Interaction between endophytic Proteobacteria strains and *Serendipita indica* enhances biocontrol activity against fungal pathogens

Alejandro del Barrio-Duque · Abdul Samad · Ole Nybroe · Livio Antonielli · Angela Sessitsch · Stéphane Compant

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Abstract

**Aims** Plants host communities of fungal and bacterial endophytes, establishing a complex network of multipartite interactions, but the mechanisms whereby they interact are poorly understood. Some fungi, such as the beneficial mycorrhiza-like fungus *Serendipita (=Piriformospora) indica*, can be helped by bacteria for establishment, survival and colonization. Although this fungus harbors a *Rhizobium* as an endofungal bacterium, we hypothesized that other bacteria might also establish associations with the fungus and combining *S. indica* with bacteria might enhance plant growth and health.

**Methods** The interactions among *S. indica* and four endophytic Proteobacteria belonging to *Methylobacterium*, *Tardiphaga*, *Rhodanobacter* and *Trinickia* spp. were characterized in vitro and for their effect on tomato growth and biocontrol of *Fusarium oxysporum* and *Rhizoctonia solani*. Possible mechanisms behind these interactions were described based on genome and microscopic analyses, using fungal and bacterial strains tagged with fluorescent markers.

**Results** All bacteria stimulated *S. indica* growth in vitro. Moreover, several of the bacteria stimulated growth of tomato plants, but co-inoculations with *S. indica* and bacteria did not perform better than single inoculations. Contrarily, combinations of *S. indica* and bacteria significantly reduced disease progression of fungal pathogens. These microbes seem to cooperate in the process of root colonization for instance by increasing fungal sporulation and hyphae expansion, showing multipartite interaction between microbes and plants. Interestingly, the strain of *Trinickia* internally colonizes spores of *S. indica* as an endofungal bacterium during in vitro-co-culturing, suggesting further that the fungus might acquire formerly unrecognized genera of bacteria and genome analysis of the bacteria revealed many genes potentially involved in fungal and plant growth stimulation, biocontrol and root colonization, highlighting putative mechanisms of plant-fungal-bacterial interaction.

**Conclusions** Our study represents an important step towards unraveling the complex interactions among plants, *S. indica*, endophytic bacteria and fungal pathogens, and indicates that adding bacteria to fungal inoculum could have a remarkable impact on the plant-*S. indica* symbiosis.

**Keywords** *Serendipita indica* · Bacterial endophytes · Bacterial genome · Tripartite interactions · Biocontrol · Endofungal bacterium
Introduction

Arbuscular mycorrhizal fungi (AMF) and rhizobacteria are among the most abundant components of the plant microbiota, and hence crucial drivers of plant evolution. They thrive in the plant rhizosphere in all natural and managed ecosystems but can also penetrate the roots and establish as endophytes (Compan et al. 2005b). In both niches, these microorganisms can improve plant nutrition, increase tolerance to abiotic stresses and enhance protection against phytopathogens (Bonfante and Anca 2009; Martínez-Arias et al. 2019). Some bacteria are capable of boosting colonization and establishment of endophytic and mycorrhizal fungi, and thus they have been regarded as mycorrhizal helper bacteria (Garbaye 1994; Frey-Klett et al. 2007). The physical connection between them can range from apparently disordered microbial assemblies to specific symbiotic associations of fungal hyphae and bacterial cells (Frey-Klett et al. 2011). Already in 1970, endofungal bacteria were detected living inside hyphae and spores of the AM fungus Endogone (Mosse 1970). Since then, several genera of bacteria have been identified residing inside endomycorrhizal fungi (Bianciotto et al. 1996; Hoffman and Arnold 2010) and seem to play a crucial role in the mycorrhizal symbiosis with plants (Partida-Martínez and Hertweck 2005; Salvioli et al. 2016). However, the complexity of the fungal-bacterial-plant interactions is still far from being fully elucidated. Understanding the functioning of the plant microbiome is an essential step for the exploitation of beneficial microorganisms, both fungi and bacteria, for the improvement of crop productivity and biological control of plant pathogens.

Serendipita indica, formerly known as Piriformospora indica, is an AM-like fungus originally isolated in the Indian Thar desert (Verma et al. 1998). It can endophytically colonize a wide range of plants, but unlike mycorrhizal fungi, this fungus depends on the host cell death for successful colonization (Deshmukh et al. 2006). S. indica promotes plant growth, induces tolerance to abiotic stresses, and confer resistance to shoot and root pathogens (Waller et al. 2005) by hormone-manipulation of the host plant, enhancement of induced systemic resistance (ISR) and activation of the antioxidative system (reviewed in Franken 2012; and Gill et al. 2016). Intriguingly, this fungus hosts an endofungal bacterium, Rhizobium radiobacter F4 (Sharma et al. 2008). In comparison to obligate endofungal bacteria, the genome of RrF4 is not reduced and the bacterium is able to live independently of its host and to thrive as an endophyte, thus forming a facultative symbiosis with the fungus (Glaeser et al. 2016). It hints at the interaction is not specific and other bacteria might also colonize and establish a facultative symbiosis with S. indica. Moreover, the endofungal bacterium RrF4 possesses plant growth-promoting properties and the ability to confer resistance to fungal plant pathogens. Therefore, it has been hypothesized that some of the beneficial properties attributed to S. indica might stem from the effect exerted by the bacterium (Qiang et al. 2012a; Glaeser et al. 2016).

It appears likely that other bacteria with plant growth-promoting (PGP) traits might also contribute to further increase fungal fitness and the PGP and biocontrol effects reported by this fungus. For instance, the model bacterium Bacillus subtilis and the strain WR5 of Azotobacter chroococcum were shown to enhance the growth and sporulation of S. indica in vitro (Jiang et al. 2018; Bhuyan et al. 2015). Some researchers have also investigated the effect of combining S. indica with plant growth-promoting rhizobacteria (PGPR) on plant growth: co-inoculation of S. indica with the phosphate-solubilizing bacterium Pseudomonas striata increased plant growth in chickpea (Meena et al. 2010), and a talcum-based formulation of S. indica and Pseudomonas sp. R81 enhanced plant biomass in tomato (Sarma et al. 2011). However, it has also been reported that dual inoculations of S. indica with Paenibacillus or pseudomonads did not improve (Nautiyal et al. 2010) or even displayed lower plant dry weight than single inoculations (Sarma et al. 2011). One way of improving biocontrol may be to combine microbes, especially if they exhibit complementary modes of action and abilities to colonize root microsites (Whipps 2001). Although some combinations of bacteria and fungi have demonstrated to enhance biocontrol (Guetsky et al. 2001; Sarma et al. 2011), there are several studies in which combinations of various bacteria and fungi provided poorer biocontrol activities than the microbes used singly (Hervás et al. 1998; Xu et al. 2011). In light of these incompatibilities, there is a need to further investigate new fungal-bacterial synergisms and the mechanisms involved in this interplay for an efficient exploitation and engineering of the plant microbiome.

In a previous study, we detected several endophytic bacterial strains from very diverse families with the ability to stimulate S. indica growth in vitro. In that
study, we focused on Mycolicibacterium strains, as this genus showed to comprise the highest number of strains having a beneficial effect on S. indica (del Barrio-Duque et al. 2019). However, strains from other taxa might have similar beneficial properties on S. indica but potentially employing different mechanisms. Among those bacteria showing the most outstanding stimulatory effects on S. indica growth, we selected four proteobacterial strains of Methylobacterium sp., Tardiphaga sp., Rhodanobacter sp., and Trinickia (=Paraburkholderia) sp. The uniquely reported endofungal bacterium of S. indica (RrF4) belongs to the proteobacterial order Rhizobiales (Sharma et al. 2008), and perhaps other bacteria belonging to Rhizobiales, like Methylobacterium or Tardiphaga, might also establish a synergistic interaction with S. indica. Members of the order Burkholderiales (e.g. Trinickia) are among the most frequently identified endofungal bacteria (Compant et al. 2008). Bacteria of the genus Rhodanobacter (Xanthomonadales) seem further to be implicated in shaping communities of AMF (Svenningsen et al. 2018) and several members of the same order have been considered as ‘fungiphile’ bacteria (Simon et al. 2015). These features make the selected strains potential candidates for studying fungal-bacterial interactions. Moreover, the four selected strains are phylogenetically closely related to other strains of PGPR described to enhance plant growth or to be involved in the biocontrol of pathogens (Tani et al. 2015; Berg 2009; De Clercq et al. 2006; Sessitsch et al. 2005). We therefore hypothesized that strains belonging to these taxa may also influence the fitness, establishment and colonzation of S. indica and enhance the effects of this symbiotic fungus on plant growth and biocontrol. We further speculated that they might internally colonize S. indica and behave as endofungal bacteria.

Materials and methods

Microorganisms

_Serendipita (=Piriformospora) indica_ strain DSM 11827 (Varma et al. 1999) was provided by Prof. Philipp Franken (Erfurt Research Centre for Horticultural Crops, Germany). The fungus was kept at −80 °C in a mixture 3:1 of Potato Dextrose Broth (PDB) (Carl Roth, Germany) and glycerol, and grown on Potato Dextrose Agar (PDA) plates or in liquid culture containing _Aspergillus_ complete medium (Pontecorvo et al. 1953).

_Rhizoctonia solani_ AG-3 was provided by Dr. Rosanna C. Hennessy (University of Copenhagen, Denmark) and _Fusarium oxysporum_ f.sp. _lycopersici_ Fol4287 by Maria E. Constantin (University of Amsterdam, Netherlands) (Di Pietro and Roncero 1996). Both fungal pathogens were maintained at −80 °C in a mixture of 3:1 PDB + glycerol.

The endophytic bacterial strains Methylobacterium sp. P1–11, Tardiphaga sp. P9–11, Rhodanobacter sp. T12–5, and Trinickia sp. T12–10 were obtained from the AIT (Austrian Institute of Technology, Tulln, Austria) strain collection. They were isolated from tomato (T) or potato (P) roots (del Barrio-Duque et al. 2019).

Effect of bacterial strains on mycelial growth of _S. indica_ and fungal pathogens

The growth of the beneficial fungus _S. indica_ or the pathogens _F. oxysporum_ and _R. solani_ in interaction with bacteria was assessed by measuring hyphae development on agar plates as described in del Barrio-Duque et al. (2019). For these experiments, bacteria were precultured for four days on Nutrient Agar No2 (NA) (Sigma-Aldrich, St. Louis, USA) and then streaked individually on 1 cm² in the center of a Petri dish (9 cm Ø) containing 15 ml of PDA or a mixture 1:1 of PDA + NA. _S. indica_ was precultured for two weeks on PDA and then a 0.5 cm² agar plug of active mycelium was placed upside down onto the streaked bacterium. Additionally, _S. indica_ was co-cultured on PDA with bacteria streaked 4 cm away from the agar plug. All the co-cultures were replicated four times. As a control, the fungus was grown alone. Plates were incubated in darkness at 26 °C and mycelial growth was monitored after 3, 6, 9 and 11 days of dual-culturing, depending on the growth rate of each fungus. The surface of the Petri dish covered with mycelium was then quantified with ImageJ 1.48 software (https://imagej.nih.gov/ij/).

For estimation of dry fungal biomass and spore yield during interactions with bacteria, seven PDA plugs from the periphery of a two-week old culture of _S. indica_ were transferred to 100 ml Erlenmeyer flasks with 25 ml PDB and incubated at 26 °C and 150 rpm. After six days, 1 ml of a bacterial culture grown on Nutrient Broth (NB) (Difco, Detroit, MI) for two days (OD 0.3 at 630 nm) was added to each flask and co-cultured for additional four days. Mycelium was then washed thrice.
with distilled water and collected by centrifugation (Thermo Scientific Heraeus Megafuge 40R) at 4600 rpm for 5 min. Dry fungal weight was determined after drying the mycelium in the oven at 60 °C for three days. Spore production was estimated in a hemocytometer after grinding and sonicating the fungal pellet according to Kumar et al. (2011).

Co-inoculation of *S. indica* and endophytic bacteria for tomato growth promotion

To produce fungal inoculum, agar plugs from the active mycelium of a 2-week old PDA culture of *S. indica* were transferred to 250 ml Erlenmeyer flasks containing 100 ml of *Aspergillus* complete medium and incubated at 26 °C for 3 weeks at 150 rpm in a shaking incubator. Mycelium and spores were harvested by centrifugation (4500 rpm, 5 min) and washed 3 to 5 times with sterile phosphate-buffered saline (PBS) of pH 6.5. The mixture of mycelium and spores was crushed with a homogenizer Ultra Turrax T25 (IKA®, Staufen, Germany) in intervals of 30 s for 3 min. Spores + mycelium fragments were measured with a hemocytometer (NanoEnTek, Seoul, Korea) and the viability of the spores + mycelium fragments was confirmed by plating on PDA. Final inoculum ratios were adjusted with PBS.

For bacteria, they were grown for 60 h in round-bottom Falcon tubes containing 5 ml of NB (26 °C, 190 rpm), then centrifuged (4600 rpm, 6 min, room temperature) and washed thrice with sterile PBS to discard traces of media. Bacterial growth and CFU were preliminarily assessed by measuring the optical density at 630 nm and plating standard serial dilutions on NA, then the amount of inoculum was calibrated to the final concentration of bacterial cells with PBS.

For plant experiments, tomato seeds cv. “Moneymaker” (Austrosaat, Vienna, Austria) were germinated in a Whatman® filter paper (110 mm Ø) for 4 days and then transferred to Falcon tubes (50 ml) containing 15 ml of PBS and either i) spores + hyphae fragments of *S. indica* (5 x 10⁵ CFU/ml), ii) bacteria (5 x 10⁷ CFU/ml) or iii) a blend of *S. indica* (5 x 10⁵ CFU/ml) with bacteria (5 x 10⁷ CFU/ml). Falcon tubes were kept for 30 min in a horizontal Tube Roller RS-TR5 (Phoenix Instrument GmbH, Garbsen, Germany). Control seeds were immersed in 1xPBS. Seeds were then sown in 1-l pots containing the substrate “Fruhstorfer Erde Typ Nullerde” (Hawita Gruppe, Vechta, Germany) at 1 cm depth. The plants were grown in the greenhouse with a 12 h light/dark photoperiod, day/night temperature of 22/21 °C, and a relative humidity 50/35%, and watered twice a week with tap water. Plants were harvested 6 weeks after planting and shoot fresh weight and leaf area were measured (using ImageJ software). The experiment consisted of 10 treatments: ± *S. indica* and ± bacteria (four strains) with 14 replicates (7 pots × 2 plants per pot) for each treatment. Additionally, the experiment was repeated with a high-nutrient soil composed of a mixture (1:1:1 v/v) of substrate “Tonsubstrat ED63 Special” (Einheitserde, Germany), perlite and sand. After harvesting, shoot fresh and dry weight (after oven-drying for 3 days at 70 °C) were measured.

Biocontrol of *F. oxysporum* and *R. solani*

Protection against *F. oxysporum* was determined by pot experiments in which tomato plants (cv. Moneymaker) were infected with the pathogen and with single or combined inoculations of *S. indica* and bacteria. *Fusarium* inoculum was produced according to Constantin et al. (2019). For this, a PDA plug from a 6-day old culture of *Fusarium* was transferred to 100 ml minimal media (3% sucrose, 1% KNO₃ and 0.17% Yeast Nitrogen Base without ammonia and amino acids) and incubated for 5 days in the dark (26 °C, 190 rpm). Spores were separated with a Miracloth filter (Millipore), rinsed twice with sterile PBS and adjusted to 10⁷ spores/ml. Bacterial and *S. indica* inoculum production as well as seed inoculation were performed as explained above. Inoculated seeds were then planted in 1 l pots containing a mixture (1:1:1 v/v) of the substrate “Tonsubstrat ED63 Special” (Einheitserde, Germany), perlite and sand. Ten replicated plants were assessed per treatment (5 pots × 2 seeds per pot). Ten-day-old tomato seedlings were uprooted, trimmed (leaving around 1 cm of root) to assist Fusarium infection (de Lamo et al. 2018) and placed in the Fusarium spore suspension for 30 min, and then immediately repotted. After five weeks, shoot fresh weight and the extent of disease progression were measured. Disease index (DI) was scored as described before (Gawehns et al. 2014) but adding DI = 5 when plants were dead. Briefly, DI 0 = no symptoms; 1 = one brown vessel above the soil; 2 = one-two brown vascular bundles at the cotyledon level; 3 = three or more brown vessels and growth deformation, 4 = all vessels brown or the plant is dwarfed and wilted, 5 = dead plant.

The effects of combinations of individual bacteria and *S. indica* against *R. solani* were determined by a
germination test in sterile closed systems. Transparent Steri Vent Containers (107 × 94 × 96 mm, Duchefa Biochemie b.v, Haarlem, Netherlands) were filled with 120 g of a sterile (2x, 20 min, 121 °C) mixture (4:1, w/w) of distilled water and vermiculite (2–3 mm, Sigma-Aldrich). Tomato seeds were surface-sterilized for 5 min with 2.5% sodium hypochlorite and washed eight times with sterile water. Seeds were inoculated with ± S. indica and ± individual bacterial strains as earlier described. Five 5 mm plugs of a 3-week-old culture of Rhizoctonia grown in 1/5 PDA were placed in a row in the center of each container, and 2 rows of tomato seeds (5 seeds per row) were sown in both sides of the pathogen row 2 cm away from the pathogen. Control was prepared with 1/5 PDA plugs. The seeds and the Rhizoctonia agar plugs remained at 5 mm depth. Three replicated containers were included per treatment and kept in the greenhouse (see above) for 4 months. Germination and disease progression of the seedlings were monitored regularly and assessed by counting seedlings as follows: 1 = Plant germinated and no Rhizoctonia symptoms; 0.5 = Plant germinated and alive, but with necrotic areas in stem and leaves; 0 = dead plant.

Bacterial genome sequencing, assembly and annotation

To determine genome contents of the four Proteobacteria strains, bacterial genomic DNA was extracted with a phenol-chloroform based protocol. Briefly, bacteria were cultivated for 3 days in NB and harvested by centrifugation. The bacterial pellets were then resuspended in lysis buffer (0.2 mg/mL. Proteinase K, 50 mM Tris-Cl, 1% SDS, 5 mM EDTA pH 8, 0.5 M NaCl) and incubated overnight (65 °C, 400 rpm). DNA was extracted 2 times using 1 volume of phenol-chloroform-isooamylalcohol (25:24:1) and collected by centrifugation (5000 rpm, 2 min). Genomic DNA was then purified with Amicon Ultra 0.5 mL 30 K Centrifugal Filter Units (Millipore, Cork, Ireland) and resuspended in water. Whole-genome shotgun sequencing was performed on an Illumina HiSeq (GATC Biotech, Konstanz, Germany), producing 2 × 150 bp reads.

Illumina reads were screened for the presence of PhiX using Bowtie 2 (v2.3.4.3) (Langmead and Salzberg 2012) and adapters were trimmed with fastp (v0.19.5) (Chen et al. 2018b). Sequence length distribution and quality were checked via FastQC (Andrews 2010). Genome assembly was performed with SPAdes v3.13.0 (Bankevich et al. 2012) and low-abundant (< 2x), short (< 500 bp) contigs discarded. Contaminant contigs were examined using BlobTools and alien contigs were eventually filtered out. Genome assembly quality was inferred using QualiMap v2.2 (Okonechnikov et al. 2015) and QUAST v5.0.0 (Gurevich et al. 2013) and then genome completeness reconstruction was evaluated with CheckM v1.0.18 (Parks et al. 2015). Gene annotation was carried out using Prokka v1.12 (Seemann 2014) and NCBI Prokaryotic Genome Annotation Pipeline (PGAP) (Tatusova et al. 2016). The presence of plasmids was ascertained by using Mash v2.1 against the PLSDB database (Galata et al. 2018). Putative plasmid contigs were screened for the presence of genes for replication initiator proteins, repA. For this purpose, a curated FASTA file with ~8000 repA genes was generated from plasmidial genome sequences in NCBI. These repA gene sequences were used to build a database against which the selected contigs were BLASTed. Functional annotation was performed using the ClassicRAST (Rapid Annotation using Subsystem Technology) web server (http://rast.nmpdr.org) (Aziz et al. 2008) and the hierarchical orthology framework EggNOG 4.5 (Huerta-Cepas et al. 2015). CAZy families were ascertained with dbCAN2 based on the DIAMOND database (Galata et al. 2018). CAZy families were ascertained with dbCAN2 based on the DIAMOND database (Galata et al. 2018). Putative plasmid contigs were screened for the presence of genes for replication initiator proteins, repA. For this purpose, a curated FASTA file with ~8000 repA genes was generated from plasmidial genome sequences in NCBI. These repA gene sequences were used to build a database against which the selected contigs were BLASTed. Functional annotation was performed using the ClassicRAST (Rapid Annotation using Subsystem Technology) web server (http://rast.nmpdr.org) (Aziz et al. 2008) and the hierarchical orthology framework EggNOG 4.5 (Huerta-Cepas et al. 2015). CAZy families were ascertained with dbCAN2 based on the DIAMOND database (Galata et al. 2018).
draft genome sequences of the strains P1–11, P9–11, T12–5 and T12–10 are available at NCBI, BioProject PRJNA393298, with the DDBJ/ENA/GenBank accession numbers NPKS00000000, NPKQ00000000, NPNK00000000 and NPKM00000000 respectively.

Quantification of root colonization by *S. indica*

To study the effect of bacteria on the colonization of the roots by *S. indica*, tomato seedlings were grown in Transparent Steri Vent Containers as described above, but without addition of *Rhizoctonia* agar plugs, and inoculated with *S. indica* alone and in combination with bacterial strains P1–11 or T12–10 wild-types as earlier described. For every treatment, 3 seedlings from each transparent (×3) container were sampled 10 days and 8 weeks after planting and maintained at −20 °C. Roots were stained according to a modified protocol of Venneman et al. (2017). Briefly, roots were abundantly rinsed under running tap water and incubated at 80 °C in 10% KOH solution for 30 min, and then at room temperature for an additional 30 min. After a rinsing step, roots were acidified for 20 min at room temperature in 1% acetate solution before adding droplets of china blue ink (2 droplets per 25 ml). This mixture was incubated for 30 min at 80 °C and at room temperature for an additional 30 min. The roots were then rinsed with water and immersed in a glycerol solution for 10 min (140 ml glycerol +46 ml water +14 ml 1% acetate) before to be rinsed again. From every stained root, fragments of 1 cm length were cut and observed with a microscope Nikon Eclipse E200, and the presence of hyphae and spores of *S. indica* were quantified. In total 5 to 7 plantlets per treatment were analyzed to have 20 root sections to quantify.

Fluorescent labeling of the endophytic bacteria

The four strains were tagged with the red fluorescent marker mCherry by transformation with the self-replicating broad-host-range plasmid pSEVA237R (Silva-Rocha et al. 2013). This plasmid does not contain transposons and is not inserted chromosomally but can replicate itself in the recipient cell. Wild-type strains were grown in 10 ml of a PDB + NB mixture (1:1) in 50 ml Falcon tubes for 24 h in a shaking incubator (28 °C, 180 rpm). Cells were harvested by centrifugation (4700 rpm, 10 min, 4 °C), washed thrice with 300 mM sucrose, and then resuspended in 400 μl of 300 mM sucrose. One hundred μl of the competent cells were mixed with 500 ng of purified pSEVA237R plasmid. The mixture was electroporated with a Gene Pulser II (Bio-Rad, Richmond, Calif.) by a pulse of 25 μF, 200 Ω, 2.5 kV, and then 1 ml of PDB + NB was immediately added to the mixture. The mixture was incubated for 2 h at 28 °C in a shaking incubator and then plated on PDA + NA amended with 50 μg/ml kanamycin (Km). Colonies and cells of the mCherry-marked strains were examined by using a fluorescence microscope (Olympus model SZX16; Tokyo, Japan). Transformant stability was assessed by culturing tagged strains in PDB + NB with or without kanamycin for over five generations, and then plating cells on PDA + NA w/o antibiotics. Moreover, the growth pattern as well as colony and cell morphologies of wild-type and mCherry-tagged bacteria were compared.

Confocal laser scanning microscopy (CLSM) analysis

CLSM assays were performed with GFP-labelled *S. indica* (gift from Prof. Alga Zuccaro, University of Cologne, Germany) and the red fluorescent-labelled bacterial strains *Methylobacterium* sp. P1–11:mCherry and *Trinickia* sp. T12–10:mCherry that were successfully transformed. Interactions between the fungus and each bacterium were examined in Petri dishes and on roots. For the in vitro analysis, *S. indica* and bacterial strains were dual-cultured in direct contact on PDA as earlier described. No antibiotics were included for the bacterial preculturing. After 11 days of co-culturing, fragments of fungal mycelium were sampled with a scalpel from the surface of the agar plate, roughly 0.5 cm away from the spot at which the fungal agar plug and the bacteria were initially confronted. For each bacterial-fungal combination, samples were retrieved and observed from four replicated Petri dishes.

Aiming at investigating root colonization patterns of *S. indica* and bacteria, tomato seedlings were grown in Transparent Steri Vent Containers as described above. Inoculum production of transformed microbes was obtained with the same protocol as for the wild type strains, but to avoid loss of the plasmid-born fluorescence, the strain *Methylobacterium* sp. P1–11:mCherry (but not *Trinickia* sp. T12–10:mCherry) was grown on NB + 50 μg/ml Km. The seedlings were harvested 14 dap. For every treatment (+ *S. indica* and + individual bacterial strains) at least 3 seedlings from each transparent (×3) container were sampled and observed.
The samples were then examined under a confocal microscope (Olympus Fluoview FV1000 with multiline laser FV5-LAMAR-2 HeNe(G) and laser FV10-LAHEG230–2). X, Y, Z pictures were acquired with 20X, 40X or 60X objectives at 405, 488, 594 nm wavelengths and then merged (RGB) using the software of the microscope provider (Olympus FV1000). Confocal stacks are presented as single optical slices or maximum projections (Cardinale and Berg 2015). The light/contrast balance was improved to better observe the image details when image acquisition remained very dark (as described in Glassner et al. 2015). Three-dimensional reconstructions (Cardinale and Berg 2015) were modeled using the software Imaris® 7.2 (Bitplane AG, Zürich, Switzerland).

Statistical analysis

The data on in vitro bacterial-fungal dual-inoculations as well as plant biomass and leaf area from the growth enhancement experiments were analyzed in R 3.5.1 (Team RC 2019). Data distributions were evaluated with the fitdistrplus package (Delignette-Muller and Dutang 2015) and linear or linear mixed-effects models (nlme R package) (Pinheiro et al. 2019), when necessary, were created. After graphical validation of homogeneity assumption, ANOVA test was carried out on the generated models, followed by pairwise comparisons (Tukey’s method, \( P = 0.05 \)) calculated by Estimated Marginal Means (emmeans R package) (Lenth 2019). Quantitative data were processed with dplyr package (Wickham et al. 2019) and displayed with boxplots using ggplot2 package (Wickham and Chang 2016). The data from the biocontrol experiments were analyzed using PRISM 8.0 (GraphPad, https://www.graphpad.com). Datasets from the germination test with Rhizoctonia were analyzed using one-way ANOVA and Tukey’s test \( (P = 0.05) \). Regarding the Fusarium assay, a non-parametric Mann–Whitney \( U \)-test was applied on the shoot fresh weight and disease index data (de Lamo et al. 2018; Constantin et al. 2019).

Results

Effect of bacterial strains on mycelial growth of \( S. \) indica and fungal pathogens

The four tested strains, Methylobacterium sp. P1–11, Tardiphaga sp. P9–11, Rhodanobacter sp. T12–5 and Trinickia sp. T12–10, significantly stimulated \( S. \) indica growth when co-cultured on PDA for all the measurements of hyphae expansion over the Petri dish (Fig. 1a–c and Suppl. Fig. 1). The stimulating effect was also observed on PDA + NA (Suppl. Fig. 1). When \( S. \) indica and the bacterial strain were placed at different spots of the Petri dish, the fungal growth was not impaired; rather the strains P1–11, P9–11 and T12–5 significantly promoted hyphal extension (Suppl. Fig. 1). In the experiments on liquid medium, strains P1–11 and T12–10 significantly increased fungal dry weight and sporulation (Fig. 1d–f). Strains P9–11 and T12–5 only exhibited a moderate non-significant increase in fungal biomass and spore yield, probably due to their slow growth in PDB.

In contrast to the observed growth stimulation of \( S. \) indica, mycelial growth of the phytopathogen \( F. \) oxysporum and \( R. \) solani was not stimulated by bacteria (Suppl. Fig. 2). In contrast, the strains P1–11 on PDA, and T12–5 on PDA and PDA + NA, significantly reduced Fusarium growth, while the strain P9–11 significantly impaired the growth of Rhizoctonia on PDA + NA.

Co-inoculation of \( S. \) indica and endophytic bacteria for tomato growth promotion

Tomato plants received single or dual inoculations of \( S. \) indica and the bacterial strains. In no case microbial application impaired plant growth (Fig. 2a–c). Strain P1–11 only exhibited neutral effects and strain P9–11 yielded non-significant 2.0- and 4.2-fold increments of shoot fresh weight and leaf area respectively (Fig. 2b and c). Strains T12–5 and T12–10 significantly promoted plant growth, exhibiting 3.9- and 4.0-fold increments in shoot fresh weight, respectively (Fig. 2b). Inoculation with \( S. \) indica resulted also in notable increased fresh weight and leaf area. Dual inoculations of \( S. \) indica and bacteria significantly enhanced plant growth in comparison to control plants, however these combinations did not perform better than single inoculations. Interestingly, dual inoculation of \( S. \) indica + T12–5 displayed significant lower performance than inoculation of either microbe alone. This greenhouse trial was repeated in soil with a higher content of nutrients, but under these conditions no significant differences were detected between treatments of control plants and plants inoculated with microbes (Suppl. Fig. 3).
Biocontrol of *F. oxysporum* and *R. solani*

The tomato plants treated with *F. oxysporum* showed typical symptoms as leaf yellowing, defoliation, stunting, wilting, necrotized xylem bundles, and death (Suppl. Fig. 4). Inoculations with the bacterial strains individually did not protect plants against the pathogen, except for strain T12–10, that significantly reduced the *Fusarium* disease index (Fig. 3a). Although not significantly, the magnitude of disease symptoms seemed to be mitigated when the plants were inoculated singly with the beneficial endophyte *S. indica*. Intriguingly, *Fusarium* symptoms (DI) were only significantly reduced when the plants were dual-inoculated with *S. indica* + P1–11, *S. indica* + P9–11 and *S. indica* + T12–10. The negative effect of *Fusarium* wilt on fresh weight was not significantly mitigated upon dual inoculation of *S. indica* + bacteria (Fig. 3b).

In the case of *Rhizoctonia* damping-off, the symptoms were first recorded 11 days after planting (dap). Single inoculation of bacteria did not restrain pathogen attack (Fig. 3c). Inoculation of seeds with *S. indica* typically conferred resistance against the pathogen, but the degree of plant protection in comparison to *Rhizoctonia* control was significant throughout the whole experiment exclusively when *S. indica* was co-inoculated with bacteria. More specifically, treating the plants with *S. indica* plus T12–10 significantly increased protection in comparison to the treatment with *S. indica* alone.

Effects of bacteria on root colonization by *S. indica*

In general, the extent of root colonization by *S. indica* was low, although similar results have been reported in tomato plants when *S. indica* was grown under the same growing conditions (Andrade-Linares et al. 2013). Ten days after planting, addition of *Methylobacterium* P1–
and Trinickia T12–10 to the S. indica inoculum significantly increased the extent of root colonization in comparison to single inoculations (Figs. 4a-b). Eight weeks after planting, inoculation of S. indica with T12–10 significantly increased the number of hyphae aggregates respect to control plants (Fig. 4c). However, in all the sampled root fragments we detected the presence of the fungus and there were no differences in terms of root areas colonized by S. indica among the different treatments 8 weeks after planting (Fig. 4d). Regarding the number of spores inside the roots we observed a modest increment in plants dual-inoculated with S. indica and T12–10 (Suppl. Fig. 5). The number of spores detected in the treatment S. indica + T12–10 was significantly increased when compared to the treatment S. indica + P1–11, but not when compared to control plants.

However, more hyphae producing spores were detected in S. indica + T12–10 compared to control (data not shown).

Stability of the mCherry-marked strains and comparison with the wild-type strain

The growth patterns and the colony and cell morphologies of the mCherry-tagged strains of Methylobacterium sp. P1–11 and Trinickia sp. T12–10 on PDA + NA w/o antibiotics were similar to those of the wild-type strains, but the colonies displayed a characteristic purple color (Suppl. Fig. 6a,b). Colonies and cells of these mCherry-marked strains were notably fluorescent under orange/red light when cultured on PDA + NA amended with kanamycin (50 μg/ml). When
Fig. 3 Effect of strains *Methylobacterium* sp. P1–11, *Tardiphaga* sp. P9–11, *Rhodanobacter* sp. T12–5 and *Trinickia* sp. T12–10 in combination or not with *Serendipita indica* (Si) on tomato plants against fungal pathogens. a,b) Biocontrol of *Fusarium oxysporum* (Fol). a Disease index (DI) score. b Fresh weight. Boxes represent standard deviation with median. Whiskers represent the Min to Max of all the values. Analysis of FW and DI was performed using the non-parametric Mann–Whitney U test (*P < 0.05, **P < 0.01, ns = non significant; n = 10); c Biocontrol of *Rhizoctonia solani* (Rs). Points represent mean values, and whiskers standard deviations. Same letters represent nonsignificantly different mean values, according to Tukey’s test (P < 0.05) after ANOVA (n = 3). ns = non significant
growing without kanamycin, few colonies of the strain *Methylobacterium* P1–11 (but not of *Trinickia* T12–10) lost the fluorescence due to plasmid loss (data not shown). The strains *Tardiphaga* sp. P9–11 and *Rhodanobacter* sp. T12–5 were not considered for further experiments as the stability of the transformation was extremely low (Supp. Figure 6c).

Physical interaction between *S. indica* and bacteria and root colonization patterns

To study if bacterial strains affect the morphological structure of *S. indica*, gfp-tagged *S. indica* was grown alone and in combination with mCherry-tagged bacteria. When the fungus was cultured alone, typical *S. indica* structures like tubular and moniliform hyphae and chlamydospores (Kost and Rexer 2013) were observed (Figs. 5a-d). During dual culture of *S. indica* with strains P1–11 or T12–10, fungal structures were not affected (Figs. 5e-w). Hyphae were not disrupted nor deformed, and the thickness of the hyphae remained unaltered. Intriguingly, the fungus seemed to be stimulated, and the mycelium density and the number of chlamydospores appeared to be enhanced in both co-cultures (Figs. 5e and j). Regarding their physical associations, both bacteria were spotted loosely scattered among the hyphae or attached to them (Figs. 5e-g and j-i). Sometimes bacteria aggregated around the hyphae (Figs. 5h and i), which hints at chemotaxis towards the mycelium. The strain P1–11 was often observed attached to the surface of chlamydospores, but never inside them (Figs. 5f-g). However, bacterial cells of strain T12–10 were not only found attached to the surface (Figs. 5j-m) but most strikingly, we repeatedly detected them colonizing the chlamydospores of *S. indica* internally (Figs. 5n-w). Some of the colonized spores were alive as a strong gfp fluorescence was detected (Figs. 5j-k and l-o) and nuclei as well as vesicles were observed (Figs. 5n-o) while others looked as empty spores with bacteria (Figs. 5j-k and p-t). Nonetheless, empty spores were also found in *S. indica* grown alone. Bacteria were not detected inside fungal hyphae but were further observed in living germinated spores (Suppl. Fig. 7). Moreover, in these pairings, no specialized morphological structures were observed upon colonization.

Colonization patterns of tomato plantlets by single and dual inoculations of bacteria and *S. indica* were investigated under sterile conditions and using marked strains. The roots of control plants (untreated) did not differ from plants inoculated with microbes, in terms of structure, cell-wall thickness and fluorescence produced by the secretion of phenolic compounds upon microbe colonization (Compant et al. 2005a, 2005b) (Figs. 6a-c). Strains P1–11 and T12–10 followed similar colonization patterns (Figs. 6d-o). Both bacteria were most abundantly detected in the upper part of the root (Figs. 6j and 7e), and considerably less abundant bacteria were found close to the root tips (Figs. 6f and n). Strain T12–10 was spotted more abundantly than P1–11 in lower

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**Fig. 4** Effect of bacterial strains on root colonization by *S. indica*. Ten days after planting (a–b), both strains increased the root areas colonized by *S. indica*. Eight weeks after planting (c–d), *S. indica* was detected in every root fragment for every treatment (d), however the number of hyphae aggregates was significantly increased when *S. indica* was co-inoculated with *Trinickia* T12–10 (e). The boxplots show all data points, the horizontal line represents the median, surrounded by the upper (25th) and lower (75th) percentiles. The bar plots represent the means of colonization and the whiskers the standard deviations. Same letters represent non-significantly different mean values, according to Tukey’s test (P < 0.05), after ANOVA. n = 20
Fig. 5 Interaction in vitro between *S. indica* (in green) and bacteria (in red). **a** Tubular hyphae and chlamydospores. **b** Representation of (a) in 3D. **c** Moniliform hyphae and ontogenies of chlamydospores. **d** Representation of (c) in 3D. **e** Bacteria loosely scattered among the hyphae and spores. **f** Bacteria attached to the surface of hyphae and spores, without penetrating them, and its 3D representation (g). **h** Bacteria attracted to the hyphae, and its 3D representation (i). **j** High density of chlamydospores and bacteria loosely scattered among the spores. **l** Bacteria attached to the spores, without penetrating them. **k** and **m** Representations of (j) and (l) in 3D. **n** Bacteria colonizing a spore at the germination site. **o** Magnification of (n) in 3D. **p** and **s** Bacteria colonizing fungal spores internally. **q** and **r** Representations of (p) in 3D. **t** Transversal plane of (s) represented from the *y* axis and its 3D representation (u). **v** and **w** Magnifications in 3D of the spore with arrow in (u). a, c, e-f, h, j, l, n, p, s and t: confocal images. b, d, g, i, k, m, o, q-r and u-w: 3D modeling using Imaris software
parts of the roots. Analysis of colonization patterns revealed that these strains settle at the surface of the roots, and aggregate in the intercellular spaces and at lateral root emergence sites (Figs. 6d-g, 6m-n and 7b-c). They were also found internally colonizing epidermis cells (Figs. 6f-g, 5m and 7a-c) and root hairs (Figs. 6h-i, 1 and o). Unlike other bacteria, they were not detected at the root tip level (Compant et al. 2005a, 2005b). Very often bacteria were observed in aggregates forming biofilm-like structures on the surface of the roots (Figs. 6j-k and 7e). In the root-stem transition zone, bacteria used the stomata to gain access to the inner parts of the plants (Figs. 6d-e).

Mycelium of S. indica was detected along the entire length of the roots (Figs. 6p-t), but more abundantly in the upper parts. Hyphae were loosely found among the root hairs (Fig. 6p-s), attached to their surface (Figs. 7f and i) and colonizing them internally (Figs. 6t, 7d and h). Sometimes the fungus colonized in high numbers the lateral root emergence zones. The fungus also colonized the surface of the roots (Figs. 6t and 7o) as well as the interior of exodermis cells, although the hyphae displayed low fluorescence (Figs. 6p and 7j-k). No spores were detected in S. indica control plants.

When S. indica was inoculated with strain P1–11, both microbes followed the same patterns of colonization as when single inoculated (Figs. 7a-i). No differences in extent of colonization were observed. Bacteria and fungus did not exclude each other, and they were found colonizing the same areas. Bacteria were occasionally detected attached to the hyphae (Figs. 7d and g). No fungal spores were observed. Comparably, dual inoculation with S. indica and strain T12–10, resulted in similar colonization patterns as single inoculations (Figs. 7j-o). However, the area of the fungal mycelium colonization seemed to be increased in comparison to the S. indica control (data not shown). The dispersion of bacteria was increased, and they were detected more abundantly in areas close to the root tips as compared to plants single-inoculated with bacteria (Figs. 7n-o). S. indica and strain T12–10 colonized the same root zones, and bacteria were often detected adhered to hyphae (Figs. 7j-k and n-q). In contrast to plants inoculated with S. indica alone, in plants dual-inoculated we repeatedly observed spores attached to hyphae of S. indica on the surface of the roots, and these spores were often surrounded by bacteria (Figs. 7l-m and p-q).

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**Genome analysis**

**Genomic features and taxonomic affiliation**

The genomes of strains Methylobacterium sp. P1–11, Tardiphaga sp. P9–11, Rhodanobacter sp. T12–5 and Trinickia sp. T12–10 were analyzed to determine which gene clusters could be linked to plant and fungal colonization. They have a total size of 6.77, 6.06, 3.96 and 6.22 Mb, and an average G + C content of 68.97%, 61.28%, 65.22% and 62.87%, respectively (Table 1). No repA genes were found in the genomes and therefore we found no evidence of plasmids in any of the strains. The taxonomical analysis only assigned species names to the strains P1–11 and T12–10, closely related to Methylobacterium sp. UNC378MF and Trinickia soli GP25–8 respectively (Suppl. Table 1). The latter strain has recently been moved to the genus Trinickia gen. Nov. (Estrada-de los Santos et al. 2018) and was previously included in the genus Paraburkholderia. Strains P9–11 and T12–5 did not reach the 95% ANI threshold required for species circumscription and were just classified at the genus level as Tardiphaga sp. and Rhodanobacter sp. respectively. The genomic features of the four genomes are summarized in Table 1.

**Genes and proteins predicted to stimulate S. indica growth**

Based on analogies to genes previously reported to be involved in fungal stimulation, we detected here several genes potentially involved in the stimulation of S. indica growth (Fig. 8, see Suppl. Table 2 with detailed numerical values and protein annotation names). Some vitamins (e.g. B12) can solely be synthesized by bacteria (Fang et al. 2017), thus we hypothesize that vitamin production could be involved in fungal growth stimulation. We found numerous genes in all the strains implicated in the synthesis of vitamins: thiamin (B1), riboflavin (B2), pyridoxin (B6), biotin (B7), folate (B9) and cobalamin (B12) (not in T12–5); vitamins K: menaquinone (K1) and phylloquinone (K2) as well as ubiquinone (Q10) (not in P1–11). Moreover, important genes reported to be involved in the provision of nitrogen to the fungus were identified in the four genomes, like those coding for nitrate and nitrite reductase (not in T12–5) (Zuccaro et al. 2011), ammonification and glutamine synthase, as well as the gene coding for nifU in the strains P1–11 and P9–11 (Suppl. Table 3), that is...
involved in nitrogen fixation (Minerdi et al. 2001). Furthermore, various genes predicted to participate in trehalose biosynthesis (Dupponnois and Kisa 2006) and protein and polysaccharide secretion have been detected (Suppl. Table 2).

**Genes and proteins related to plant growth promotion traits**

Several genes related to plant growth promotion (PGP) features were detected in all genomes based on RAST, antiSMASH and protein functional annotation (eggNOG) analysis. The four genomes harbor genes involved in auxin biosynthesis as well as solubilization and transport of phosphates (Table 2, see Suppl. Table 4 with detailed protein and gene annotation names). Genes related to the synthesis of polyamines, phytohormone-like compounds involved in plant growth and stress mitigation (Chen et al. 2018a), were also detected. The strains *Methylobacterium* P1–11 and *Tardiphaga* P9–11 possess genes involved in nitrogen fixation (Suppl. Table 3) and genes participating in the production of nitrilases, which are implicated in the synthesis of plant hormones (Park et al. 2003). Genes involved in siderophore biosynthesis and receptors were detected; potentially contributing to the iron supply to the plant. Strains P1–11 and T12–10 encode genes for aerobactin (Table 2) and vibrioferrin (Suppl. Table 3) biosynthesis (Tanabe et al. 2003), and the antiSMASH analysis further identified biosynthetic gene clusters (BGCs) involved in the synthesis of enterobactin, pyoverdine, taywachelin and malleobactin (Suppl. Table 5). Furthermore, strains P1–11, P9–11 and T12–10 encode the gene acdS for production of 1-amino-cyclopropane-1-carboxylate (ACC) deaminase (Suppl. Table 3) related to plant growth enhancement by lowering plant ethylene levels (Hardoim et al. 2015).

**Resistance to antibiotics and production of antibiotic compounds**

Genes for production of antibiotic compounds as well as for antibiotic resistance may protect the plant against other pathogenic microbes. We identified several genes involved in the production of the antibiotic compounds colicin and bacteriocins (Table 2 and Suppl. Table 4). Strain *Trinickia* T12–10 encodes genes for type IV pili, a bacterial virulence mechanism that appears operational during pathogenesis of fungal hosts (Dörr et al. 1998), genes to produce aminoglycoside antibiotics and a cyclic lipopeptide identified as bananamide (Nguyen et al. 2017) (Suppl. Table 5). Besides, the four genomes encode proteins involved in the synthesis of phenazines (Suppl. Table 3) (Thomashow and Weller 1988), as well as BGCs identified as lipopolysaccharide (LPS) and O-antigen (glycan polymer contained within an LPS) (Suppl. Table 5). Furthermore, CAZy analysis detected enzymes involved in the breakdown of fungal cell-wall components like β-hexosaminidases, exo-β-glucosaminidase and chitooligosaccharide deacetylases (Table 4, see Suppl. Table 6 for detailed information), that have been described to degrade chitin, chitosan and chitooligosaccharides (Mekasha et al. 2016; Katta et al. 2013; Zhao et al. 2010). The RAST analysis further characterized in the genomes of T12–5 and T12–10 several genes under the subsystem “Chitin and N-acetylglucosamine utilization” (data not shown). In a similar way, the four genomes contain genes involved in multidrug resistance efflux pumps, resistance to the bactericide fluoroquinolone and synthesis of β-lactamases, enzymes that confer resistance to...
Fig. 7 Root colonization patterns of dual inoculations of *S. indica* (in green) with *Methylobacterium* P1–11 (in red) and *Trinickia* T12–10 (in red). a Exodermis cell heavily colonized by bacteria through intercellular spaces. b and c 3D representations of (a). d Colocalization of fungus and bacteria during colonization of root hairs (arrow), and bacteria attached to hyphae (asterisk). e Bacteria aggregating in the surface of the upper part of the root, forming biofilm-like structures. f Fungus and bacteria spotted together in lower parts of the roots. g Representation in 3D of (d). h Root hair colonized by fungus (arrow) and bacteria (asterisk). i Representation in 3D of (f). j Hyphae and bacterial aggregates colonizing the surface of the upper part of the root, and mycelium penetrating the exodermis (arrow). k Representation in 3D of (j). l Spores aggregated in the surface of the upper part of the root, surrounded by bacteria (arrow), and a 3D magnification (m). n Hyphae (arrow) and bacterial aggregates (asterisk) colonizing the surface of the root and root hairs in the lower part of the root, and its 3D representation (o). p Colocalization of bacteria, hyphae and spores (arrow) in the surface of roots close to the root tips. q 3D representation of (p). a, d-f, h, j, l, n and p: confocal images. b-c, g, i, k, m, o and q: 3D modeling using Imaris software.
β-lactam antibiotics (Berg et al. 2010) (Table 2 and Suppl. Table 4).

**Genes and proteins related to stress tolerance**

Numerous genes identified in these genomes participate in multiple pathways of protection against environmental stress (Table 2 and Suppl. Tables 4 and 5), and this protection may ultimately result in plant growth promotion (Liu et al. 2017). For instance, genes involved in cell protection against oxidative stress, like those for the synthesis of peroxidases, superoxide dismutases, hydrogen peroxide reductases, catalases, glutathione S-transferases and carotenoids (Suppl. Table 5). They further harbor genes for detoxification, like those coding for cyanide hydratases/nitrilases (Howden and Preston 2009) and rhodanese (Chaudhary and Gupta 2012).

In addition, the four strains might enhance plant resistance to salt stress by producing the osmoregulator glycine betaine (Table 2 and Suppl. Table 4) (Ashraf and Foolad 2007) and the disaccharide trehalose (Penna 2003), as well as genes that operate directly in the protection of membranes and proteins, and functionally in plant immunity, like cold and heat shock proteins, chaperones and glucans (Park and Seo 2015). Moreover, these strains contain numerous copies of genes implicated in plant resistance to heavy metals/metalloids including arsenic, mercury, chromium, copper, cadmium, cobalt and zinc (Table 2 and Suppl. Table 4).

**Table 1** Summary of the genomic features of strains *Methylobacterium* sp. P1–11, *Tardiphaga* sp. P9–11, *Rhodanobacter* sp. T12–5 and *Trinickia* sp. T12–10

| Feature            | P1–11          | P9–11          | T12–5          | T12–10         |
|--------------------|----------------|----------------|----------------|----------------|
| Length (bp)        | 6,774,627      | 6,056,105      | 3,960,066      | 6,220,544      |
| Coverage           | 185.5 ± 75.8   | 480.6 ± 83.0   | 763.9 ± 117.9  | 358.9 ± 115.5  |
| Completeness (%)   | 100.00         | 99.6           | 99.95          | 99.31          |
| G + C content (%)  | 68.97          | 61.28          | 65.22          | 62.87          |
| Contamination (%)  | 0.94           | 0.72           | 1.20           | 0.57           |
| Contigs            | 68             | 18             | 16             | 88             |
| Total genes        | 6596           | 5776           | 3516           | 5741           |
| Predicted CDS      | 6509           | 5699           | 3445           | 5657           |
| rRNA number        | 3              | 3              | 3              | 2              |
| tRNA number        | 63             | 49             | 50             | 55             |
| misRNA number      | 19             | 24             | 17             | 26             |
| tmRNA number       | 2              | 1              | 1              | 1              |
Interestingly, strains P1–11, P9–11 and T12–10 encode the gene oxc implicated in the catabolism of oxalate (Table 2), which has been reported to be required for successful plant colonization by the endophyte *Paraburkholderia phytofirmans* PsJN (Kost et al. 2014). Furthermore, these strains harbor numerous genes for ROS-detoxification (Table 2), required for overcoming the plant defense response upon colonization (Kandel et al. 2017). All these genomic features presumably permit these bacteria to colonize roots and adapt to the endophytic lifestyle.

Fig. 8 Circular visualization of genomes using the Blast Ring Image Generator (BRIG). Track 1: (innermost) genome map; Track 2: GC%; Track 3: GC Skew; Track 4: contig boundaries as alternating red and black color; Track 5: genome positions coding for genes potentially involved in the stimulation of fungal growth. Vitamins of the B complex (in red) are indicated by B (e.g. B12 stands for Vitamin B12); Genes involved in nitrogen supply (in black) include nitrate reductase (NR1), nitrite reductase (NR2), and nitrate/nitrite/ammonium transporters (NT). Trehalose biosynthesis (in green) is indicated by TR. Numbers (2,3,4..) before the abbreviation of genes indicated if more than one gene are present at a certain position.
Table 2 Protein encoding genes predicted to be involved in plant growth promotion and resistance of strains Methylobacterium sp. P1–11, Tardiphaga sp. P9–11, Rhodanobacter sp. T12–5 and Trinickia sp. T12–10 determined by RAST. (n° R: number of subsystem roles/ proteins; n° G: number of protein encoding genes (peg))

| Category                      | Subsystem                                      | P1–11 | P9–11 | T12–5 | T12–10 |
|-------------------------------|------------------------------------------------|-------|-------|-------|--------|
|                               |                                                | n°R   | n°G   | n°R   | n°G    |
| Phosphate solubilization uptake and transport | Phosphatase                                    | 3     | 4     | 6     | 7      |
|                               | Low-affinity inorganic phosphate transport system | 2     | 3     | 6     | 7      |
|                               | High-affinity phosph. Trans. system            | 8     | 10    | 9     | 8      |
| Siderophore                   | Aerobactin biosynthesis                        | 4     | 4     |       |        |
|                               | Siderophore receptors and transport            | 4     | 5     | 4     | 5      |
| Polyamine                     | Putrescine/ spermidine synthesis               | 4     | 5     | 4     | 5      |
|                               | Putrescine/ spermidine transport               | 8     | 12    | 6     | 10     |
| Plant hormone                 | Auxin biosynthesis                             | 4     | 4     | 5     | 4      |
| Resistance to antibiotics     | Polymyxin resistance                           | 6     | 6     |       |        |
|                               | Colicin tolerance                              | 1     | 1     | 1     | 1      |
|                               | Fluoroquinolone resistance                     | 4     | 5     | 4     | 4      |
|                               | Beta-lactamase                                 | 3     | 9     | 3     | 8      |
|                               | Vancomycin resistance                          | 3     | 9     | 3     | 8      |
|                               | Multidrug resistance efflux pumps              | 9     | 19    | 7     | 20     |
|                               | Tripartite multidrug resistance system         | 3     | 8     | 3     | 17     |
|                               | Oxalate catabolism                             | 1     | 1     | 1     | 1      |
|                               | 4-hydroxybenzoate degradation                 | 1     |       |       |        |
| Resistance to heavy metals    | Cobalt-zinc-cadmium resistance                 | 9     | 29    | 7     | 19     |
|                               | Copper homeostasis/ tolerance                  | 9     | 13    | 8     | 13     |
|                               | Arsenic resistance                             | 3     | 5     | 4     | 5      |
|                               | Mercury resistance and detoxification          | 1     | 1     | 1     | 1      |
|                               | Chromium compounds resistance                  | 1     | 3     | 2     | 3      |
| Antibiosis compounds          | Production of colicin and bacteriocin          | 6     | 7     | 6     | 7      |
|                               | Flavonoid (apigenin)                           | 1     |       |       |        |
|                               | Type IV pilus                                  |       |       |       |        |
| Resistance to oxidative stress| Peroxidase                                     | 3     | 5     | 3     | 6      |
|                               | Catalase                                       | 1     | 1     | 1     | 2      |
|                               | Superoxide dismutase                           | 2     | 2     | 2     | 1      |
|                               | Glutathione S-transferase                      | 6     | 21    | 5     | 26     |
|                               | Hydroperoxide reductase                        | 1     | 1     | 1     | 2      |
| Heat/cold shock               | Heat shock protein/ chaperone                  | 9     | 11    | 9     | 11     |
|                               | Cold shock protein                             | 3     | 8     | 3     | 8      |
| Cyanide detoxif.              | Rhodanese                                      | 1     | 1     | 1     | 1      |
| Salt tolerance                | Choline/ betaine uptake & synthesis            | 5     | 8     | 8     | 16     |
|                               | Synthesis of osmoregulated periplasmic glucans | 4     | 8     | 3     | 3      |
|                               | Trehalose biosynthesis                         | 9     | 12    | 9     | 11     |
Discussion

The interactions between fungi and bacteria have often been overlooked, but they seem to be more widespread than expected and their dynamics may be crucial in natural ecosystems (Deveau et al. 2018). Here, these endophytic bacteria are suspected to positively interact with the functioning of the fungal symbiosis, therefore they might be considered as mycorrhizal helper bacteria (Frey-Klett et al. 2007). These strains greatly promoted fungal growth in vitro, enhanced plant growth, and their combination with *S. indica* could effectively confer resistance against plant pathogens. We previously found other strains within the genus *Mycolicibacterium* exhibiting similar traits although the functioning underlying the tripartite interactions seem to be partly distinct.

Effective microbial inoculum must colonize plant roots and overcome the plant immune system (Compant et al. 2005a; Liu et al. 2017). In the genomes of the tested bacterial strains the presence of multiple genes typically considered to be involved in root colonization and endophytism was congruent with microscopy observations in which root exodermis cells and hairs were internally colonized by strains *Methylobacterium* P1–11 and *Trinickia* T12–10. These strains were detected in different root zones, and particularly strain T12–10 expanded more throughout the root, probably due to the higher number of genes involved in flagellar motility and secretion systems. In addition to the bacterial genomic features, it has been reported that plants might also facilitate the entry of bacteria into periplasmic spaces in root cells, so that plants might acquire nutrients from bacterial protoplasts in a process known as rhizophagy cycle (White et al. 2018). In the process of root colonization *S. indica* and bacteria did not exclude each other but they were commonly co-localized, and rather than competing they seem to cooperate, since the addition of bacteria resulted in increased hyphae expansion, fungal sporulation and root zones colonized by the fungus. Interestingly, we often found bacteria attached to fungal hyphae, hence bacteria might employ hyphae as vectors to colonize new niches along the root (Kohlmeier et al. 2005; Miquel Guennoc et al. 2018).

Most striking was to find cells of *Trinickia* T12–10 colonizing spores of *S. indica*, thus the endosymbiotic associations among Sebacinales and bacteria could be much broader in nature than we previously thought. Congruently with this finding, bacteria from the order Burkholderiales are among the most frequently identified intracellular bacteria in fungal hyphae (Partida-Martinez and Hertweck 2005; Frey-Klett et al. 2011) and several *Burkholderia* sensu lato strains are endosymbionts of beneficial endophytic fungi (Compant et al. 2008). Moebius et al. (2014) revealed that a type 2 secretion system (T2SS) of *Burkholderia rhizoxinica* is required for the endosymbiosis with *Rhizopus microsporus*. In that study, mutants deficient in the genes *gspC* and *gspD* (general secretion pathways) were unable to internally colonize the fungus and induce sporulation. Among these bacteria, strain T12–10 contains the highest number of genes involved in T2SS,

| Table 3 | Protein encoding genes predicted to be involved in colonization and establishment of endophytes in the host of strains *Methylobacterium* sp. P1–11, *Tardiphaga* sp. P9–11, *Rhodanobacter* sp. T12–5 and *Trinickia* sp. T12–10 determined by RAST |
|---------|--------------------------------------------------|
| Function/Strain (n° genes) | P1–11 | P9–11 | T12–5 | T12–10 |
| Motility & Chemotaxis | Bacterial chemotaxis | 49 | 37 | 20 | 54 |
| | Flagellar assemblage | 49 | 81 | 51 | 67 |
| | Flagellar motility | 27 | 36 | 25 | 41 |
| Cell Wall & Capsule | Exopolysaccharide | 11 | | | |
| | Lipopolysaccharide | | 24 | 23 | |
| Membrane Transport | Type IV pilus | | 27 | 21 | |
| | Secretion system, Type I | | | 6 | |
| | Secretion system, Type II | 15 | 15 | 11 | 57 |
| | Secretion system, Type V | | | 3 | |
| | Secretion system, Type VI | 18 | | 49 | |
| | Secretion system, Type VII | | 2 | | |
| | Secretion system, Type VIII | | | 6 | |

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| CAZy family | Substrate | Annotation | EC number | Copy number |
|-------------|-----------|------------|-----------|-------------|
| CE1 | Polysaccharides | esterase | 3.1.1.- | 2 1 1 1 |
| CE4 | Polysaccharides | polysaccharide deacetylase | 3.5.1.- | – – 1 1 |
| Chitoooligosaccharide | chitooligosaccharide deacetylase | 3.5.1.- |
| CE9 | Polysaccharides/ Chitoooligosaccharides | N-acetylglucosamine 6-phosphate deacetylase | 3.5.1.25 | – – 1 1 |
| CE11 | Lipopolysaccharide | UDP-3-O-acetyl N-acetylglucosamine deacetylase | 3.5.1.- | 2 1 1 1 |
| GH1 | Cellulose | β-glucosidase | 3.2.1.21 | – 1 1 1 |
| Pectin (rhamnogalacturonan I) | β-galactosidase | 3.2.1.23 |
| GH2 | Hemicellulose (galactomannan) | β-mannosidase | 3.2.1.25 | 1 1 3 3 |
| Pectin (rhamnogalacturonan I) | β-glucuronidase | 3.2.1.31 |
| Chitosan | exo-β-glucosaminidase | 3.2.1.165 |
| GH3 | Chitoooligosaccharides | β-N-acetylhexosaminidase | 3.2.1.52 | 1 1 3 3 |
| Cellulose | glucan 1,4-β-glucosidase | 3.2.1.74 |
| GH4 | Cellulose | exo-β-1,4-glucanase | 3.2.1.4 | 1 – – – |
| GH8 | Cellulose | cellulase | 3.2.1.4 | 1 – – – |
| GH13 | Polysaccharides | α-amylase | 3.2.1.11 | 8 9 6 6 |
| pullulanase | 3.2.1.41 |
| trehalose synthase | 5.4.99.16 |
| GH15 | Polysaccharides | glucomylase | 3.2.1.3 | 2 1 1 1 |
| GH17 | Polysaccharides | glucan 1,3-β-glucosidase | 3.2.1.39 | – 2 2 2 |
| exo-beta-1,3-glucanase | 3.2.1.- |
| GH19 | Peptidoglycans/Chitin | lysozyme | 3.2.1.17 | – – – 1 |
| GH20 | Polysaccharides/Chitin | β-hexosaminidase | 3.2.1.52 | – – 2 – |
| GH23 | Peptidoglycans | peptidoglycan lyase | 4.2.2.n1 | 6 4 2 3 |
| GH25 | Peptidoglycans/Chitin | lysozyme | 3.2.1.17 | 1 – – – |
| GH30 | Polysaccharides | O-Glycosyl hydrolase | 3.2.1.- | – – 1 – |
| GH31 | Polysaccharides | α-glucosidase | 3.2.1.20 | – – 1 – |
| GH37 | Trehalose | α,α-trehalase | 3.2.1.28 | – – 2 2 |
| GH39 | Hemicellulose (xylan) | β-xylosidase | 3.2.1.37 | – – 2 – |
| GH42 | Pectin (rhamnogalacturonan I) | β-galactosidase | 3.2.1.23 | – – – 2 |
| GH50 | Agar | β-agarase | 3.2.1.81 | – – 1 – |
| GH65 | Polysaccharides | trehalose phosphorylase | 2.4.1.64 | 1 1 3 – |
| β-phosphoglucomutase | 5.4.2.6 |
| GH73 | Peptidoglycan/Chitin | endo-β-N-acetylhexosaminidase | 3.2.1.96 | – – 1 2 |
| lysozyme | 3.2.1.17 |
| GH77 | Polysaccharides | 4-α-glucanotransferase | 2.4.1.25 | – 1 – 1 |
| GH92 | Oligosaccharides | α-mannosidase | 3.2.1.24 | – – 5 – |
| GH94 | Polysaccharides | cellubiose phosphorylase | 2.4.1.20 | – – 2 1 |
| cyclic β-1,2-glucan synthase | 2.4.1.321 |
| GH102 | Peptidoglycans | peptidoglycan lyase | 3.2.1.- | 1 1 – 1 |
| GH103 | Peptidoglycans | peptidoglycan lyase | 3.2.1.- | 4 6 1 1 |
including gspC and gspD, which presumably enables this strain to adhere and colonize fungal structures. Strain P1–11, however, lacks these genes and harbors fewer T2SS genes. This could explain why T12–10 could internally colonize the spores and induce sporulation, while P1–11 did not enter spores. Strain T12–10 encodes also genes involved in fungal cell wall degradation (e.g. β-hexosaminidase), although it lacks the gene (chi) for chitinase production, which was shown to be crucial for entering fungal hyphae in the B. rhizoxinica-R. microsporus symbiosis. Consequently, it is not yet understood how Trinickia T12–10 gains access to the spores of S. indica. Likewise, the mechanism by which RfF4 invades S. indica is yet unresolved. This bacterium lacks the chi gene and even T2SS genes, albeit concomitant with T12–10, genes for T4SS, involved in adhesion, are present (Glaeser et al. 2016). It has been hypothesized that germinating spores of S. indica can be potential entry sites for bacteria (Guo et al. 2017), and for penetrating growing hyphal tips where walls are thin, other enzymes different from chitinases (e.g. glucanases) might suffice. We detected T12–10 in apparently empty spores, but also inside those with an intact cytoplasm as well germinated spores. Nevertheless, it claims further investigation if the bacterium T12–10 could be vertically transmitted to the next generation of spores.

The rhizosphere and the roots of most plants are host of complex interactions between different microorganisms, but the precise nature and mechanisms of these interactions are yet to be elucidated. To obtain further understanding on potential mechanisms, we analyzed the genomes of these bacterial helpers. We detected a great number of genes related to vitamin production seemingly involved in fungal growth stimulation. For instance, cobalamin (vitamin B12) produced by bacteria was reported to be implicated in the promotion of fungal growth and fitness (Ghignone et al. 2012). Jiang et al. (2018) found that S. indica is auxothophic for thiamine (vitamin B1), and the bacterium Bacillus subtilis complemented S. indica’s auxotrophy for thiamine and promoted its growth. S. indica holds biotrophic-associated genomic adaptations, hence some genes and protein families have undergone contraction, like those for nitrate and nitrite reductase, amino acid and ABC transporters, and is thus predicted to suffer from some metabolic deficiencies. Congruently, the fungus barely grows on nitrate as sole N source (Zuccaro et al. 2011). The genomes of the bacterial strains tested in this study harbor several genes coding for nitrate and nitrite reductase, amino acid and peptide ABC transporters, ammonium transporters and glutamine synthase as well as nitrogen fixation, which might enhance fungal nitrogen metabolism, palliating the predicted deficiencies. Furthermore, Salvioli et al. (2016) revealed that an endobacterium of Gigaspora margarita enhanced fungal sporulation and increased the bioenergetic capacity of the mycorrhizal fungus by inducing oxidative phosphorylation and ATP production. Investigating S. indica, Bhuyan et al. (2015) detected that genes coding for enolase-I and urease D, which are involved in the activation of several hexose sugar transporters and in the glycolytic pathway for generating ATP, were up-regulated in the presence of the fungal-stimulating bacterium Azotobacter chroococcum WR5. Similarly, the bacteria tested in this study, and particularly strain T12–10 that can internally colonize fungal structures, might up-regulate sugar uptake, glycolysis and oxidative phosphorylation, prompting ATP production and increased fungal fitness. There is still no clear evidence if only one, or several genomic features detected in these bacteria are responsible for fungal growth stimulation. Nevertheless, these isolates did not stimulate the growth of the fungal pathogens, as opposed to S. indica. Hence there seems to exist specificity in the stimulation of fungal growth, as it has been earlier reported for other mycorrhizal helper bacteria (Frey-Klett et al. 2007). To better elucidate the molecular basis behind the fungal stimulation, further studies should focus on the functional role of these bacterial factors by knock-out mutants, and transcriptomics of fungus and bacteria to detect genes up and down-regulated during co-cultivation.

The tested bacteria harbor also many genes involved in plant growth promotion, which confirms the potential of these strains as plant biostimulants. Correspondingly, strains Tardiphaga P9–11, Rhodanobacter T12–5 and Trinickia T12–10 increased plant fresh weight and leaf area, although strain Methylobacterium P1–11 did not affect plant growth. In agreement with the widely reported beneficial effect of S. indica on plant growth (Franken 2012), S. indica significantly increased tomato growth in this experiment. Dual inoculations, however, did not further improve the growth-promoting effect exerted by microbes inoculated singly. Interestingly, inoculation of S. indica + T12–5 significantly displayed lower performance than single inoculations. Certainly, there are several reports of combinations of fungi and
bacteria providing no better or, in some cases, worse plant promotion than the isolates used singly (Whipps 2001). In our previous study, combinations of \textit{S. indica} with strain P9–22 or P9–64 of \textit{Mycolicibacterium} negatively affected plant growth in comparison to single inoculations (del Barrio-Duque et al. 2019). The same negative effect was described for co-inoculation of pseudomonad R62 and \textit{S. indica} (Sarma et al. 2011). This incompatibility of fungal-bacterial inoculants has been ascribed to competition for nutrients and niches in the rhizosphere (Whipps 2001), although in this study \textit{S. indica} and bacteria seemed to cooperate during colonization. The lack of synergism might also be caused by alterations in the plant hormone levels. Rather low levels of exogenous auxins stimulate plant growth whereas high concentration may cause a negative impact (Sarwar and Frankenberger 1994). Since these bacteria and \textit{S. indica} can produce auxins (Sirrenberg et al. 2007), the dual inoculation might lead to an imbalance of auxins levels and consequently less growth promotion. In analogy, a further explanation may be over-colonization of roots by the fungus. \textit{S. indica} begins colonization with a biotrophic growth phase invading living cells, followed by a cell death-dependent phase, in which root cells are actively killed by the fungus (Qiang et al. 2012b). If the number of dead cells transcends a threshold, \textit{S. indica} could manifest a negative influence on plant growth (Fakhro et al. 2010). Our isolates greatly stimulated fungal growth in vitro, and by microscopy observations we detected enhanced hyphae expansion in plants dual-inoculated with \textit{S. indica} and strain T12–10. Bacteria might stimulate fungal growth and root colonization until the threshold is surpassed, shifting the fungus thus from mutualistic to saprotrophic and impairing plant development.

On the basis of the accelerated depletion of the currently available fungicides, and the huge amounts of crops affected worldwide by plant pathogens, the upcoming generation of bioinoculants shall combine plant growth enhancement with biological control of plant pathogens (Collinge et al. 2019). Notwithstanding that these bacterial strains harbor genes involved in biocontrol of plant diseases, bacteria inoculated individually were generally not able to control fungal pathogens. This fact complies with the in vitro tests in which confrontations between bacteria and the pathogens did not display a direct suppression of fungal growth. Only the strain \textit{Trinickia} T12–10 significantly reduced the progression and severity of Fusarium wilt. In agreement, numerous strains of \textit{Burkholderia} sensu lato have been described to induce plant resistance against fungal pathogens (Compant et al. 2008).

It has been reported that \textit{S. indica} colonization results in the activation of defense related genes and ISR, which consequently improves plant tolerance against biotic stresses (reviewed in Franken 2012; Gill et al. 2016). Consistently, in this study \textit{S. indica} also reduced disease progression caused by fungal pathogens. However, the highest degree of protection was observed when the plants were dual-inoculated with combinations of \textit{S. indica} and bacteria. This synergistic effect was also observed in our previous research, in which Fusarium wilt’s severity was significantly reduced by dual-inoculations of \textit{S. indica} with \textit{Mycolicibacterium} strains P1–18 and P9–22 in comparison to single inoculations, and biocontrol of \textit{Rhizoctonia} resulted most effective by combining \textit{S. indica} with \textit{Mycolicibacterium} P9–64 (del Barrio-Duque et al. 2019). Interestingly, the endofungal bacterium of \textit{S. indica} (\textit{Rhizobium} radiobacter F4) was found to induce resistance responses in barley and confer resistance against different pathogens (Sharma et al. 2008; Glaeser et al. 2016). Consequently, it was speculated that the beneficial effect exerted by this fungus might actually be attributable to the bacterium (Qiang et al. 2012a; Glaeser et al. 2016). Following this line, it is likely that other bacterial strains might also contribute to further increase the beneficial effect reported for this fungus. There might be a synergism in rising the plant ISR (defense) when \textit{S. indica} and certain bacteria are inoculated. Genome analysis of the bacterial strains tested in this study revealed several genes involved in the activation of the plant ISR, like those involved in the synthesis of PCWDEs, phenazines, LPS and O-antigens, type IV pili, antibiotics, EPS and flagella (Pieterse et al. 2014; Van Loon et al. 1998; Lerouge and Vanderleyden 2002) that might complement the \textit{S. indica}-triggered ISR. In analogy, there might be a synergism in increasing antioxidant activity in plant cells. \textit{S. indica} induces systemic resistance by increasing the concentration of antioxidants, glutathione and ascorbate in the plants to mitigate the oxidative stress caused by pathogens (Waller et al. 2005; Gill et al. 2016). Since our bacterial strains encode numerous genes involved in the resistance to oxidative stress, like glutathione S-transferase, peroxidase, and genes participating in the synthesis of carotenoids (Stahl and Sies 2003), we can hypothesize that with the addition of bacterial helpers to the fungal inoculum, the plant has additional “tools” to
cope with the oxidative stress caused by pathogens. Besides, on the basis of ethylene as a stress hormone, potential production of ACC-deaminase by these bacteria might have an additional asset to protect plants (Wang et al. 2000). Furthermore, the genomes of the tested bacteria encode also genes involved in direct biocontrol of pathogens, like genes involved in siderophores synthesis and receptors that deprive the pathogens from iron, enzymes involved in the degradation of fungal cell-wall structures, phenazines or cyclic lipopeptides (Whipps 2001; Compant et al. 2005a). The outcome of the multipartite interaction among these beneficial microbes, the pathogens and the plant might rely on several of these mechanisms. Perhaps the combined effect of competition for nutrients and space, S. indica-triggered ISR and the bacterial features directly and indirectly involved in biocontrol might justify the increased resistance of plants inoculated with S. indica + bacteria.

All in all, what really happens inside the roots and the rhizosphere seems to be a complex network of multitrophic cooperation and several factors drive the outcome of the plant-microbe interaction. When searching for plant growth enhancement, the application of beneficial microbes should consider the nutrient conditions of the soil. In this study, dual and single inoculations of fungus and bacteria in rich soil did not promote shoot growth while on low-nutrient soil it did. This nutrient-dependent effect upon S. indica inoculation on tomato was previously reported (Fakhro et al. 2010). This research group also demonstrated that the concentration of S. indica inoculum was crucial, and negative effects were observed when high cell doses were applied. Similarly, inoculation of S. indica with a high concentration of its endobacterium resulted in harmful, if not lethal, effects to the fungus (Sharma et al. 2008). It is evident that the optimal concentration of S. indica, and the bacteria to accompany the fungal inoculum, should be further investigated and additional research is needed to further elucidate the interaction of the growth-promoting fungus and its bacterial partners. Soil condition, plant determinants (physiology, species and cultivar), environmental factors, dosage of the inoculum and microbial compatibility must be addressed for the development of new biostimulants and biocontrol agents.

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Author contribution statement Alejandro del Barrio-Duque carried out the experiments and wrote the manuscript. Abdul Samad provided expertise in data analysis. Livio Antonielli was responsible for genome assemblage and helped with statistics. Ole Nybroe provided guidance for bacterial transformation. Alejandro del Barrio and Stéphane Compant designed the experiments. All the authors gave also intellectual input and critically revised the manuscript.

Compliance with ethical standards

Conflict of interest SC is section editor of Plant and Soil. However, this does not interfere with the independent review process.

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Author contribution statement Alejandro del Barrio-Duque carried out the experiments and wrote the manuscript. Abdul Samad provided expertise in data analysis. Livio Antonielli was responsible for genome assemblage and helped with statistics. Ole Nybroe provided guidance for bacterial transformation. Alejandro del Barrio and Stéphane Compant designed the experiments. All the authors gave also intellectual input and critically revised the manuscript.

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