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1 Crying out for help with root exudates: adaptive mechanisms by 2 which stressed plants assemble health-promoting soil microbiomes.

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10 Highlights

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- The cry-for-help model states that stressed plants assemble protective rhizobiomes.
- Plant attacked by pathogens or herbivores change their root exudation chemistry.
- Specific rhizosphere signals alter the composition and activity of the rhizobiome.
- The modified rhizobiome protects plants via direct and indirect mechanisms.
- Legacy effects on the soil microbiome can benefit the next generation of plants.
- 17
- 18 Abstract
- 19

20 Plants employ immunological and ecological strategies to resist biotic stress. Recent evidence 21 suggests that plants adapt to biotic stress by changing their root exudation chemistry to 22 assemble health-promoting microbiomes. This so-called 'cry-for-help' hypothesis provides a 23 mechanistic explanation for previously characterized soil feedback responses to plant disease, 24 such as the development of disease-suppressing soils upon successive cultivations of take all-25 infected wheat. Here, we divide the hypothesis into individual stages and evaluate the 26 evidence for each component. We review how plant immune responses modify root exudation chemistry, the impact this has on microbial activities, and the subsequent plant 27 28 responses to these activities. Finally, we review the ecological relevance of the interaction, 29 along with its translational potential for future crop protection strategies.

- 31 Short title: A systems review of the cry-for-help hypothesis.
- 32 Key words: plant immune responses; root exudates; rhizosphere signals; rhizobiome; root-
- 33 microbe interactions; induced systemic resistance; soil feedback responses.

1 Introduction

2

3 Soil is a critical resource for agricultural crop production. While agri-technological progress 4 has made spectacular progress over recent decades, most innovations are based on 5 agrochemicals and crop breeding technologies. By comparison, soil as a target for crop 6 improvement has largely been overlooked, despite ample evidence for its plant protective 7 activities [1]. The ability of soil to suppress plant diseases is a common characteristic of soil 8 health and is determined by the soil- and root-associated microbiome [2*,3]. While crop 9 rotation, conservation tillage and soil organic amendments improve soil health [1], these 10 practices are not always financially feasible for farmers. However, disease-suppressive soil 11 activity can also develop in high-intensity production systems that rely on successive crop 12 monocultures. The classic example is take-all decline, during which continued wheat 13 cultivation in soil infested with the pathogenic take-all fungus Gaeumannomyces graminis 14 pv. tritici initially leads to increased disease, followed by a progressive decline in disease [4]. 15 There are ample other examples whereby prolonged disease exposure leads to disease-16 suppressing soil activity [2^{*}]. These observations have led to the hypothesis that disease- and 17 herbivore-exposed plants employ a strategy that involves active selection and/or recruitment 18 of disease-suppressing soil microbiomes. This adaptive strategy not only benefits the plant 19 that is under attack, but also subsequent plant generations, which is why these feedback 20 responses are sometimes referred to as 'legacy' or 'soil memory' effects [5-8]. Analogous to 21 aboveground multitrophic interactions between plants and arthropods [9], the mechanisms 22 initiating this long-term adaptation are encompassed by the 'cry-for-help' hypothesis (Figure 23 1). This concept gained significant traction in the rhizosphere research community after a 24 pioneering study that identified shifts in the microbial community structure of a disease-25 suppressive soil following prolonged cultivation of Rhizoctonia solani-infected sugar beet 26 [10^{**}]. In subsequent years, various other studies have confirmed enrichment of disease-27 suppressing microbes in disease-suppressive soils [2*]. In addition, there is an impressive 28 body of evidence to support that root exudation chemistry is critical for the assembly of plant 29 health-promoting microbiomes [11]. However, there remain knowledge gaps in the 30 successive stages predicted by belowground cry-for-help model. In this review, we evaluate 31 the evidence for each stage of the process, after which we will discuss the ecological relevance 32 and translational opportunities of this long-term plant adaptation strategy.



Figure 1: Model of the successive stages of the 'cry-for-help' hypothesis. Local and systemic signals elicited by pathogens or herbivores activate root immune responses (stage 1), which alter root exudation profiles of primary and secondary metabolites with biocidal and/or semiochemical activities (stage 2). Altered root exudation profiles influence the microbiome by recruiting and selecting specific microbiota and inducing microbial activities (stage 3). Some of these activities involve direct and indirect mechanisms that antagonize plant attackers, such as antibiosis, nutrient competition and induced systemic resistance (ISR; stage 4).

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8 Stage I: root immune responses to below- and aboveground attackers

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10 Of all plant tissues, roots are exposed to the highest microbial density and diversity [12]. In 11 that regard, it is unsurprising that immune responses by roots differ from those by above-12 ground tissues [13]. Detailed studies of root responses to microbe-associated molecular 13 pattern MAMPs have revealed that defence-related gene expression is spatially restricted to 14 specific cell types, which vary according the applied MAMP [14**,15*]. The immunological 15 differences between roots and shoots may result from the lack photosynthesising 16 chloroplasts in the roots, which generate high concentrations of defence-enhancing reactive 17 oxygen and nitrogen species [16]. Furthermore, although pathogen-infected roots are 18 capable of accumulating salicylic acid (SA) [17], the initial biosynthetic steps occur in 19 chloroplasts, indicating phloem-mediated transport of SA and/or derivatives from shoot to 20 root tissues thereof [18,19]. Indirect evidence that jasmonic acid (JA-) and SA-dependent 21 immune reactions in roots generate rhizosphere-active signals is based on rRNA amplicon 22 sequencing experiments, showing that exogenous hormone treatments or mutations in these 23 pathways influence the root-associated microbiome [20*,21*]. In addition, systemic immune 24 responses to aboveground pests and defence elicitors have been reported to alter root 25 interactions with belowground microbes in a SA-dependent manner [22,23]. In the following 26 section, we will review how root immune responses lead to exudation and accumulation of 27 rhizosphere-active metabolites and derivatives thereof.

1 Stage II: stress-induced changes in root exudation of antimicrobials and 2 semiochemicals

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4 Roots release primary metabolites, such as carbohydrates, amino acids, organic acids and 5 membrane lipids, which provide energy and nutrients to the soil microbiome [24]. The 6 concentration and composition of these compounds in root exudates changes upon exposure 7 to biotic stress and can have specific signalling effects in the rhizosphere. For instance, foliar 8 infection of Arabidopsis by Pseudomonas syringae increases L-malic acid exudation, leading 9 to increased root colonisation by resistance-inducing Bacillus subtilis [25*]. In cucumber, local 10 root infection by pathogenic Fusarium oxysporum alters concentrations of 89 mostly primary 11 metabolites in exudates from distal roots, of which increased tryptophan and reduced 12 raffinose correlated with root colonisation by beneficial Bacillus amyloliquefaciens [26**]. 13 However, it seems unlikely that primary metabolites alone are responsible for the assembly 14 of disease-suppressive root/soil microbiomes. Secondary root metabolites seem equally, if 15 not more important, since they are often inducible by biotic stress, are less quickly 16 metabolized by microbes, and typically have antimicrobial and/or signalling activities. Based 17 on previous studies reviewed by [27-29], Figure 2 provides an overview of the main 18 biochemical pathways controlling pathogen- and herbivore-inducible secondary metabolites 19 with antimicrobial and/or signalling activity. It is important to note that rhizosphere 20 chemistry, rather than root (exudation) chemistry, is responsible for shaping root- and soil-21 associated microbiomes. Rhizosphere chemistry is the sum of root exudation chemicals, their 22 breakdown products and microbial products of soil-derived chemicals. A recent study 23 developed a new method for chemically profiling non-sterile rhizosphere soil, providing a 24 powerful technique to identify semiochemicals in non-sterile rhizosphere soil and link them 25 to rhizobiome activities [30^{**}].

26 MAMP-treated Arabidopsis roots increase the expression of CYP71A12 and MYB51 [14^{**}], 27 which control biosynthesis of tryptophan-derived defence compounds, such as camalexin and 28 indolic glucosinolates. These stress-responsive metabolites often have both antimicrobial and 29 signalling activities [31,32], and are commonly detected in root exudates [14**,33,34**]. 30 Their increased exudation from defence-expressing and/or damaged roots can influence the 31 root-associated microbes, which was recently demonstrated for camalexin [34^{**}]. Similarly, 32 parasitic nematodes increase strigolactone (SL) biosynthesis in tomato roots [35]. Irrespective 33 the exact role of SLs in plant immune signalling [36], exudation of SLs can influence the root-34 associated microbiome by stimulating hyphal branching and infection by arbuscular 35 mycorrhizal fungi (AMF) [37]. Stress-induced rhizodeposition of defence hormones may also 36 play an important role in shaping the soil- and root-associated microbiome [38*]. In 37 particular, SA is commonly detected in plant root exudates [39*] and can attain 38 concentrations in soil sufficient to induce resistance in neighbouring plants [40^{*}]. Because SA 39 can be incorporated in iron-chelating siderophores by rhizosphere bacteria [39^{*}], it is likely 40 that rhizosphere accumulation of SA selects for siderophore-producing rhizobacteria, which 41 contribute to disease suppression [41].

1 Like pathogens, herbivores can induce exudation of rhizosphere-active root metabolites. 2 Infestation of maize roots by larvae of Diabrotica vergifera induces emission of the 3 sesquiterpene (E)- β -caryophyllene (E β c), which recruits soil-borne entamopathogenic 4 nematodes [42]. Over-expression of an EBc synthase gene in the genetic background of a EBc-5 deficient variety [43] revealed that Eßc increases growth and susceptibility to the soil-borne 6 fungal pathogen Colletotrichum graminicola [43,44*], suggesting that EBc may have wider-7 ranging impacts on soil microbes. In cereals, herbivory and wounding induce increase the 8 accumulation of aglycone benzoxazinoids, such as methoxy-2H-1,4-benzoxazin-3(4H)-one 9 (DIMBOA) [45]. Three recent studies have used maize mutants in BX production to determine 10 the extent by which these metabolites influence root- and soil-associated microbiomes 11 [46^{**},47^{**},48^{*}], all reporting significant effects on plant- and soil-associated microbiomes. 12 Hu et al. [46^{**}] demonstrated that soil conditioned by BX-producing maize induces JA-13 dependent resistance against herbivores, which was linked to the presence and activity of 6-14 methoxy-benzoxazolin-2-one (MBOA). Since DIMBOA acts as a within-plant defence signal 15 [49], Cotton et al. [47^{**}] investigated whether BX biosynthesis genes influence the 16 composition of the wider root metabolome. They reported that the bx1 and bx2 mutations 17 have major impacts on the secondary metabolite profiles in roots, suggesting that the 18 effects of BXs on root-associated microbes could partially be caused by BX-controlled root 19 exudates, rather than BXs themselves. Indeed, correlation analysis between differentially 20 abundant metabolites and bacterial taxa pointed to a dominant role of BX-controlled root 21 metabolites, including compounds with known signalling activities in the rhizosphere, such 22 as flavonoids [47^{**}]. More research is needed to determine the (in)direct signalling 23 activities of BXs in the soil. Does biotic stress increase DIMBOA exudation and MBOA 24 accumulation in the soil? If so, does MBOA act as a stress-induced soil-mobile signal that 25 alters root exudation patterns in systemic roots and roots of neighbouring plants? And 26 finally, does the belowground signalling activity of BXs extend to other plant species, such 27 as wheat, raising the possibility that BXs could act as the regulatory signals driving take-all 28 decline?

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30



1 Figure 2: Scheme of the shikimate (A) and terpenoid (B) pathways, generating stress-inducible secondary

2 metabolites in plant roots with previously reported anti-microbial and/or semiochemical activity in the soil.

3 Coloured boxes show examples of compounds within each class.

1 Stage III Impacts of root exudates on the root- and soil-associated microbiome

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The 'cry-for-help' hypothesis postulates that specific components of root exudates from stressed plants favour recruitment of beneficial microbes and constrain the development of pathogens. This reshaping of the rhizosphere involves multiple mechanisms. Exudates may include substrates for microbial growth, elicit chemotactic responses and facilitate root colonisation, while antimicrobial compounds limit development of susceptible microbial communities. Exudates may also interact with microbial quorum sensing systems (QSS) or be processed by community members, eliciting the release of microbially-derived metabolites.

10 As mentioned above, Liu et al. showed that local infection of cucumber roots by Fusarium 11 oxysporum increases tryptophan exudation and reduces raffinose exudation [26^{**}], resulting 12 in increased colonization by beneficial Bacillus amyloliquefaciens SQR9 (BaSQR9) and reduced 13 pathogen colonisation, thus pushing the rhizobiome towards plant-beneficial associations. 14 They furthermore showed that the growth-promoting activity of BaSQR9 results from 15 tryptophan-dependent auxin production, indicating further feedback loops between plant 16 and bacteria. Other studies have implicated organic acids as important signals, acting as 17 recruitment signals for plant growth-promoting rhizobacteria (PGPR) in cucumber, tomato, 18 banana, watermelon and Arabidopsis. [25*,50-53]. As reviewed above, BXs are important 19 antimicrobial metabolites [45,49]. Neal et al. [54*] found that DIMBOA is chemo-attractive to 20 plant-beneficial P. putida KT2440 bacteria, activating genes associated with bacterial motility, 21 QSS and breakdown of N-heteroaromatic compounds. Such selection for BX tolerance can 22 also influence potentially hostile organisms. Sanders et al. [55] reported that BOA, a toxic 23 degradation product of DIBOA, selects for BX-resistant Fusarium sp. in maize with the 24 potential for grain contamination by mycotoxins.

25 For many microbial responses to root exudation metabolites, bacterial stress seems a 26 recurrent theme. Exposure of PGPR to root exudates activates genes associated with nutrient 27 responses and motility, but also the production of antibacterial and antifungal substances, 28 degradation of aromatic compounds and microbial stress responses [56*,57,58*]. Thus, while 29 many root exudates act as nutrients and recruitment factors, other root exudates induce 30 microbial stress that lead to plant-beneficial activities. For instance, quorum sensing signals 31 (QSS) activate transcriptional stress responses in bacteria, once a certain population density 32 has been reached. Given the ubiquity of QSS, it is unsurprising that plants have evolved to 33 respond to QSS molecules and manipulate QSS responses [59]. Sweet basil releases 34 rosmarinic acid (RA) when infected by pathogenic P. aeruginosa PA01 and PA14 [60]. RA is 35 toxic to bacteria at high concentrations, but also binds to the response regulator RhIR triggering premature QSS responses [61**]. This QSS system is commonly found in 36 37 Pseudomonads including PGPR, and may therefore also regulate PGPR responses, such as 38 biofilm formation and antibiosis. Indeed, the protective effect of Pseudomonas aureofaciens 39 strain 30-84 against take-all disease has been attributed to phenazine antibiotic production 40 that is regulated by QSS [62,63]. Bacterial stress responses in the rhizosphere can also be an 41 indirect consequence of microbial competition. For instance, saprotrophic fungi consume

1 root exudates rapidly, which reduces nutrient availability to rhizobacteria that in turn triggers

2 rhizobacterial production of antifungal compounds [64].

3 The effects of rhizosphere chemistry on the beneficial microbiome activities in the soil can be 4 long-lived. Yuan et al. found that five generations of Arabidopsis plants inoculated with 5 Pseudomonas syringae DC3000 (Pst) leads to disease suppression in the sixth generation, 6 which was associated with changes in soil microbial community [65**]. This study 7 furthermore showed that >50 root exudation compounds changed upon infection. Soil 8 complementation experiments with mixtures of components identified long chain organic 9 acids as the underpinning soil signals, stimulating microbiome-mediated induced systemic 10 resistance (ISR). Similarly, Hu et al. reported soil feedback responses that were linked to soil 11 accumulation of MBOA, which induced JA-dependent resistance in maize plants of the next 12 generation [46^{**}]. Finally, Berendsen et al [66^{**}] isolated three community members that 13 accumulated in soils of downy mildew-infected Arabidopsis plants, and found that this 14 assemblage interacted to induce biofilm formation and ISR in subsequent plant generations. 15 Notably, in all three examples, the response of the soil microbiome was critical for the 16 beneficial ISR response of the host plant.

17 Just as plants have evolved to respond to microbial signals, microbes have evolved to respond 18 to plant signals, including plant growth regulators involved in biotic stress responses. 19 Treatment of both plants and soil with SA, JA and ethylene (ET) induces changes in root 20 exudates and rhizosphere communities, whereas mutations in plant JA signalling reduces root 21 exudates associated with PGPR chemotaxis or that act as growth substrates for PGPR and N-22 fixing diazotrophs [20*,38*,67]. The emerging pattern suggests a complex network of 23 interactions between plants and soil/plant-associated microbiomes, which are mediated by a 24 multitude of chemicals signals that are derived from both plants and microbes. Bruto et al. 25 [68] attempted to identify plant-beneficial function contributing (PBFC) genes in 26 Proteobacterial PGPR. Interestingly, none of these genes were found in all PGPR, and many 27 were found in non-PGPR. However, combinations of PBFC genes were only found in particular 28 taxonomic subgroups of PGPR, indicating that specific assortments were associated with the 29 beneficial trait. It is therefore plausible that similar, if not greater, complexity exists in PGPR 30 responses to root exudates with combinations of signals and signalling mechanisms 31 contributing to microbial recruitment and development.

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Stage IV: mechanisms by which the root- and soil-associated microbiome suppress pests and diseases

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The mechanisms underpinning disease-suppressing soil activity are complex [2*]. Apart from direct mechanisms, such as parasitism and the production of biocidal compounds, beneficial rhizosphere microbes suppress soil-borne attackers indirectly through competition for (micro)nutrients and elicitation of ISR. Of these, ISR provides protection against both belowand above-ground attackers [69]. Much knowledge about the mechanisms underpinning ISR come from the interaction between Arabidopsis and *Pseudomonas simiae* WCS417. Early

1 studies have shown that ISR is controlled by a SA-independent signalling pathway that primes 2 distal tissues for JA- and ET-dependent defence genes and cell wall-based defences [70,71]. 3 While there are exceptions, ISR in other plant-microbe interactions often follows a similar 4 signalling signature [69]. This commonality could be explained by the fact that ISR-eliciting 5 microbes trigger a general nutrient deficiency response that results in systemic up-regulation 6 of ISR-related immune pathways. Castrillo et al. recently demonstrated that inoculation of 7 Arabidopsis with a synthetic rhizobacterial community induces a phosphate starvation 8 response (PSR), which modulates systemic plant immunity and that is under control by the 9 regulatory gene PHR1 [72**]. Interestingly, PHR1 has previously been reported to control 10 ISR-related immune pathways, including JA signalling [73] and production of callose-11 stimulating glucosinolates [74]. If these mechanisms apply to other plant species, the PSR by 12 ISR-eliciting microbiota could lead to increased exudation of SLs and recruitment of 13 endophytic fungi, such as AMF, which in turn alter root-associated microbial populations 14 [75,76]. Furthermore, a recent study of the Arabidopsis-WCS417 model system found that 15 bacterial induction of the ISR-regulatory transcription factor MYB72 and downstream beta-16 glucosidase BGLU42 induce an iron-deficiency response that is associated with increased root 17 exudation of scopoletin [77**]. This iron-mobilizing metabolite has selective impacts on the 18 root-associated microbiome, including biocidal activity on soil-borne pathogenic fungi. A 19 recent study by Vogel et al. confirmed the importance of selected scopoletin derivatives in 20 shaping synthetic rhizobiome communities of Arabidopsis via redox-mediated mechanisms [78^{**}] Together, these recent studies illustrate that interactions between roots and disease-21 22 suppressing bacteria trigger a succession of signalling events, resulting in a range of disease-23 suppressive mechanisms, including ISR, recruitment of biocontrol fungi, (micro)nutrient 24 competition, and antibiosis (Figure 3).



Figure 3: emerging roles for phosphate (Pi; purple) and iron (Fe; orange) starvation responses in the orchestration of diseasesuppressive mechanisms in the microbial biosphere of the plant. The model is based on recent evidence for reciprocal signalling events between nutrient starvation signalling in the host. systemic immune responses (ISR), and diseasesuppressing activities by the root- and soil-associated microbiome [72**, 77**).

1 Conclusions: ecological relevance and translational opportunities

2

3 There is ample evidence to support the individual components of the cry-for-help hypothesis. 4 However, the outcome of the response is not always adaptive and can come with ecological 5 trade-offs. The signals used by plants to recruit plant-beneficial organisms in the soil can be 6 hijacked by parasitic organisms. For instance, emission of Eßc increases infection by the soil-7 borne fungal pathogen Colletotrichum graminicola [44*], exudation of BXs enhances damage 8 by pathogenic fungi and the western corn rootworm [55,79], and exudation of AMF-recruiting 9 SLs can be exploited by pathogenic nematodes and parasitic weeds to locate their host [80]. 10 We speculate that these ecological trade-offs are determined by soil quality. Healthy soils 11 with high microbial biodiversity more likely contain robust networks of beneficial 12 rhizobacteria than poorer soils with low biodiversity [81]. Once an interaction with beneficial 13 microbiota has been initiated, the subsequent signalling cascade leads to the establishment 14 of a chemical and biological environment that is mutually beneficial to both partners. In 15 situations where the soil fails to provide fast-responding beneficials, due to loss of biodiversity 16 by overfertilization, soil compaction, or soil inversion, the cry-for-help is more likely to be 17 hijacked by parasitic microbes and arthropods. Recent evidence that plant and microbial 18 nutrient starvation responses control the establishment of plant health-promoting 19 microbiomes [72^{**},77^{**}] is directly antagonistic to the often excessive amounts of fertilizer 20 applied in modern agriculture [82^{**}]. Furthermore, human selection for aboveground yield 21 under high fertilizer input have resulted in plant varieties with rudimentary root systems that 22 communicate less effectively with the soil microbiome [82^{**}]. While the importance of soil 23 microbiomes is increasingly being recognised by farmers and the wider agri-tech sector, a 24 better mechanistic understanding of the individual components of the cry-for-help hypothesis 25 is necessary to reliably exploit the benefits of soil-preserving land management, biocontrol 26 inoculations, and crop breeding programmes selecting for soil-health promoting root traits.

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1 References

- 2
- 3 1. Larkin RP: Soil health paradigms and implications for disease management. Annu Rev 4 Phytopathol 2015, **53**:199-221. 5 2. Gómez Expósito R, de Bruijn I, Postma J, Raaijmakers JM: Current Insights into the Role of 6 Rhizosphere Bacteria in Disease Suppressive Soils. Front Microbiol 2017, 8. 7 8 * This review provides an overview of the literature about disease-suppressive soils. The 9 paper includes a comprehensive table outlining the microbial taxa associated with disease-10 suppresive activity, which is based on recent 'omics' studies. 11 12 3. Dubey A, Malla MA, Khan F, Chowdhary K, Yadav S, Kumar A, Sharma S, Khare PK, Khan 13 ML: Soil microbiome: a key player for conservation of soil health under changing 14 climate. Biodivers Conserv 2019, 28:2405-2429. 15 4. Kwak YS, Weller DM: Take-all of Wheat and Natural Disease Suppression: A Review. 16 Plant Pathol J 2013, 29:125-135. 17 5. Berendsen RL, Pieterse CM, Bakker PA: The rhizosphere microbiome and plant health. 18 Trends Plant Sci 2012, **17**:478-486. 19 6. Raaijmakers JM, Mazzola M: Soil immune responses. Science 2016, 352:1392-1393. 20 7. Wilkinson SW, Mageroy MH, Lopez Sanchez A, Smith LM, Furci L, Cotton TEA, Krokene P, 21 Ton J: Surviving in a Hostile World: Plant Strategies to Resist Pests and Diseases. 22 Annu Rev Phytopathol 2019, 57:505-529. 8. Kong HG, Song GC, Ryu CM: Inheritance of seed and rhizosphere microbial communities 23 24 through plant-soil feedback and soil memory. Environ Microbiol Rep 2019, 11:479-25 486. 26 9. Turlings TCJ, Erb M: Tritrophic Interactions Mediated by Herbivore-Induced Plant 27 Volatiles: Mechanisms, Ecological Relevance, and Application Potential. Annu Rev 28 Entomol 2018, **63**:433-452. 29 10. Mendes R, Kruijt M, de Bruijn I, Dekkers E, van der Voort M, Schneider JH, Piceno YM, 30 DeSantis TZ, Andersen GL, Bakker PA, et al.: Deciphering the rhizosphere 31 microbiome for disease-suppressive bacteria. Science 2011, 332:1097-1100. 32 33 ** The authors profiled operational taxonomic units (OTUs) of microbial communities in a 34 disease suppressive soil and identified 17 microbial taxa that are closely associated with 35 disease suppressing activity, including bacterial isolates producing antimicrobial peptides. 36 37 11. Olanrewaju OS, Ayangbenro AS, Glick BR, Babalola OO: Plant health: feedback effect of 38 root exudates-rhizobiome interactions. Appl Microbiol Biotechnol 2019, 103:1155-39 1166. 40 12. Hassani MA, Durán P, Hacquard S: Microbial interactions within the plant holobiont. 41 Microbiome 2018, 6:58. 42 13. Chuberre C, Plancot B, Driouich A, Moore JP, Bardor M, Gügi B, Vicré M: Plant Immunity 43 Is Compartmentalized and Specialized in Roots. Front Plant Sci 2018, 9. 44 14. Millet YA, Danna CH, Clay NK, Songnuan W, Simon MD, Werck-Reichhart D, Ausubel FM: 45 Innate immune responses activated in Arabidopsis roots by microbe-associated 46 molecular patterns. Plant Cell 2010, 22:973-990.

1	
2	** Using Arabidopsis promoter: GUS reporter lines to visualise spatial distributions of PAMP-
3	induced immune responses, the authors demonstrate that expression of innate immunity in
4	roots follows spatially distinct patterns. The authors also demonstrate that flg22-induced
5	immunity in the root elongation zone requires components of the ethylene response
6	pathway and production of tryptophan-derived secondary metabolites.
7	
8	15. Poncini L, Wyrsch I, Dénervaud Tendon V, Vorley T, Boller T, Geldner N, Métraux J-P,
9	Lehmann S: In roots of Arabidopsis thaliana, the damage-associated molecular
10	pattern AtPep1 is a stronger elicitor of immune signalling than flg22 or the chitin
11	heptamer. PLOS One 2017, 12 :e0185808.
12	
13	* This study used a range of promoter:YFP lines to visualise activities of jasmonic acid-,
14	salicylic acid-, ethylene-, and reactive oxygen species-dependent signalling pathways in
15	roots following treatments with different MAMPs. The results confirm that root immune
16	responses are spatially compartmentalised, depending on the applied MAMPs.
17	
18	16. Serrano I, Audran C, Rivas S: Chloroplasts at work during plant innate immunity. J Exp
19	Bot 2016, 67 :3845-3854.
20	17. Balmer D, de Papajewski DV, Planchamp C, Glauser G, Mauch-Mani B: Induced
21	resistance in maize is based on organ-specific defence responses. Plant J 2013,
22	74 :213-225.
23	18. Strawn MA, Marr SK, Inoue K, Inada N, Zubieta C, Wildermuth MC: Arabidopsis
24	isochorismate synthase functional in pathogen-induced salicylate biosynthesis
25	exhibits properties consistent with a role in diverse stress responses. J Biol Chem
26	2007, 282 :5919-5933.
27	19. Fragniere C, Serrano M, Abou-Mansour E, Metraux JP, L'Haridon F: Salicylic acid and its
28	location in response to biotic and abiotic stress. FEBS Lett 2011, 585:1847-1852.
29	20. Carvalhais LC, Dennis PG, Badri DV, Tyson GW, Vivanco JM, Schenk PM: Activation of the
30	jasmonic acid plant defence pathway alters the composition of rhizosphere
31	bacterial communities. PLoS One 2013, 8 :e56457.
32	
33	* The authors have used 16S rRNA gene sequencing to demonstrate that elicitation of JA-
34	dependent immunity in Arabidopsis by volatile methyl-jasmonate changes root-associated
35	bacterial communities.
36	
37	21. Lebels SL, Paredes SH, Lundberg DS, Breakfield N, Genring J, McDonald M, Malfatti S,
38	Glavina del Rio T, Jones CD, Tringe SG, et al.: Salicylic acid modulates colonization of
39	the root microbiome by specific bacterial taxa. Science 2015, 349:860-864.
40	* The outhous compound 1/C xDNA core converses between Archidensis wild type plants
41 42	and solidie asid (SA) signalling mutants, demonstrating that SA dependent immunity
4Z 42	determines the abundance and diversity of reat endephytic bacterial communities
43 44	determines the abundance and diversity of root-endophytic pacterial communities.
45	22, Song GC, Lee S, Hong J, Choi HK, Hong GH, Bae DW, Mysore KS, Park YS, Ryu CM:
46	Aboveground insect infestation attenuates belowground Agrobacterium-mediated
47	genetic transformation. New Phytol 2015, 207:148-158.

1	23. Yang JW, Yi H-S, Kim H, Lee B, Lee S, Ghim S-Y, Ryu C-M: Whitefly infestation of pepper
2	plants elicits defence responses against bacterial pathogens in leaves and roots
3	and changes the below-ground microflora. J Ecol 2011, 99:46-56.
4	24. Canarini A, Kaiser C, Merchant A, Richter A, Wanek W: Root Exudation of Primary
5	Metabolites: Mechanisms and Their Roles in Plant Responses to Environmental
6	Stimuli . Front Plant Sci 2019, 10 .
7	25. Rudrappa T, Czymmek KJ, Pare PW, Bais HP: Root-secreted malic acid recruits beneficial
8	soil bacteria . Plant Physiol 2008, 148 :1547-1556.
9	
10	* This is the first study that provides experimental evidence that aboveground disease alters
11	root exudation of a primary root metabolite (L-malic acid), resulting in increased root
12	colonization by a plant-beneficial rhizobacterial strain. The authors propose that L-malic acid
13	acts as a specific signal for chemotaxis and recruitment in the rhizosphere.
14	
15	26. Liu Y, Chen L, Wu G, Feng H, Zhang G, Shen Q, Zhang R: Identification of Root-Secreted
16	Compounds Involved in the Communication Between Cucumber, the Beneficial
17	Bacillus amyloliquefaciens, and the Soil-Borne Pathogen Fusarium oxysporum. Mol
18	Plant Microbe In 2017, 30 :53-62.
19	
20	** The authors show that inoculation of part of the cucumber root system changes
21	numerous root exudates and interactions with pathogens and PGPRs in distal roots, which
22	was linked to increased exudation of tryptophan and decreased exudation of raffinose.
1 2	
23 24	27 Massalha H. Korenhlum F. Tholl D. Abaroni A: Small molecules below-ground: the role
2 4 25	of specialized metabolites in the rhizosphere Plant 12017 90.788-807
25	28 Stringlis IA de longe R Dieterse CMI: The Age of Coumarins in Plant-Microbe
20	Interactions Plant Cell Physiol 2019 60:1405-1419
28	29 Hassan S. Mathesius U: The role of flavonoids in root-rhizosphere signalling :
29	opportunities and challenges for improving plant-microbe interactions JExp Bot
30	2012. 63 :3429-3444.
31	30. Petriaco P. Williams A. Cotton A. McFarlane AF. Rolfe SA. Ton J: Metabolite profiling of
32	non-sterile rhizosphere soil. Plant J 2017. 92 :147-162.
33	•
34	** This study presents an adjustable experimental system for characterising rhizosphere
35	chemistry. The method involves an experimental system to collect, extract and analyse
36	samples by mass spectrometry. Subsequent statistical comparison of extracts between
37	plant-free and plant-containing soil allows inference of rhizosphere-enriched chemistry.
38	
39	31. Bednarek P, Osbourn A: Plant-Microbe Interactions: Chemical Diversity in Plant
40	Defense . Science 2009, 324 :746-748.
41	32. Stotz HU, Sawada Y, Shimada Y, Hirai MY, Sasaki E, Krischke M, Brown PD, Saito K,
42	Kamiya Y: Role of camalexin, indole glucosinolates, and side chain modification of
43	glucosinolate-derived isothiocyanates in defense of Arabidopsis against Sclerotinia
44	sclerotiorum . Plant J 2011, 67 :81-93.

1	33. Monchgesang S, Strehmel N, Schmidt S, Westphal L, Taruttis F, Muller E, Herklotz S,
2	Neumann S, Scheel D: Natural variation of root exudates in Arabidopsis thaliana-
3	linking metabolomic and genomic data. Sci Rep-UK 2016, 6:29033.
4	34. Koprivova A, Schuck S, Jacoby RP, Klinkhammer I, Welter B, Leson L, Martyn A, Nauen J,
5	Grabenhorst N, Mandelkow JF, et al.: Root-specific camalexin biosynthesis controls
6	the plant growth-promoting effects of multiple bacterial strains. Proc Natl Acad Sci
7	USA 2019, 116 :15735-15744.
8	
9	** The authors performed a genetic screen of 172 Arabidopsis accessions for impacts on
10	microbial sulfatase activity in the soil, which led to the identification of the CYP71A27 gene.
11	The study furthermore shows that this gene controls camalexin production by roots, which
12	influences the interaction with plant growth-promoting bacteria.
13	
14	35 Xu X Eang P Zhang H Chi C Song L Xia X Shi K Zhou Y Zhou I Yu I: Strigolactones
15	nositively regulate defense against root-knot nematodes in tomato / Evn Bot 2019
16	70.1325-1337
17	36 Escudero Martinez CM Guarneri N Overmars H van Schaik C Bouwmeester H Buvter-
10 12	Spira C. Goverse A: Distinct roles for strigolactones in cyst nematode parasitism of
10	Arabidonsis roots Eur Plant Pathol 2019 154.129-140
20	27 Akiyama K. Matsuzaki K. Hayashi H: Plant sesquiternenes induce hyphal branching in
20	37. Akiyama K, Matsuzaki K, Hayasin H. Flant sesquiterpenes induce hypital branching in arbuscular mycorrhizal fungi. Natura 2005. 135 :824-827
21	28 Canvalhais I.C. Donnis D.C. Schonk DM: Blant defence inducers ranidly influence the
22 22	diversity of bacterial communities in a potting mix. Applied Soil Ecol 2014, 84:1-5
23	diversity of bacterial communities in a potting mix. Applied Soll Ecol 2014, 04.1-5.
24 25	* The authors have used 16S rPNA gene sequencing to demonstrate that treatment of
25 26	common potting soil with SA MeIA ARA and ET has differential impacts on the microbial
20	community structure
21 つ0	community structure.
20 20	20 Bakker DAHM Ban L. Marcada Planca & Phizabactorial caliculate production provoker
27	boodschool Diant Coil 2014 292:1 14
3U 21	Headaches: Piant 3011 2014, 302 :1-10.
31 22	* This comprohensive review covers the role and impact of SA on rhizebasteria. Apart from
ວ∠ າາ	incorporation of SA into bacterial siderenbores, the authors review evidence that SA can
33 24	influence other besterial estivities including entibiotic resistance, biofilm formation and
34 25	minuence other bacterial activities, including antibiotic resistance, biomin formation and
35	molility.
30	40. Charles and C. Cim III. Kin CO. Due CM. Beat we diste drive alternative in a frantamic
3/	40. Cheol Song G, Sim HJ, Kim SG, Kyu CM: Root-mediated signal transmission of systemic
38	acquired resistance against above-ground and below-ground pathogens. Ann Bot
39	2016.
40	
41	* This study demonstrates that chemical elicitation of systemic acquired resistance (SAR) in
42	tobacco induces SAR in neighbouring plants via soil-transmitted SA.
43	
44	41. Aznar A, Dellagi A: New insights into the role of siderophores as triggers of plant
45	immunity: what can we learn from animals? J Exp Bot 2015. 66:3001-3010.

1	42. Rasmann S, Kollner TG, Degenhardt J, Hiltpold I, Toepfer S, Kuhlmann U, Gershenzon J,
2	Turlings TC: Recruitment of entomopathogenic nematodes by insect-damaged
3	maize roots. Nature 2005, 434 :732-737.
4	43. Degenhardt J, Hiltpold I, Kollner TG, Frey M, Gierl A, Gershenzon J, Hibbard BE, Ellersieck
5	MR, Turlings TC: Restoring a maize root signal that attracts insect-killing nematodes
6	to control a major pest. Proc Natl Acad Sci USA 2009, 106:13213-13218.
7	44. Fantaye CA, Kopke D, Gershenzon J, Degenhardt J: Restoring (E)-beta-Caryophyllene
8	Production in a Non-producing Maize Line Compromises its Resistance against the
9	Fungus Colletotrichum graminicola. J Chem Ecol 2015, 41:213-223.
10	
11	* The authors demonstrate that constitutive production of (E)- β -caryophyllene by maize
12	roots enhances growth and virulence of the hemi-biotrophic soil fungus Colletotrichum
13	graminicola.
14	
15	45. Niemeyer HM: Hydroxamic acids derived from 2-hydroxy-2H-1,4-benzoxazin-3(4H)-
16	one: key defense chemicals of cereals. J Agric Food Chem 2009, 57:1677-1696.
17	46. Hu L, Robert CAM, Cadot S, Zhang X, Ye M, Li B, Manzo D, Chervet N, Steinger T, van der
18	Heijden MGA, et al.: Root exudate metabolites drive plant-soil feedbacks on growth
19	and defense by shaping the rhizosphere microbiota. Nature Commun 2018, 9 :2738.
20	
21	** Using rRNA gene seqeuncing, the authors compared soil- and root-associated microbial
22	communities of field-grown wild-type maize to that of the BX-deficient bx1 mutant of maize
23	and found significant effects by benzoxazinoids (BXs). They furthermore show that BX-
24	producing wild-type plants augment the soil's ability to induce JA-dependent resistance
25	against herbivores. Chemical analyses and complementation experiments indicated that this
26	soil legacy effect is caused by DIMBOA breakdown product 6-methoxy-benzoxazolin-2-one
27	(MBOA), which could still be detected in soil at 6 months after cultivation.
20	47 Catton TEA Datriaga D. Camaran DD. Masalmani MA. Schwarzanhachar D. Dalfa SA. Tan
27 20	47. Cotton TEA, Pethacq P, Cameron DD, Mesennani MA, Schwarzenbacher R, Rohe SA, Ton I: Metabolic regulation of the maize rhizobiome by benzovazinoids. <i>ISME 1</i> 2019
30	13.1647_{-1658}
32	13.1047 1030.
33	** This study compared the rhizohiomes (rRNA gene sequencing) of different root types
34	between wild-type maize and three by mutants in different steps of the benzovazinoid (BX)
35	pathway. Untargeted mass spectrometry analysis of the root types furthermore revealed a
36	substantial impact of the bx mutations on the root metabolome, supporting a within-plant
37	regulatory function of BXs. Subsequent correlation analysis between BX-dependent root
38	metabolites and bacterial taxa suggested a dominant role for BX-regulated metabolites,
39	particularly flavonoids, in constraining and stimulating root-associated bacterial taxa.
40	
41	48. Kudjordjie EN, Sapkota R, Steffensen SK, Fomsgaard IS, Nicolaisen M: Maize synthesized
42	benzoxazinoids affect the host associated microbiome. Microbiome 2019, 7:59.
43	
44	* Using rRNA gene sequencing of microbial communities associated with below- and
45	aboveground tissues of wild-type maize and three different <i>bx</i> mutants, this study provides
46	a detailed characterisation the effects of BXs on microbial communities associated with the
47	rhizosphere, roots and shoot tissues of maize.

1	
2	49. Ahmad S, Veyrat N, Gordon-Weeks R, Zhang Y, Martin J, Smart L, Glauser G, Erb M, Flors
3	V, Frey M, et al.: Benzoxazinoid metabolites regulate innate immunity against
4	aphids and fungi in maize. Plant Physiol 2011, 157 :317-327.
5	50. Liu Y, Zhang N, Qiu M, Feng H, Vivanco JM, Shen Q, Zhang R: Enhanced rhizosphere
6	colonization of beneficial Bacillus amyloliquefaciens SQR9 by pathogen infection.
7	FEMS Microbiol Lett 2014. 353 :49-56.
8	51. Tan S. Yang C. Mei X. Shen S. Raza W. Shen O. Xu Y: The effect of organic acids from
9	tomato root exudates on rhizosphere colonization of Bacillus amyloliquefaciens T-
10	5 . Applied Soil Ecol 2013, 64 :15-22.
11	52. Ling N, Raza W, Ma J, Huang Q, Shen Q: Identification and role of organic acids in
12	watermelon root exudates for recruiting Paenibacillus polymyxa SQR-21 in the
13	rhizosphere. Eur J Soil Biol 2011, 47:374-379.
14	53. Yuan J, Zhang N, Huang Q, Raza W, Li R, Vivanco JM, Shen Q: Organic acids from root
15	exudates of banana help root colonization of PGPR strain Bacillus
16	amyloliquefaciens NJN-6. Sci Rep-UK 2015, 5.
17	54. Neal AL, Ahmad S, Gordon-Weeks R, Ton J: Benzoxazinoids in Root Exudates of Maize
18	Attract Pseudomonas putida to the Rhizosphere. PLOS One 2012, 7:e35498.
19	
20	* This study demonstrated that DIMBOA in root exudates of young maize seedlings induces
21	transcriptomic changes in plant-beneficial P. putida KT2440 bacteria, which are associated
22	with catabolism of aromatic metabolites, quorum sensing and chemotaxis. Using maize
23	genotypes affecting in DIMBOA production, the authors furthermore show that maize BX
24	production has a positive effect on rhizosphere colonization by <i>P. putida</i> KT2440.
25	
26	55. Saunders M, Kohn LM: Evidence for alteration of fungal endophyte community
27	assembly by host defense compounds. New Phytol 2009, 182:229-238.
28	56. Fan B, Carvalhais LC, Becker A, Fedoseyenko D, von Wiren N, Borriss R: Transcriptomic
29	profiling of Bacillus amyloliquefaciens FZB42 in response to maize root exudates.
30	BMC Microbiol 2012, 12 .
31	
32	* Transcriptomic analysis of <i>Bacillus amyloliquefaciens</i> FZB42 revealed that root exudates
33	activate bacterial genes associated with nutrient metabolism, motility, antibiotic production
34	and signalling.
35	
36	57. Matilla MA. Espinosa-Urgel M. Rodríguez-Herva JJ. Ramos JL. Ramos-González MI:
37	Genomic analysis reveals the major driving forces of bacterial life in the
38	rhizosphere. Genome Biol 2007. 8:R179.
39	58. Kierul K. Voigt B. Albrecht D. Chen XH. Carvalhais LC. Borriss R: Influence of root
40	exudates on the extracellular proteome of the plant growth-promoting bacterium
41	Bacillus amyloliquefaciens FZB42. Microbiol 2015. 161:131-147.
42	
43	* Proteomic analysis of <i>B. amyloliquefaciens</i> FZB42 revealed that root exudates increase
44	bacterial production of proteins associated with polymer degradation and bacterial stress.

1 2	59. Schenk ST, Stein E, Kogel K-H, Schikora A: Arabidopsis growth and defense are modulated by bacterial quorum sensing molecules. <i>Plant sig behav</i> 2012, 7 :178-
3	181.
4 5	60. Walker TS, Bais HP, Déziel E, Schweizer HP, Rahme LG, Fall R, Vivanco JM: <i>Pseudomonas</i> <i>aeruginosa-</i> Plant Root Interactions. Pathogenicity, Biofilm Formation, and Root
6	Exudation. Plant Physiol 2004, 134 :320-331.
/ 8	61. Corral-Lugo A, Daddaoua A, Ortega A, Espinosa-Urgel M, Krell T: Rosmarinic acid is a homoserine lactone mimic produced by plants that activates a bacterial quorum-
9	sensing regulator. Sci Signal 2016, 9:ra1.
10	** The systems are vide in vitre and in vive systems that as provining a side
11 12	The authors provide in vitro and in vivo evidence that rosmanning acid, a
12	plants triggers homosoring lastone dependent querum sensing responses in <i>P. geruginesa</i>
17	bacteria
14	Dactella.
16	62 Pierson IS Keppenne VD Wood DW: Phenazine antibiotic biosynthesis in
17	Pseudomonas aureofaciens 30-84 is regulated by PhzR in response to cell density
18	Bacteriol 1994. 176 :3966-3974.
19	63. Wood DW. Gong F. Davkin MM. Williams P. Pierson LS: N-acyl-homoserine lactone-
20	mediated regulation of phenazine gene expression by Pseudomonas aureofaciens
21	30-84 in the wheat rhizosphere . J Bacteriol 1997, 179 :7663-7670.
22	64. de Boer W, Hundscheid MPJ, Gunnewiek PJAK, de Ridder-Duine AS, Thion C, van Veen
23	JA, van der Wal A: Antifungal Rhizosphere Bacteria Can increase as Response to the
24	Presence of Saprotrophic Fungi. PLOS One 2015, 10.
25	65. Yuan J, Zhao J, Wen T, Zhao M, Li R, Goossens P, Huang Q, Bai Y, Vivanco JM, Kowalchuk
26	GA, et al.: Root exudates drive the soil-borne legacy of aboveground pathogen
27	infection. Microbiome 2018, 6 .
28	
29	** This study has characterised soil feedback responses to five successive generations of
30	Arabidopsis stress by <i>P. syringae</i> DC3000. The authors report that soil conditioned by
31	diseased plants induces resistance as a result from changes in the soil microbiome, which
32	are mediated by long chain organic acids.
33	
34	66. Berendsen RL, Vismans G, Yu K, Song Y, de Jonge R, Burgman WP, Burmolle M,
35	Herschend J, Bakker P, Pieterse CMJ: Disease-induced assemblage of a plant-
36	beneficial bacterial consortium. ISME J 2018, 12:1496-1507.
37	
38	** This study demonstrates that downy mildew-infected Arabidopsis conditions the soil for
39	resistance-inducing activity, providing protection against this pathogen in a second
40	population of plants. The authors furthermore show that the downy mildew-infected
41	Arabidopsis selects for three bacterial species in the rhizosphere, which interact
42	synergistically in biofilm formation and elicitation of induced systemic resistance.
43	
44	67. Carvalhais LC, Dennis PG, Badri DV, Kidd BN, Vivanco JM, Schenk PM: Linking Jasmonic
45	Acid Signaling, Root Exudates, and Rhizosphere Microbiomes. Mol Plant Microbe
46	Interact 2015, 28 :1049-1058.

1	68. Bruto M, Prigent-Combaret C, Muller D, Moënne-Loccoz Y: Analysis of genes
2	contributing to plant-beneficial functions in plant growth-promoting rhizobacteria
3	and related Proteobacteria. Sci Rep-UK 2014, 4:6261.
4	69. Pieterse CM, Zamioudis C, Berendsen RL, Weller DM, Van Wees SC, Bakker PA: Induced
5	systemic resistance by beneficial microbes. Annu Rev Phytopathol 2014, 52:347-
6	375.
7	70. Pieterse CM, van Wees SC, van Pelt JA, Knoester M, Laan R, Gerrits H, Weisbeek PJ, van
8	Loon LC: A novel signaling pathway controlling induced systemic resistance in
9	Arabidopsis. Plant Cell 1998, 10 :1571-1580.
10	71. Van der Ent S, Van Hulten M, Pozo MJ, Czechowski T, Udvardi MK, Pieterse CM, Ton J:
11	Priming of plant innate immunity by rhizobacteria and beta-aminobutyric acid:
12	differences and similarities in regulation. New Phytol 2009, 183:419-431.
13	72. Castrillo G, Teixeira PJ, Paredes SH, Law TF, de Lorenzo L, Feltcher ME, Finkel OM,
14	Breakfield NW, Mieczkowski P, Jones CD, et al.: Root microbiota drive direct
15	integration of phosphate stress and immunity. Nature 2017, 543:513-518.
16	
17	** This study demonstrates that rhizobacterial communities trigger a phosphate starvation
18	response (PSR) in Arabidopsis, which controlled by the transcriptional regulator PHR1 and its
19	weakly redundant paralogue, PHL1. Both regulators induce microbome-enhanced responses
20	to phosphate limitation and modulate SA-dependent immunity in the host.
21	
22	73. Khan GA, Vogiatzaki E, Glauser G, Poirier Y: Phosphate Deficiency Induces the
23	Jasmonate Pathway and Enhances Resistance to Insect Herbivory. Plant Physiol
24	2016, 171 :632-644.
25	74. Pant BD, Pant P, Erban A, Huhman D, Kopka J, Scheible WR: Identification of primary
26	and secondary metabolites with phosphorus status-dependent abundance in
27	Arabidopsis, and of the transcription factor PHR1 as a major regulator of metabolic
28	changes during phosphorus limitation. Plant Cell Environ 2015, 38:172-187.
29	75. Miransari M: Interactions between arbuscular mycorrhizal fungi and soil bacteria. Appl
30	Microbiol Biotechnol 2011, 89 :917-930.
31	76. Cameron DD, Neal AL, van Wees SC, Ton J: Mycorrhiza-induced resistance: more than
32	the sum of its parts? Trends Plant Sci 2013, 18:539-545.
33	77. Stringlis IA, Yu K, Feussner K, de Jonge R, Van Bentum S, Van Verk MC, Berendsen RL,
34	Bakker P, Feussner I, Pieterse CMJ: MYB72-dependent coumarin exudation shapes
35	root microbiome assembly to promote plant health. Proc Natl Acad Sci USA 2018,
36	115 :E5213-E5222.
37	
38	** This detailed study of the Arabidopsis – P. simae WCS417 interaction revealed that the
39	ISR-regulatory genes MYB72 and BGLU42 induce an iron-starvation response in the plant,
40	which in turn increases root exudation of the anti-micorbial and iron-chelating metabolite
41	scopoletin that influence root-associated microbial populations.
42	
43	78. Voges M, Bai Y, Schulze-Lefert P, Sattely ES: Plant-derived coumarins shape the
44	composition of an Arabidopsis synthetic root microbiome. Proc Natl Acad Sci U S A
45	2019, 116 :12558-12565.
46	

- 1 ** Using a gnotobiotic growth system of Arabidopsis and a 22-member synthetic community 2 of soil bacteria, the authors screened different Arabidopsis mutants for impacts on bacterial 3 root colonization. The results showed that redox-active scopoletin derivates, such as 4 esculetin, fraxetin, and sideretin, have specific impacts on selected members of the 5 synthetic rhizobiome community. 6 7 79. Robert CA, Veyrat N, Glauser G, Marti G, Doyen GR, Villard N, Gaillard MD, Kollner TG, 8 Giron D, Body M, et al.: A specialist root herbivore exploits defensive metabolites 9 to locate nutritious tissues. Ecol Lett 2012, 15:55-64.
- 80. Lopez-Raez JA, Shirasu K, Foo E: Strigolactones in Plant Interactions with Beneficial and
 Detrimental Organisms: The Yin and Yang. Trends Plant Sci 2017, 22:527-537.
- 81. Berg G, Köberl M, Rybakova D, Müller H, Grosch R, Smalla K: Plant microbial diversity is
 suggested as the key to future biocontrol and health trends. FEMS Microbiol Ecol
 2017, 93.
- 82. Hartman K, van der Heijden MGA, Wittwer RA, Banerjee S, Walser JC, Schlaeppi K:
 Cropping practices manipulate abundance patterns of root and soil microbiome
 members paving the way to smart farming. *Microbiome* 2018, 6:14.
- ** The authors have compared soil- and root-associated microbial communities of wheat
 under conventional and organic field management that varied in agrochemical input and
- tillage. The study discovered that 10% of the variation in microbial communities is affected
- 21 by field management. The study concludes that future agricultural strategies should
- 22 consider the soil microbiome to foster specific microbiota with agricultural benefits.



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Figure 2: Simplified schemes of the plant shikimate (A) and terpenoid (B) biosynthesis pathways, which generate stress-inducible secondary metabolites in roots with anti-microbial and/or semio-chemical activities [22-24]. Coloured boxes show representative examples of compounds within each class.



Figure 3: emerging roles for phosphate (purple) and iron (orange) starvation responses in the orchestration of diseasesuppressive mechanisms in the microbial biosphere of the plant. The model is based on recent evidence for reciprocal signalling events between nutrient starvation signalling in the host, systemic immune responses (ISR), and diseasesuppressing activities by the root- and soil-associated microbiome [67**, 72**).