most often observed in *Pseudomonas* spp. These *Pseudomonas* isolates with strong quorum quenching activity also showed biocontrol activity against *P. carotovorum* subsp. *carotovorum in vitro* and on potato tubers.

**Competition for Nutrients and Space**

Phyllosphere bacterial community sizes are limited by low carbon availability on the leaf surface (Mercier and Lindow, 2000). Therefore, carbon competition will likely play an important role in the community structure. Microcosm experiments show that “invaders,” such as introduced biocontrol agents, with a similar metabolism as the resident species are strong competitors in environments with a low resource availability, whereas fast-growing species have an advantage when resource availability is high (Yang et al., 2017).

The dominant carbohydrates available on the leaf surface are sucrose, fructose and glucose. These sugars are specifically altered after epiphytic leaf colonization by *Sphingomonas melonis* or the pathogen *P. syringae pv. tomato*, but only to a minor extent by *Methylobacteria* (Ryffel et al., 2016). Phyllosphere bacteria have developed different strategies to utilize all possible carbon sources available. Methylotrophs, such as *Methylobacteria*, have specialized in the utilization of single carbon compounds, such as methane and methanol. Therefore, they do not rely as much on the available sugars on the phyllosphere (Kutscheria, 2007). Methyllobacteria even modulate the release of methanol, which is released as plant cells expand, by encouraging plant growth via the production of plant hormones (see further, section “Plant Hormones”) (Kutscheria, 2007). The *mxaF* gene, which contains the active site of a methanol oxidation complex, was found to be highly conserved among methylotrophs and is an appropriate probe to screen for methylotrophy (Mcdonald and Murrell, 1997). Methylotrophy is thus an important adaptation factor for some phyllosphere bacteria. However, methylotrophs are not likely to inhibit pathogens by competing for nutrients. Nevertheless, Methyllobacteria can possess other biocontrol mechanisms such as antimicrobial metabolites (Kwak et al., 2014) or indirect mechanisms by triggering plant immunity (see further in section “Plant Immunity”) (Madhaiyan et al., 2006).

Another adaptation strategy is the ability to scavenge for a wide variety of carbon sources. The presence of a high variety of TonB receptors in the phyllosphere proteome has been suggested as an indication that the residing species can metabolize a wide variety of carbon compounds (Delmotte et al., 2009). Indeed, TonB receptors are involved in the transport of carbohydrates, siderophores, and vitamin B12, in Gram-negative bacteria (Schauer and Kutschera, 2013). Blanvillain et al. (2007) noted that bacteria expressing a high variety of TonB receptors, but belonging to various taxonomical lineages, share the ability to metabolize a wide variety of carbohydrates. The overrepresentation of TonB receptors in *Xanthomonas* spp. appears to facilitate their survival in the phyllosphere by making them competitive nutrient scavengers (Blanvillain et al., 2007). Additionally, community proteogenomics of the phyllosphere of *Arabidopsis*, clover, and soybean assigned a high proportion and great variety of TonB receptors to *Sphingomonas* species. This high abundance of TonB receptors is thought to allow *Sphingomonas* spp. to be more successful than other Gram-negative bacteria to withstand the carbon-stressed environment and account for their success on the phyllosphere in terms of their relative abundance (Delmotte et al., 2009). Innerer and et al. (2011) tested 17 *Sphingomonas* strains on the phyllosphere of *A. italicana* for their ability to suppress disease symptoms of the pathogen *P. syringae pv. tomato* DC3000. All seven phyllosphere isolates, and four out of five rhizosphere isolates, protected the plant against developing disease symptoms. On the other hand, four out of five *Sphingomonas* non-plant isolates (isolated from air, dust, or water), did not protect the host plant against *P. syringae* infection. Carbon-source profiling of two protective and two non-protective strains suggested that substrate competition plays a role in the observed antagonistic effect. It would be interesting to verify whether the difference in carbon-source utilization is a result of a higher TonB diversity and whether plant-associated *Sphingomonas* spp. typically have a higher TonB diversity in comparison to other *Sphingomonas* spp. The niche-overlap index (NOI) is a measure that can be used to quantify the similarity in carbon source profile of two microbes (Wilson and Lindow, 1994). Wilson and Lindow (1994) calculated the NOI as the number of carbon sources that both strains utilize as a proportion of the total number of carbon sources utilized by one strain. They demonstrated that the NOI of the epiphytic bacteria *Pseudomonas fluorescens*, *P. agglomerans*, *Stenotrophomonas maltophilia*, and *Methylobacterium organophilum* correlated inversely with their ability to coexist with the pathogen *P. syringae* on the phyllosphere of beans (*Phaseolus vulgaris*). In another study, the NOI of 36 non-pathogenic phyllosphere bacteria were correlated with the ability to suppress disease caused by *P. syringae pv. tomato* (Ji and Wilson, 2002). These studies
confirm the hypothesis made by Lindow (1987) that “antagonism due to competition of one strain with another would increase proportionally to the overlap of their ecological niche.” This hypothesis was formulated based on a field study where ice nucleation-deficient \( P. syringae \) mutants successfully antagonized the \( P. syringae \) wild-type strain in field conditions when the mutant was applied to the plants two days before the wild-type strain (Lindow, 1987). Under such conditions, the mutants could successfully outcompete the wild-type strain and a reduction of the frost injury to the plants was noted. However, the mutants had the advantage of being able to occupy the ecological niche first. Priority effects play an important role in competition between microbes and in the assembly of phyllosphere communities (Maingrien et al., 2014). Therefore, some biocontrol agents are more effective as a preventive measure and less so as a treatment.

Berg and Koskella (2018) tested the antipathogenic properties of both a natural phyllosphere community and a simplified synthetic phyllosphere community (comprising of 12 bacterial strains), against \( P. syringae \) pv. \( tomato \). Both the natural as the synthetic community protected the plant against the pathogen. The authors observed that addition of fertilizer to the soil canceled the observed pathogen protection of the synthetic community, but not of the natural community. Microbial loads on the leaves did not increase significantly due to fertilization. The authors hypothesize that fertilization resulted in an increase in phyllosphere nutrient availability. The synthetic communities were all cultured on KB medium before application on the plants. This is a medium on which \( P. syringae \) also grows well. This might have caused selection for metabolically similar strains, which would increase antagonism due to nutrient competition (cfr. Lindow, 1987). Nutrient competition might therefore play a more prominent role in the synthetic communities than in the more diverse natural communities, where other modes of action could possibly dominate. This hypothesis on nutrient-dependent effects provokes two novel research questions. Firstly, does soil fertilization increase nutrient availability in the phyllosphere and secondly, how does this have an impact on biocontrol in the phyllosphere in field conditions?

**Siderophores**

Apart from carbon sources, iron is often a limiting element in phyllosphere microbial communities. Siderophores are secreted by microorganisms to bind and transport iron into the cell. Siderophore production is essential for the epiphytic fitness of \( P. syringae \) pv. \( glycinea \) 22d/93, a strain with biocontrol activity against the pathogen \( P. syringae \) pv. \( glycinea \) (Wensing et al., 2010). Interestingly, when inoculated in wounded leaves, siderophore production by the commensal had no effect on its own population size nor on the population size of the pathogen. This indicates that iron was not a limiting element in wounded plant cells. Siderophore production is thus not a biocontrol mechanism of importance for \( P. syringae \) pv. \( glycinea \), when the pathogen rapidly penetrates living tissue. However, siderophore production is an important adaptation factor for biocontrol agent \( P. syringae \) pv. \( syringae \) 22d/93, as 10 days post inoculation, the population size of a siderophore-negative mutant was 2 orders of magnitude lower than that of the wild-type. Furthermore, a role for siderophores in the induced systemic resistance (ISR) (see section “Induced Systemic Resistance”) has been reported in several systems (Bakker et al., 2007). It is not excluded that the wounding in the experiment by Wensing et al. (2010) triggered ISR, via host jasmonic acid (JA) and ethylene mediated pathways (see section “Plant Hormones”). The wounding switched off the necessity for an additional siderophore triggered ISR and the strain did not exert any biocontrol activity in the wounded plants.

Siderophores can have alternative functions in addition to iron scavenging, such as non-iron metal transport, sequestration of toxic metals, signaling, protection from oxidative stress, and antibiotic activity. The latter occurs by attaching a bactericidal ‘warhead’ on a siderophore which is then taken up by the antagonized bacterium (Kramer et al., 2019). The siderophore enanto-pyochelin, produced by \( Pseudomonas protegens \) CS1, isolated from the lemon tree phyllosphere, showed antagonistic activity in vitro and in the phyllosphere of lemon plants against the pathogen \( Xanthomonas citri \) subsp. \( citri \) (Michavila et al., 2017). Additions of iron and ascorbic acid indicated that not competition for iron but oxidative stress, induced by the formation of reactive oxygen species (ROS) from pyochelin, was the mechanism of action for the observed antimicrobial activity. Indeed, ascorbic acid was able to counteract the antimicrobial activity of ROS while addition of iron had almost no effect. In contrast, experiments with \( P. orientalis \) F9 and a mutant deficient in the production of siderophore pyoverdine (also see section “Antibiotic Metabolites”) showed that the mutant was still able to antagonize \( E. amylovora \) and three \( P. syringae \) pathovars in vitro, as well as \( E. amylovora \) on a detached flower assay and \( P. ultimum \) in a soil microcosm assay (Santos Kron et al., 2020). This indicates that pyoverdine did not play a role in the biocontrol mechanism by \( P. orientalis \) F9. Another function of siderophores on the phyllosphere was demonstrated by Ruiz et al. (2015). The siderophores pyoverdine and enanto-pyochelin, synthetized by \( P. protegens \), were responsible for its resistance against the mycotoxin fusaric acid. Fusaric acid is produced by pathogenic fungi of the \( Fusarium \) genus and is toxic to plants and bacteria through mechanisms that are not yet fully understood. Finally, Burbank et al. (2015) showed that mutations in the iucA and iutA genes, responsible for siderophage and receptor biosynthesis respectively, results in a loss of surface motility of the xylem-dwelling pathogen \( Pantoea stewartia \), and reduced virulence in sweet corn. This indicates that siderophores also play a role in adaptation by mediating motility. However, this mechanism has not been described yet as an adaptation strategy for phyllosphere biocontrol agents.

**INDIRECT INTERACTIONS**

Next to direct interactions, biocontrol agents can inhibit pathogens indirectly, by modulating the plant’s immune system or hormone levels (Figure 1). Microbe-plant interactions that protect the plant against pathogen infection are discussed here as indirect interactions.
Plants have evolved a complex immune system to prevent infection by recognizing potential intruders and responding with an appropriate defense response. Reversely, pathogens evolve continuously to evade recognition or to interfere with the defense response. This action and counteraction are described by Jones and Dangl (2006) in the “zigzag model.” A schematic representation of the host immune system as well as mechanisms by which biocontrol agents can interact with it is given in Figure 2. The host plant recognizes microbe-associated molecular patterns (MAMPs) by specific pattern-recognition receptors (PRRs), which leads to pattern-triggered immunity. One of the best studied MAMPs is flagellin, more specifically the epitope flg22, which is recognized by the PRR FLS2. Other MAMPs are lipopolysaccharides from Gram-negative bacteria and N-acetylglucosamine-containing glycans, such as bacterial peptidoglycan, generally more prominently in Gram-positive bacteria, fungal chitin, or rhizobacterial nodulation factors. Also volatile compounds emitted by beneficial bacteria such as Bacillus and Pseudomonas spp. can trigger the plant’s immune system, however the receptors remain to be identified (Tyagi et al., 2018). An overview of PRRs, the specific MAMPs that are recognized, and the molecular basis of the following pattern triggered immunity has been reviewed by Saijo et al. (2018). Of importance here, both pathogens and mutualistic microbes are detected through MAMP-PRR interactions and detection generally leads to relatively weak immune responses. Hacquard et al. (2017) argues that the pattern-triggered immunity does not discriminate between a beneficial or pathogenic attack, but mainly functions by restricting the microbial load. The immune response can become stronger when additional virulence factors are present, such as tissue damage or plant hormones modulation (discussed further in this section) (Jones and Dangl, 2006; Hacquard et al., 2017).

Microbes can overcome this first line of defense by modifying MAMPs or by secreting effectors into the cytoplasm of host cells that interfere with the triggered immune signaling. Consequently, plants have evolved additional mechanisms to detect these microbial effectors: effector-triggered immunity (Jones and Dangl, 2006; Figure 2). Gram-negative bacteria use type III secretion systems to deliver effector molecules into the cytoplasm to suppress the immune system. When such a secretion system is inactivated (through mutations in hrp genes, which are required for a functional type III secretion system and elicitation of a hypersensitive response in plants) in pathogens, disease symptoms are reduced (Hanemian et al., 2013). Such mutants are then unable to overcome pattern-triggered immunity and are unable to infect host tissue. These mutants often reside in the apoplast without causing harm and can even protect the host against invasion of the wild-type pathogen (Hanemian et al., 2013). Mutants and wild-type pathogens colonize separate cells/niches when co-inoculated. In some studies, co-inoculation led to protection (Faize et al., 2006), while in others, it was necessary to inoculate the non-virulent mutant prior to the pathogen (Feng et al., 2012). Therefore, it is debatable whether competition for nutrients and space is an important mode of action. However, multiple studies show that inoculation with hrp mutants induces changes in gene expression which lead to a defense response and increased resistance against the pathogen (Faize et al., 2006; Feng et al., 2012; Hanemian et al., 2013).

Type III secretion systems and effectors have mostly been described in pathogens. However, some commensals also interact more actively with the plant’s immune response via effectors. For example, nodulating rhizobacteria secrete effectors using type III and type VI secretion systems to activate nodulation in the host plant (Deakin and Broughton, 2009). Recently, Stringlis et al. (2019) identified type III secretion system gene clusters in beneficial rhizosphere Pseudomonas spp. These gene clusters were highly similar to type III secretion systems in other beneficial bacteria, but distinct compared to phytopathogenic P. syringae. Also on the phyllosphere, type III secretion systems have been identified in the genome of a non-pathogenic Paraburkholderia isolate (Herpell et al.,...
However, more research is needed to determine the presence of type III secretion systems in other beneficial phyllosphere bacteria, to identify effectors associated with pathogens versus biocontrol effectors, and to determine the role of these type III secretion systems in beneficial host–microbe interactions.

A group of molecules often forgotten that have an effect on the host immune system are N-acyl-homoserine lactones (AHLs) (Schenk and Schikora, 2015). AHLs regulate the behavior of Gram-negative pathogens (see section “Quorum Sensing and Quenching”). The host plant could benefit from the ability to respond to or interfere with this quorum sensing signal. Indeed, exposing roots to AHLs or AHL-producing bacteria has been shown to trigger the upregulation of defense genes in the plant and inducing systemic resistance via salicylic acid (SA) signaling (described further) (Schenk and Schikora, 2015). One of the AHL-induced defense reactions of Arabidopsis plants was stomatal closure, a common first defense reaction to limit the entry of pathogens in the apoplast. Furthermore, plants react by degrading the AHLs (Delalande et al., 2005) or transporting the AHLs into the vascular system to remove them from the bacteria that produced them (Sieper et al., 2014). Both actions could be described as a plant’s equivalent to quorum quenching. Detection of AHLs in Arabidopsis plants is mediated through the G-protein coupled receptor encoded by AtGPA1. However, more research is needed to identify AHL receptors in other plants.

Finally, the plant’s immune system can be triggered by the detection of host-derived danger-associated molecular patterns (DAMPs), such as oligogalacturonides which are released from the plant cell wall during pathogen invasion. Detection of DAMPs in combination with pattern or effector triggered immunity, will result in a more severe immune response against the invader (Hacquard et al., 2017). It is unlikely biocontrol agents trigger the host immune system through the release of DAMPs.

As described here, both biocontrol as pathogenic microorganisms interact with the host immune system. Similarly to pathogen and commensal host interactions in humans, the final host response depends on the sum of the interactions with host receptors. Commensal bacteria do not trigger a strong defense reaction in the host as they lack additional virulence factors, such as triggering the formation of DAMPs, modulating plant hormone levels or secreting effectors into the host cells (Lebeer et al., 2010).

Plant Hormones

The recognition of beneficial or pathogenic microbial attacks, as described above, leads to the activation of signaling hormones in the plant, as counterparts of immune modulating cytokines in human and animal cells. Relevant plant hormones include SA, JA, and ethylene, where SA and JA are considered to act antagonistic (Koornneef and Pieterse, 2008; Pieterse et al., 2014). JA and ethylene are usually involved in the defense response against necrotrophic pathogens (feeding on killed host cells), or after wounding, while SA is involved in the defense reaction against biotrophic or hemibiotrophic pathogens (feeding on living tissue) (Glazebrook, 2005). Experiments spraying bacterial produced rhamnolipids on leaves of Arabidopsis (see section “Direct Interactions”) showed that SA plays a central role in rhamnolipid-mediated disease resistance (Sanchez et al., 2012).

A first example on how phyllosphere microbes can directly modulate plant hormone levels, is through the enzyme 1-aminocyclopropane-1-carboxylate (ACC) deaminase that degrades the ethylene precursor ACC. It has been detected in plant-growth promoting rhizosphere bacteria such as Azospirillum, Rhizobium, and Pseudomonas spp. (Gamalero and Glick, 2015; Nascimento et al., 2019), as well as in phyllosphere bacteria, such as several Methyllobacterium spp. (Kwak et al., 2014) and Rhodococcus fascians (Chinnadurai et al., 2009; Francis et al., 2016). 1-Aminocyclopropane-1-carboxylate deaminase activity lowers ethylene levels, reducing the plant’s defense responses and thereby facilitating symbiotic microorganisms. ACC deaminase also results in the promotion of plant growth, since plants become more resilient against environmental stress such as drought, flooding, salt stress or pathogen pressure (Gamalero and Glick, 2015; Nascimento et al., 2018; Saghafi et al., 2020). Direct evidence for a role of ethylene in modulating the community composition of the phyllosphere is given by Bodenhausen et al. (2014), as ethylene-insensitive plant mutants harbored a different phyllosphere community. Moreover, evidence is rising for a direct role of ACC in regulating plant development (Van de Poel and Van Der Straeten, 2014; Vanderstraeten and van Der Straeten, 2017) and defense responses (Tsang et al., 2011).

Levels of phytohormones that are primarily involved in plant growth, such as cytokinins and auxins, can also be modulated by microbes (Leach et al., 2017). Both production and degradation of the auxin indole-3-acetic acid (IAA) have been observed in both plant growth-promoting and pathogenic bacteria (Duca et al., 2014; Nascimento et al., 2019). Degradation of IAA can be advantageous for phyllosphere microbes in two ways. On the one hand, IAA is a good source of carbon and nitrogen (Leveau and Lindsey, 2005; Nascimento et al., 2019). On the other hand, manipulation of IAA levels induces physiological changes in the plant, such as cell wall-loosening and the release of nutrients that benefit the survival or colonization of the microbe (Vanderhoef and Dute, 1981). P. putida 1290 is able to grow on IAA as a sole source of carbon, nitrogen, and energy (Leveau and Lindsey, 2005). This ability of P. putida 1290 is encoded by the iac gene cluster. Homologs of the iac gene cluster have been identified in strains from various genera, such as P. putida GB-1, Marinomonas sp. MWYL1, Burkholderia sp. 383, Sphingomonas wittichii RW1, Rhodococcus sp. RHA, Acinetobacter baumannii ATCC 19606, and Lelliottia sp. (Leveau and Gerards, 2008; Lin et al., 2012). On the other hand, high levels of IAA, produced by plant or bacterium, can play an important role in disease development [e.g., by gall forming pathogens Rhodococcus fascians (Sies et al., 2012) (see further in text), knot development by P. savastanoi (Sorico et al., 1985) or suppression of the host defense system by P. syringae pv. syringae DC3000 (McClerklin et al., 2018)]. Bacterial degradation of IAA has so far not
been directly linked with antipathogenic effects. However, IAA degradation is an important adaptation mechanism of bacteria on the phyllosphere.

Besides degradation, IAA can also be produced by plant-associated bacteria. Production of IAA can occur via several pathways, as reviewed by Duca et al. (2014). The presence of these pathways can be detected by the presence of the following essential genes: ipdC and aldH for the indole-3-pyruvate pathway (encoding the enzymes necessary for the decarboxylation of indole-3-pyruvate and subsequent oxidation, respectively), dce and aldH for the tryptamine pathway (encoding the enzymes necessary for the decarboxylation of tryptophan and subsequent oxidation, respectively), iaaM and iaaH for the indole-3-acetamide pathway (encoding for tryptophan-2-monoxygenase and indole-3-acetamide hydrolase, respectively), and nthA for the indole-3-acetonitrile pathway (encoding for nitrile hydratase α) (de Souza et al., 2019). It is important to note that although the indole-3-acetamide pathway was considered as being exclusive for the excessive IAA production by gall-forming bacteria like P. savastanoi, Erwinia spp., and Agrobacterium transformed plant tissue (Jameson, 2000), these genes are also present in methylotrophic rhizosphere microorganisms (Li et al., 2019). The ipdC gene is of special interest since it was demonstrated that the specific growth conditions in the phyllosphere trigger the expression of the ipdC gene in symbiotic P. agglomerans (syn. E. herbicola) (Brandl et al., 2001). Furthermore, the phyllosphere consists of microenvironments that induce differential expression of the ipdC gene. Expression of the ipdC gene and production of IAA is induced in the rhizosphere symbiont Azospirillum brasilense Sp245 when carbon source availability is limited in batch and fed-batch cultures (Ona et al., 2005). These observations indicate that IAA production, encoded on the ipdC gene, is used by beneficial plant-associated bacteria to adapt to the phyllosphere by improving their growth conditions and availability of carbon sources. However, as with IAA degradation, IAA production is an important adaptation factor and has not yet been linked to biocontrol activity.

Members of the genus Methylobacterium enhance plant growth by producing auxins and cytokinins (Koenig et al., 2002; Kwak et al., 2014; Jorge et al., 2019; Li et al., 2019). Interaction with the host is beneficial for the symbiont’s growth since they metabolize the methanol released as the plant grows (Kutscher, 2007). Methylobacterium derived cytokinins were attributed to drought/saline stress resistance in the host (Jorge et al., 2019). Both the type of cytokinins present and the presence of a miaA gene indicate that methylobacterial cytokinin production is merely via tRNA (Koenig et al., 2002; Kwak et al., 2014; Jorge et al., 2019). Moreover, biocontrol activity has been observed, for example, seed inoculation of groundnut plants with Methylobacterium spp. increased protection against pathogens Aspergillus niger and Sclerotium rolfsii (Madhaiyan et al., 2006). The treatment with Methylobacterium spp. induced an increased activity of enzymes in the host plant that are typically associated with Induced Systemic Resistance (ISR) a state in which the plant’s immune system is triggered in order to become resistant against pathogen infection (see further for the paragraph on Induced Systemic Responses). This indicates that the applied Methylobacterium spp. interacted with the host plant’s defense system resulting in protection against A. niger and S. rolfsii. However, the specific role of microbial production of cytokinins and auxins in the plant’s defense response has not been elucidated in this study.

Bacteria that are known to modulate plant hormone levels are Rhodococcus fascians. Both pathogenic as non-pathogenic R. fascians have the ability to both produce IAA and cytokinins, and decrease ethylene levels (Francis et al., 2016). In pathogenic bacteria, the genes encoding auxin and cytokinin production are plasmid-borne (Jameson, 2000). IAA production is higher in presence of exogenous tryptophan, a precursor of IAA. Interestingly, upon inoculation of the plant with pathogenic or non-pathogenic R. fascians, the metabolism of the host plant changes and more tryptophan is accumulated, possibly stimulating bacterial production of IAA (Francis et al., 2016). On the other hand, the production of cytokinins by R. fascians induce increased auxin production in the plant. The increased auxin levels play an important role in the development of disease symptoms (Stes et al., 2012). The non-pathogenic derivative of this strain lacks the plasmid with virulence genes. The main pathogenicity factor on the plasmid is the production of modified methylated cytokinins, which are not degraded by cytokinin oxidase activity, mimic plant cytokinins, induces increased auxin production in plants and results in the development of disease symptoms (Radhika et al., 2015). Cytokinin and auxin production in pathogenic R. fascians is thus detrimental and contributes to the disease development. Reversely, cytokinin and auxin production in non-pathogenic R. fascians, as well as in other symbionts (e.g., Methylobacteria, described above), is being described as a beneficial trait since it promotes plant growth (Schauer and Kutschera, 2011; Francis et al., 2016; Romero et al., 2016).

In conclusion we can postulate that, through the ability to control the auxin steady state by producing additional auxins on the one hand, and by auxin degradation when excessive auxin production occurs in case an additional pathogen is invading the host on the other hand, a benign symbiont is possibly able to optimize its ecological niche both by improving the host growth and excluding other invaders. The cross talk between auxins and ethylene, as well as the bacterial ACC-degradation might prevent ethylene induced excessive immune and senescence responses to occur. Moreover, it is of general knowledge that cytokinins act as a sink for sugar and other metabolites (Rötsch and Ehneß, 2000), therefore we postulate that it is plausible that the microbial cytokinin production works as a sink for metabolites to the benefit of the symbiont (carbon source) as well as the host by reducing senescence and as a consequence prolonging photosynthetic activities.

**Induced Systemic Responses**

Once a microbe is detected by the plant and its presence is signaled via plant hormones, as described above, a specific immune response is triggered in the plant (Fig 2). Beneficial microbes are able to trigger the plant’s defense system at the
point of recognition and induce immunity against pathogens in the whole plant body (Pieterse et al., 2014). This phenomenon is called induced systemic resistance (ISR). Also pathogens can induce a systemic response, which then protects other parts of the plants, this is called systemic acquired resistance (SAR). The term ISR is usually used when it is triggered by rhizosphere microbes (Pieterse et al., 2014). However, in the next paragraphs, we will describe several examples of ISR by beneficial microorganisms (Pieterse et al., 2014). ISR is usually used when it is triggered by rhizosphere microbes (Pieterse et al., 2014). However, in the next paragraphs, we will describe several examples of ISR by beneficial microorganisms (Pieterse et al., 2014). However, in the next paragraphs, we will describe several examples of ISR by beneficial microorganisms (Pieterse et al., 2014). However, in the next paragraphs, we will describe several examples of ISR by beneficial microorganisms (Pieterse et al., 2014).

Vogel et al. (2016) studied the differences in gene expression in Arabidopsis plants upon inoculation with two model commensal phyllosphere bacteria; S. melonis Fr1 and Methyllobacterium extorquens PA1. Colonization by M. extorquens PA1 resulted in very little transcriptional response from the plant whereas S. melonis Fr1 changed the expression of several hundreds of genes. This corresponds with the findings of Innerebner et al. (2011) where S. melonis Fr1 decreased disease development on A. thaliana while M. extorquens PA1 did not. The transcriptional response induced by S. melonis Fr1 was similar to the response induced by an encounter with the pathogenic P. syringae DC3000 (Vogel et al., 2016). The authors hypothesize that plants detect the presence of S. melonis Fr1 in a similar way as P. syringae and respond with an expression of defense-related genes that are involved in plant protection. However, the plant's response is less severe, probably because S. melonis Fr1 lacks additional virulence factors which are present in P. syringae DC3000. The exact mechanisms still need to be unravelled, since plant mutants defective in several known defense signaling pathways, such as SA and JA signaling, and lacking the FLS2 PRR, showed that these pathways and receptors where not involved. More recently, Ryffel et al. (2016) demonstrated that S. melonis Fr1 induced camalexin production in the host plant. The pathogen P. syringae pv. tomato also induced production of camalexin, yet in higher concentrations. Production of the tryptophan-derived indole alkaloid camalexin, is a typical defense response of Arabidopsis and other plants from the Crucifereae family. Due to its lipophilic nature, camalexin is effective against a wide range of bacteria and fungi by interfering with the integrity of membranes (e.g., by binding to phospholipids). Production of camalexin by Arabidopsis, triggered by S. melonis Fr1 is thus postulated to be the mechanism behind the observed plant protection by this commensal.

The host's immune system does not only target bacterial or fungal pathogens, but can also protect against viral diseases (Lee and Ryu, 2016). Three-year field trials of foliar applications of Bacillus amyloliquefaciens 5B6 showed consistently reduced cucumber mosaic virus accumulation as measured by qPCR (Lee and Ryu, 2016). Observed changes in gene expression in the host plant suggested that activation of SA and ethylene signaling pathways played a key role in the acquired resistance. Also here, the genes upregulated by the biocontrol agent were the same as the genes upregulated in the defense response caused by several viruses, including cucumber mosaic virus (Park et al., 2004). B. amyloliquefaciens 5B6 colonized the phyllosphere successfully as their population size remained stable during seven days after administration of 10^8 CFU/ml until run-off. This contrasted with the sharp decline in population size of strain FZB42, isolated from the soil, showing that strain-specific adaptation traits are important for survival and successful biocontrol in the phyllosphere.

Another intriguing observation was made by Hong et al. (2016) on the known plant-growth promoting rhizosphere bacterium Paenibacillus polymyxa AC-1. This strain inhibited the growth of phyllosphere pathogens P. syringae pv. tomato DC3000 and P. syringae pv. tabaci in an in vitro setting. Cell-free supernatant of P. polymyxa AC-1 also suppressed these pathogens, suggesting that antimicrobial metabolites excreted by the antagonist play a direct antagonistic role (see section “Antimicrobial Metabolites”). Inoculation of the root tips of axenic Arabidopsis seedlings with bacterial suspensions of P. polymyxa AC-1 resulted in a SA and JA-dependent defense reaction. Interestingly, this inoculation of the roots of axenic plants resulted in colonization of the Arabidopsis leaf endosphere with P. polymyxa AC-1. Colonization of the leaf endosphere was 10-fold higher in Arabidopsis mutants with reduced sensitivity to JA and 10-fold lower in mutants deficient in the isoprenoid plant hormone abscisic acid, compared to wild-type plants. The colonization of the leaf endosphere by P. polymyxa AC-1 in JA-deficient plants even caused disease symptoms in the phyllosphere. This indicates that JA negatively impacts the detrimental endophytic growth of AC-1. This illustrates that the plant's defense system is important in regulating the total microbial load and preventing symbiotic bacteria to become invasive.

The mechanisms by which microbes are detected by the host and subsequently trigger the host's immune response, are similar in both non-pathogenic and pathogenic strains (Fig 2). However, non-pathogenic strains lack additional virulence factors, resulting in a milder defense response from the host. Biocontrol agents have the ability to trigger the immune system, and thereby inducing resistance to phylogenetically distinct pathogens. Sometimes, biocontrol agents are very closely related to pathogenic strains, or can even be opportunistic pathogens themselves (e.g., Pseudomonas, R. fascians, P. polymyxa), in this case the difference between ISR and SAR becomes less clear.

CONCLUSION AND FUTURE RESEARCH PERSPECTIVES

The phyllosphere harbors a diverse set of microbes. These microbes interact closely with each other and with the host plant. Amongst them are pathogens, causing disease in the host plant and reducing yields in agriculture, but also beneficial microbes which can be the key to environmentally friendly solutions to protect crops from diseases. These beneficial microbes can inhibit pathogen growth directly, by competing for nutrients and space, by interfering with their communication, by excreting antimicrobial metabolites or enzymes, or by parasitizing on the pathogen. On the other hand, beneficial microbes can trigger the plant's immune response and modulate plant hormone levels, and hereby indirectly inhibiting pathogen growth.
Knowledge on these mechanisms is often gained through \textit{in vitro} experiments using gnotobiotic or sterile plants, while the efficacy of a biocontrol agent needs to be validated in field trials. In this review, we linked these two types of studies and gave an overview of biocontrol mechanisms and adaptation mechanisms that play a role in the phyllosphere. Several mechanisms still need further validation, for example, the characterization of novel antimicrobial peptides, the role of type III secretion systems, the biocontrol potential of small peptides involved in quorum sensing and the impact of bacterially produced plant hormones on the host immune system. Next, we need to understand which mechanisms are at play in field conditions. This question has also been addressed by Köhl et al. (2019). Firstly, which biocontrol mechanisms are active on the field, e.g., some antimicrobial metabolites play a role in biocontrol \textit{in vitro} but not \textit{in vivo}, and reversely (Köhl et al., 2019; Santos Kron et al., 2020)? Secondly, is the biocontrol agent adapted to the phyllosphere in field conditions, and can it disperse in the growing plant (e.g., Wei et al., 2016)? Finally, the biocontrol agent needs to integrate in the resident microbial community. This resident community can positively or negatively influence the biocontrol activity and the success of colonization of the biocontrol agent (Massart et al., 2015).

Techniques that can help us understand the mechanisms playing a role in complex phyllosphere communities are metagenome, metatranscriptome, metaproteome, and metabolome analyses. Eventually, this understanding may allow us to go beyond the application of single strains, and toward designing communities, an idea that is often repeated in biocontrol research (Massart et al., 2015; Singh and Trivedi, 2017). Biocontrol communities possess a variety of complementary adaptation and biocontrol factors, that co-operate in suppressing the disease and surviving on the phyllosphere. Biocontrol agents and communities can be an effective and sustainable alternative to conventional pesticides, which is needed to safeguard our crop production.

**AUTHOR CONTRIBUTIONS**

ML and SL conceived and designed the manuscript. ML wrote the manuscript and prepared the figures. All authors critically reviewed the manuscript and contributed to figure improvement, with special attention toward their specific expertise ranging from phyllosphere bacterial communities, bacterial antimicrobial compounds, probiotics, bacterial genome analysis and functional prediction, plant hormones, plant immune system toward ecology in general and approved the final version of the manuscript.

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**AUTHOR CONTRIBUTIONS**

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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