Tapping into the maize root microbiome to identify bacteria that promote growth under chilling conditions

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Abstract

Background: When maize (Zea mays L.) is grown in the Northern hemisphere, its development is heavily arrested by chilling temperatures, especially at the juvenile phase. As some endophytes are beneficial for plants under stress conditions, we analyzed the impact of chilling temperatures on the root microbiome and examined whether microbiome-based analysis might help to identify bacterial strains that could promote growth under these temperatures.

Results: We investigated how the maize root microbiome composition changed by means of 16S rRNA gene amplicon sequencing when maize was grown at chilling temperatures in comparison to ambient temperatures by repeatedly cultivating maize in field soil. We identified 12 abundant and enriched bacterial families that colonize maize roots, consisting of bacteria recruited from the soil, whereas seed-derived endophytes were lowly represented. Chilling temperatures modified the root microbiome composition only slightly, but significantly. An enrichment of several chilling-responsive families was detected, of which the Comamonadaceae and the Pseudomonadaceae were the most abundant in the root endosphere of maize grown under chilling conditions, whereas only three were strongly depleted, among which the Streptomycetaceae. Additionally, a collection of bacterial strains isolated from maize roots was established and a selection was screened for growth-promoting effects on juvenile maize grown under chilling temperatures. Two promising strains that promoted maize growth under chilling conditions were identified that belonged to the root endophytic bacterial families, from which the relative abundance remained unchanged by variations in the growth temperature.

Conclusions: Our analyses indicate that chilling temperatures affect the bacterial community composition within the maize root endosphere. We further identified two bacterial strains that boost maize growth under chilling conditions. Their identity revealed that analyzing the chilling-responsive families did not help for their identification. As both strains belong to root endosphere enriched families, visualizing and comparing the bacterial diversity in these communities might still help to identify new PGPR strains. Additionally, a strain does not necessarily need to belong to a high abundant family in the root endosphere to provoke a growth-promoting effect in chilling conditions.

Keywords: Chilling temperatures, Root endosphere, Microbiome, Maize, Plant growth-promoting rhizobacteria, PGPR

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Background
Maize (Zea mays L.) is one of the most important staple crops worldwide with an annual production of approximately 1.4 billion tonnes in 2017 (www.fao.org). Due to its subtropical origin, cultivation of maize in the Northern hemisphere is rather challenging because of chilling spring temperatures [1]. These chilling temperatures, on average 17 °C during the day and 12 °C during the night, retard plant development, eventually resulting in yield losses [2–4]. As additional arable land is scarce, the proposed increase in maize needs to result from higher yields per area.

To boost plant growth, plant growth-promoting rhizobacteria (PGPR) have been proposed as an ecological additive under both normal and stress conditions [5–8]. As well in the plant roots, i.e. the root endosphere, and as in the closely surrounding soil, the rhizosphere, microbial communities are established that can affect plant fitness [6]. Deep-sequencing technologies have allowed us to obtain insight into the diversity of these microbial communities [9–13], demonstrating that the endo- and rhizospheres are occupied by a vast selection of microbes that are recruited from the soil. The plant compartment plays an important role in this process: bacterial community compositions gradually change from the soil toward the inside of the root [10–12]. The root endosphere-colonizing microbes are expected to engage in robust interactions with the plant roots, but they could also have been acquired by vertical transmission of seed microbes besides recruitment from the soil [14–16].

Varying environmental cues, including pathogen attack [17, 18] and abiotic stresses, such as drought and salt [19, 20], modify the microbial composition of the root endosphere and rhizosphere. As a result, plants have been proposed to select a microbiome that provides the best plant fitness [17–19].

Even though chilling temperature is one of the major abiotic stresses many plants have to cope with during their life cycle, especially when grown in temperate regions, its effect on the microbiome has not been investigated yet, either in maize or other plants. More effort has been put in the identification of particular PGPR strains that can alleviate chilling stress in plants. Several psychrotolerant bacterial strains belonging to different families, such as Burkholderiaceae, Bacillaceae, and Pseudomonadaceae, have been identified that can help plants such as Arabidopsis thaliana, rice (Oryza sativa), wheat (Triticum sp.), tomato (Solanum lycopersicum), and grapevine (Vitis vinifera), to cope with chilling stress [21–26]. However, to our knowledge, bacteria that can promote maize growth under chilling stress have not been identified yet.

As chilling stress restricts maize growth, research on its impact on the microbiome might guide the selection of possible growth-promoting bacteria. To test this hypothesis, in independent repeats, we identified the main endosphere microbiome of maize roots and analyzed the effects of chilling temperatures on its composition. In parallel, a bacterial collection of maize endophytes was established and screened for growth promotion of juvenile maize under chilling stress.

Materials and methods
Microbiome experimental design
Five experiments (experiments I to V) were set up with different objectives (Table 1). For all experiments, surface-sterilized hybrid uncoated maize seeds (LG30.217, Limagrain, Saint Beauzire, France) were used. Briefly, seeds were washed 5 min with sterile water, 2 min with 70% (v/v) ethanol, 20 min with a bleach solution (29 ml sterile water, 15 ml NaClO, 12–13% (v/v) stock solution, and 1 ml Tween 20), five times for 15 min with sterile water, then air-dried, and stored at 4 °C until further use.

In experiment I, seeds were sown according to common agricultural practices in an experimental field with sandy loam soil (United State Department of Agriculture classification) (50° 58’ 41’’ N, 3° 46’ 47.28’’ E; Merelbeke, Belgium). Approximately after 3 weeks of growth, until development of the sixth leaf, bulk soil samples and root endosphere samples of five plants were collected as described below.

In experiment II, top soil was collected from the same field as that for experiment I and sieved before use. Soil

| Experiment | Growth conditions | Compartment studied | Main objective |
|------------|-------------------|---------------------|---------------|
| I          | Maize in the field | Bulk soil vs. root endosphere | Define main root microbiome (field setup) |
| II         | Maize in field-soil filled pots | Bulk soil vs. root endosphere | Define main root microbiome (growth chamber setup) |
| III        | Maize in field-soil filled pots and in vitro | Root endosphere of plants grown in soil- vs. in vitro- | Define origin of root microbiome = seed or soil |
| IV         | Maize in field-soil filled pots under control and chilling conditions | Root endosphere of maize grown under control vs. chilling conditions | Define chilling-responsive families |
| V          | Maize in field-soil filled pots under control and chilling conditions | Root endosphere of maize grown under control vs. chilling conditions | Define chilling-responsive families |
characteristics were determined at the beginning of the experiment (Additional file 1: Table S1). Twelve pots were filled with freshly collected soil, six pots were kept uncultivated and used for bulk soil sampling, whereas in six other pots, sterilized maize seeds were sown and the root endosphere was sampled after development of leaf 6 (4 weeks of growth). All pots were maintained in the growth room under controlled conditions (16 h/8 h light/dark regime, 21 °C) and kept well-watered throughout the time of the experiment.

In experiment III, as in experiment II, top soil from the experimental field was collected and 12 pots were filled with the collected soil. Again, six were kept bare for bulk soil samples and in the six remaining pots, maize was sown in soil to identify the root endosphere of soil-grown plants after 2 weeks of growth. Additionally, six sterilized seeds were sown in transparent plastic boxes filled with Hoagland’s solution (0.945 g/l Ca(NO$_3$)$_2$.4H$_2$O, 0.506 g/l KNO$_3$, 0.136 g/l KH$_2$PO$_4$, 0.493 g/l MgSO$_4$.7H$_2$O, 2.5 ml/l Fe-EDTA stock (5.56 g FeSO$_4$.7H$_2$O, 7.46 g/l Na-EDTA)) supplemented with plant agar. Due to space limitation in the boxes, samples were collected after 2 weeks of growth under constant conditions (16 h/8 h light/dark regime, 21 °C).

In experiments IV and V, ten maize plants were grown as in experiment II for 5 weeks under chilling (16 h/8 h light/dark regime at 17 °C/12 °C) or control temperature conditions (16 h/8 h light/dark regime, 21 °C) and kept equally well-watered throughout the time of the experiment. The sampling of the field soil for experiment IV was repeated in experiment V, but after a 1-year gap.

To ensure that bulk soil and root endosphere samples were treated similarly, both were washed before DNA extraction. For the bulk soil, samples were washed in phosphate-buffered saline (PBS) solution for 20 min and centrifuged at 3220×g for 20 min to ensure that all bacteria and soil particles were dissolved in the pellet; however, mostly soil-particle-sorbed microbes will be captured. The supernantant was removed and the remaining soil pellet was frozen in liquid nitrogen and stored at −80 °C. For the root endosphere sample collection, adhering soil was removed by shaking the roots vigorously. Roots were washed twice in PBS by shaking in 500-ml sterile flasks with 50 ml PBS for 20 min, sonicated (10 min of 30-s cycles at 4000 Hz) to remove remaining sticking microorganisms, flash-frozen in liquid nitrogen, and stored at −80 °C. Roots were ground in liquid nitrogen before DNA extraction. DNA was isolated from all collected samples with the DNeasy PowerSoil DNA kit (QIAGEN, Hilden, Germany), whereas the V4 region (515F-806R) of the 16S rRNA gene was amplified as proposed by The Earth Microbiome Project (www.earthmicrobiome.org). Reverse bar-coded primers were used to amplify the V4 region in triplicate with the iProof High-Fidelity PCR Mix (Bio-Rad, Hercules, CA, USA) with 30 cycles of amplification at 55 °C for 30 s. Pooled PCR products of the triplicate samples were purified with Agencourt AMPure XP magnetic beads (Beckman Coulter, Pasadena, CA, USA) and DNA concentration was measured with a Qubit fluorometer (Thermo Fisher Scientific, Waltham, MA, USA). Based on previous experiments, a high amount of 16S rRNA gene reads from chloroplastic and mitochondrial origin was expected in the root endosphere, thus the bulk soil and root endosphere samples were unequally pooled in a 1:5 ratio. Sequencing was done on an Illumina MiSeq platform (v3, 2x300 bp) according to the Illumina protocols (VIB, Nucleomics Core, Leuven, Belgium). The raw sequence reads were deposited and are available for download at the NCBI sequence reads archive (SRA) with project number PRJNA523518 (experiments I, II, and III) and PRJNA524079 (experiments IV and V).

The sequencing reads were demultiplexed and primers were removed by the sequencing provider. The sequence read quality was checked by FastQC [27]. Read quality and length trimming were done with DADA2, using the default quality score (truncQ = 2); forward reads were truncated at 240 bp and reverse reads at 200 bp (truncLen = c(240,200)) and a maximum of three expected errors both in the forward and in the reverse reads was allowed (maxEE = c(3,3)). Determination of amplicon sequence variants (ASVs) and taxonomy assignments were done by means of the standard DADA2 pipeline v1.6.0 [28]. After ASV determination, each sample had approximately 100,000 quality checked read (Additional file 1: Table S3). The resulting count table was filtered to remove low abundant ASVs (less than two counts/ASV in at least five samples), reducing the read count per sample to an average of 40,000 reads (Additional file 1: Table S3). Taxonomy was assigned according to the SILVA Database version 128 [29]. Based on the taxonomy assignment, reads belonging to chloroplasts (Class Chlorella) and mitochondria (Order Rickettiales) were removed, reducing the number of reads by an average of 5% and 80% in the bulk soil and root endosphere, respectively. The resulting ASV count tables were used for further statistical analysis.

**Bacterial collection and screening assay**

To obtain a collection of endophytic bacterial strains, the maize plants were grown in soil collected from the experimental field for 4–6 weeks under chilling temperature conditions (16 h/8 h light/dark at 17 °C/12 °C). To ensure isolation of endophytes, maize roots were washed and sterilized as follows: three times for 5 min with sterile water, 30 s with 70% (v/v) ethanol, 3 min with bleach solution (12% [v/v] sodium hypochlorite; Chem-Lab, Zedelgem, Belgium), and five times with sterile water for 5 min. Roots were crushed on ice with
stereile mortar and pestle in PBS, pushed through a 70-
μm cell strainer, and diluted 1/100 and 1/1000 with PBS.

Diluted suspensions were plated on three different
commonly used bacterial media TSB, King’s B, and R2A (Addi-
tional file 1: Table S2) and incubated for maximum 2 weeks.
Growing colonies were selected, streaked until pure cultures,
and identified based on 16S rRNA gene using universal 16S
rRNA primers: 27F (AGAGTTTGATCMTGCTCAG) and
1492R (GGTTACCTTGTTACGACTT). DNA was ex-
tracted with alkalitic lysis [30]. In total, 1 μl of supernatant
DNA was used for PCR amplification with 0.2 μl polymerase
(IProofTM High Fidelity DNA Polymerase; Bio-Rad), 1 μl of
10 μM forward primer, and 1 μl of 10 μM reverse primer,
0.4 μl 10 mM dNTP mix, and 4 μl buffer 5× IProof buffer
(Bio-Rad). PCR conditions were denaturation at 98 °C for 3
min, 30 cycles of 98 °C for 10 s, 55 °C for 20 s, and 72 °C for
45 s, and a final elongation at 72 °C for 5 min. The resulting
PCR products were purified with the GeneJET PCR Purifica-
tion Kit (Thermo Fisher Scientific) and sequenced by Sanger sequencing (Eurofins Genomics, Ebersberg,
Germany). Forward and reverse sequences were merged by
means of the CLC Main Workbench 7 (CLC Bio-QIAGEN)
and taxonomy was assigned to the resulting sequences with
the SILVA database version 128.

Of the bacterial collection, isolates representing either the
main root endosphere microbiome families identified in ex-
periments I and II, the chilling-responsive endosphere fam-
ilies identified in experiments IV and V, or other endophytic
families were selected. In total, 28 of the 282 bacterial isolates
in the collection were tested for maize growth promotion.
Maize seeds were surface-sterilized as described above and
pre-germinated for 48 h in the dark at 24 °C on 1% (w/w)
agar plates. The effect of each strain was tested in two or
three repeats. For each repeat, 15 seedlings were inoculated
by shaking in a bacterial solution for 3 h. The different bac-
terial isolates were grown in liquid medium and diluted to
OD600 0.02 with PBS buffer. For each experiment, a mock-
inoculated treatment was included, for which 15 seedlings
were inoculated with PBS buffer for 3 h before sowing. Bac-
terial- and mock-inoculated seedlings were sown in sand/
perlite (50/50 volume %) and cultivated under controlled
growth conditions in the growth chamber (16 h/8 h light/
dark regime at 17 °C/12 °C). The 15 plants were cultivated in
separate pots (square pots: 7 × 7 × 8 cm); per treatment, all
pots were put together on one tray. The entire tray was
watered (500 ml) every 2 days, while nutrients (150
ml Hoagland’s solution for 15 plants) were added once
a week. After 30 days of growth, plants were
harvested and the fresh total (root and shoot) weight
was analyzed. Additionally, the effect of the two iden-
tified PGPR strains, RHG5 and RHG12, was analyzed
under control temperature conditions. The experimental
setup was the same as for the screening under
chilling conditions, with the exception that the

growth conditions differed, namely 16 h/8 h light/
dark regime and 21 °C, and the plants were harvested
after 25 days of growth.

Statistical analysis
All statistical analyses were done in R (version 3.4.0). For the
microbiome experiments, the ASV count tables, generated
by DADA2 [28], were filtered to remove low abundant ASVs.
ASVs with a count number of 2 in at least five samples were
retained. The multivariate analysis was done using the R
package vegan (version 2.0-10) [31]. The dissimilarity matrix,
based on the Bray-Curtis dissimilarity index, was calculated
from the ASV tables as generated by DADA2 [28]. By means
of the betadisper function, the homogeneity of the variances
was checked on this dissimilarity matrix. Further, the signifi-
cance of sample type (e.g., bulk soil, root endosphere of soil-
grown and in vitro-grown plants) or chilling treatment and
experiment were analyzed with PERMANOVA, in which the
Bray-Curtis dissimilarity index matrix was used as input. The
temperature experiments IV and V were considered as bio-
logical repeats. To test the effect of the experiment, the com-
bined ASV table was analyzed with PERMANOVA, showing
‘biological repeat’ as a significant factor (P < 0.001) that
shapes the bacterial communities. Hence, the statistical ana-
lyses were done separately for each experiment. Additionally,
PERMANOVA analysis demonstrated a significant inter-
action effect in both experiments between sample type (bulk
soil and root endosphere) and temperature (control and
chilling) (P < 0.001).

The differential abundance was assessed with the
likelihood-ratio tests. Analyses were done at the levels of
ASV, family, and phylum, using the ASV counts and tax-
onomy for the latter two. The data were normalized based
on the library size of the count table, resulting in analysis of
the relative abundances. All these analyses were done with
the edgeR package, version 3.18.1 [32].

In the screening assay, the statistical differences in growth
promotion due to bacterial inoculation were analyzed by
general linear models, after the assumptions of linearity, nor-
mality, and homogeneity of the data had been checked. The
experiment was repeated two or three times and an inter-
action effect between treatment and repeat was identified (P
< 0.05). Therefore, the growth-promoting effects of bacterial
strains were examined for each repeat separately. The fresh
total (root and shoot) weight total was compared between
bacteria- and mock-inoculated plants with a two-sided Stu-
dent’s t test (analysis done in R).

Results
Identification of the main root microbiome families of
maize grown in field soil
To identify the main bacterial community of the maize
root endosphere (hereafter designated root endosphere),
grown in field soil, two experiments were carried out. In
experiments I and II (Table 1), maize was grown in the field and in field soil-filled pots, respectively. Sequences assigned to chloroplasts and mitochondria were removed from quality-filtered reads, resulting in ASV count tables containing 1013 unique ASVs in experiment I and 820 in experiment II (Additional file 1: Table S3). Sample type, i.e., bulk soil or root endosphere, was the major variance driver (87.15% and 86.25% for experiments I and II, respectively) for the differences in the bacterial community composition as found by the PERMANOVA analysis ($P < 0.001$) and illustrated in the Bray-Curtis dissimilarity-based principal coordinate analysis (PCoA) plot (Fig. 1a).

In both maize grown in the field (experiment I) and in pots (experiment II), five highly abundant phyla were significantly enriched ($P < 0.001$) in the root endosphere compared to the bulk soil: *Proteobacteria*, *Bacteroidetes*, *Chloroflexi*, *Firmicutes*, and *Actinobacteria* (Additional file 1: Table S4). Two dominating phyla in the pot-grown root endosphere were *Spirochaetae* and *Cyano- bacteria*, phyla that were either not detected or at a very low abundance in the root endosphere of field-grown plants (Additional file 1: Table S4).

To characterize the main root endosphere microbiome, we compared the results of both field- and pot-grown plants. The main microbiome was defined as the bacterial families that were noteworthy abundant (relative abundance > 0.5%) and significantly enriched ($P < 0.05$) in the root endosphere of both experiments (Fig. 1b, d). We detected 12 families that met these requirements, namely *Anaerolineaceae*, *Blissiellaceae*, *Caulobacteraceae*, *Comamonadaceae*, *Cytophagaceae*, *Erysilipelotrichaceae*, *Flavobacteriaceae*, *Hyphomicrobiaceae*, *Oxalobacteraceae*, *Rhizobiaceae*, *Streptomyceae*, and an unclassified *Xanthomonodales* family (Fig. 1c; d; Additional file 1: Table S4). Although the same enrichments were found in both experiments, the abundance of the bacterial families differed for the two root endosphere communities. The identified main microbiome families accounted for 76.94% and 45.97% of the bacterial community of the root endosphere of field-grown (experiment I) and pot-grown (experiment II) plants (Fig. 1c), respectively. The most striking discrepancy occurred in the abundance of *Actinobacteria*. In the root endosphere of field-grown maize (relative abundance ± SE; 27.18 ± 2.91%), the relative abundance of this phylum was 5-fold higher than in the pot-grown maize root endosphere (5.67 ± 0.64%) (Additional file 1: Fig. S1). In both experiments, almost all actinobacterial ASVs were assigned to one family, the *Streptomycetaceae* (Additional file 1: Fig. S1). The enrichment was strong under both conditions (logFC 9.4 and 7.3 for experiments I and II, respectively) and the initial abundance in the bulk soil was low (0.035 ± 0.02% and 0.05 ± 0.01% for experiments I and II, respectively). Thus, it is tempting to speculate that the growing conditions, i.e., field or pot, affect the *Streptomycetaceae* abundance.

Although the abundance of the *Proteobacteria* did almost not vary between the pot-grown (46.09 ± 3.52%) and field-grown (52.42 ± 2.14%) root endosphere, the abundance of the proteobacterial family *Oxalobacteraceae* differed, accounting for almost 30% (27.77 ± 3.16%) of the root endosphere of field-grown maize and less than 4% (3.62 ± 0.35%) of the root endosphere of pot-grown maize. In comparison, the abundance of this family in the field bulk soil (2.6 ± 0.76%) was higher than in the pot bulk soil (0.14 ± 0.01%), whereas the fold changes between the experiments were comparable (logFC 4.26 vs. 4.86) (Additional file 1: Table S4).

To conclude, the growth conditions in field soil and field soil-filled pots affected the root endosphere microbiome. Despite differences in abundances, the recurring enrichment pattern remained similar and resulted in the identification of a main root microbiome of maize grown in Belgian field soil of 12 different families.

**Contribution of the seed-inherited root microbiome to the root microbiome**

The root endosphere is colonized by soil-inhabiting bacteria, i.e., bacteria that were recruited through horizontal transmission from the soil, and by seed-inhabiting bacteria that colonize the inner tissues of the seed and might be potentially considered vertically transmitted [14, 15, 33, 34]. In view of identifying bacterial strains for application via seed coating, e.g., the inoculation of bacteria at the outside of the seed, it is important to make out whether bacteria can colonize the root endosphere from the surrounding soil or via the seeds. To this end, we set up a third experiment (experiment III; Table 1). Here, we compared bulk soil samples, root endosphere samples of 2-week-old maize seedlings grown in vitro under gnotobiotic soil-free conditions, and root endosphere samples of seedlings grown in field soil-filled pots.

PERMANOVA analysis showed that sample type explained most of the variation in the bacterial community structure as illustrated in the PCoA plot ($P < 0.01$; based on Bray-Curtis distances) (Fig. 2a). Besides distinct clustering of the bulk soil samples as in experiments I and II, the root endosphere of the plants grown in soil and in vitro also clustered separately. This clustering can be partially explained by the different nature of the growth substrate of the plants, i.e., field soil vs. Hoagland’s solution provided with agar. Besides the substrate, the clustering can also be affected by the strong difference in read counts after filtering out the reads assigned to chloroplasts and mitochondria of the root endosphere of soil-grown and in vitro-grown maize; in total, 10% and 1%, respectively, of the reads were retained (Additional file 1: Table S3). As a result, the bacterial
diversity in the root endosphere of the in vitro-grown plants was very low (Additional file 1: Fig. S2).

In total, 21 bacterial ASVs were detected in the root endosphere of in vitro-grown plants, belonging to three different phyla: Bacteroidetes (four ASVs; relative abundance ± SE, 19 ± 1.23%), Firmicutes (one ASV; 3 ± 0.34%), and Proteobacteria (16 ASVs; 76 ± 1.24%), represented by 12 different families (Fig. 2b; Additional file 1: Table S5). Nine of these ASVs were among the 426 ASVs detected in the bulk soil bacterial community retained after filtering out chloroplast, mitochondrial, and low abundant reads. The family Moraxellaceae (Proteobacteria) contained seven and three ASVs in the root endosphere of in vitro or in soil-grown plants, respectively. However, the family was not detected in the bulk soil indicating that these ASVs most probably colonized the root endosphere through the seeds (Fig. 2b; Additional file 1: Table S5). The second most abundant family in the root endosphere of in vitro-grown plants was the Flavobacteriaceae (two ASVs; 16.2 ± 0.63%), with one ASV representing 13% that also occurred in the root endosphere of soil-grown plants. Several Flavobacteriaceae were detected in the bulk soil as well, but were not the same ASVs as those found in the root endosphere of in vitro-grown plants. Two of the ASVs detected in the root endosphere of in vitro-grown plants were present both in the bulk soil and root endosphere of soil-grown plants, one belonging to the Pseudomonadaceae family and one belonging to the Enterobacteriaceae family (Fig. 2b). Whether these bacteria came through the seeds or via the soil into the root endosphere cannot be distinguished. Based on these data, we can assume that the root endosphere was acquired mainly from the surrounding soil microbial community, although transmission through the seeds occurred as well.
The effect of chilling temperature on the maize root endosphere microbiome

To analyze the effect of chilling temperature on the root endosphere, we set up experiments IV and V (Table 1; Fig. 3a). In both experiments, maize was grown in field soil-filled pots under either ambient or chilling temperature conditions. After filtering and removal of plant-related reads, 1137 and 962 unique ASVs were detected for experiments IV and V, respectively (Additional file 1: Table S3). Besides the expected shift in community due to sample type (see experiments I and II), we observed a significant interaction effect between temperature and sample type, indicating that the bacteriological changes upon chilling varied between the bulk soil and the root endosphere ($P<0.001$). Both in the bulk soil and in the root endosphere, the temperature had a significant effect on the bacterial communities (Additional file 1: Table S6). As seen in the PCoA plots based on Bray-Curtis distances (Fig. 3b), the shift in the microbial community of the chilling-treated bulk soil was smaller in both experiments than that in the root endosphere bacterial communities (Fig. 3b; Additional file 1: Fig. S3 and Table S6). In the root endosphere, temperature was responsible for more than 40% of the detected variance (41.43% and 48.77% in experiments IV and V, respectively), whereas in the bulk soil samples, the variation differed less (38.01% and 34.48% in experiments IV and V, respectively) (Additional file 1: Table S6). In the bulk soil of experiment IV, chilling stress had the highest influence on the Chitinophagaceae, increasing with a logFC of 1.6, from $1.6 \pm 0.19$ to $4.6 \pm 0.52$% under chilling conditions ($P<0.001$). In the bulk soil of experiment V, chilling temperatures resulted in a major increase in the abundance of Blastocatellaceae, from an average of $14.08 \pm 0.63$% to $28.20 \pm 0.91$% (Additional file 1: Table S7). Thus, no recurring pattern in community shifts at the family or phylum level was demonstrated in the bulk soil.

In contrast, the root endosphere differed significantly between the control and chilling - grown maize plants. At phylum level, a significant decrease in Actinobacteria in the root endosphere upon chilling was detected in both experiments IV and V ($P<0.001$) (Additional file 1: Table S7). Additionally, we identified 15 families (relative abundance > 0.5%) that consistently responded in the root endosphere under chilling conditions, of which 12 were enriched and three were depleted in both experiments. We compared the

![Fig. 2](image-url)

**Fig. 2** Contribution of the seed-inherited root microbiome to the maize root microbiome. **A** PCoA plot based on Bray-Curtis dissimilarity indices of the microbial communities in bulk soil and root endosphere of soil-grown and of in vitro-grown maize in experiment III. **B** Presence and abundance of the 21 ASVs belonging to 11 different families, detected in the root endosphere of in vitro-grown maize plants and compared with the abundance of the root endosphere of soil-grown maize and bulk soil samples.
so-called chilling-responsive families with the identified main microbiome and detected an overlap of four bacterial families (Fig. 3c). The 12 enriched chilling-responsive families (relative abundance > 0.5%) were Enterobacteriaceae, Pseudomonadaceae, Sphingobacteriaceae, Sphingomonadaceae, Methylophilaceae, Phyllobacteriaceae, Opitutaceae, Cellibrionaceae, Flavobacteriaceae, Comamonadaceae, Caulobacteraceae, and Xanthomonadaceae (Fig. 3d; Additional file 1: Table S7). Furthermore, three other chilling-responsive families were recurrently depleted in the chilling versus the control root endosphere, i.e., Chitinophagaceae, Burkholderiaceae, and Streptomycetaceae (Fig. 3d Additional file 1: Table S7). The latter is part of the Actinobacteria phylum that also significantly decreased in the root endosphere upon chilling and consisted primarily of Streptomycetaceae (Fig. 3d Additional file 1: Table S7). The latter is part of the Actinobacteria phylum that also significantly decreased in the root endosphere upon chilling and consisted primarily of Streptomycetaceae (Fig. 3d Additional file 1: Table S7). The latter is part of the Actinobacteria phylum that also significantly decreased in the root endosphere upon chilling and consisted primarily of Streptomycetaceae (Fig. 3d Additional file 1: Table S7).

In conclusion, two independent experiments in which the same field soil was used showed that chilling temperatures affected the microbial communities in the root endosphere in a recurring pattern, both causing enrichment and depletion of certain chilling-responsive families and, to a lesser extent, in the bulk soil environment.

Screening for growth-promoting bacterial strains of maize under chilling temperatures

We hypothesized that under chilling temperatures, the root endosphere might recruit bacterial taxa that could trigger adaptation to this stress and promote plant growth. To this end, a bacterial collection of 282 maize root endophytic bacterial strains was established from plants grown in chilling temperatures (Additional file 2: Table S8A, B, C). The collection represented four well-known root endosphere phyla, i.e., Proteobacteria (n = 178), Actinobacteria (n = 29), Bacteroidetes (n = 14), and Firmicutes (n = 61) as well as 22 different endophytic families (Additional file 2: Table S8A). The bacterial collection represents more than half (n = 7) of the families of the identified main microbiome, i.e., Flavobacteraceae, Comamonadaceae, Caulobacteraceae, Streptomycetaceae, Rhizobiaceae, Oxalobacteraceae, and Cytophagaceae (Fig. 4). Four families that are both identified in the main microbiome and chilling-responsive are present in the collection: Flavobacteraceae, Comamonadaceae, Caulobacteraceae, and Streptomycetaceae (Fig. 3c and 4). Additionally, six other chilling-responsive families could be isolated, i.e., Enterobacteraceae, Pseudomonadaceae, Burkholderiaceae, Xanthomonadaceae, Sphingomonadaceae, and the Sphingobacteriaceae (Fig. 4).

A subset of the bacterial collection was screened for their growth-promoting effects on the total fresh weight of...
juvenile maize grown under chilling conditions (Fig. 5; Additional file 1: Table S9 and Fig. S4). The subset used for screening contained 14 strains belonging to main microbiome families, 11 strains belonging to the chilling-responsive families, and three strains of families enriched in the root endosphere but low abundant. Growth promotion of the bacterial strains was evaluated based on the fresh total weight (root and shoot), of bacteria-inoculated versus mock-inoculated maize. Two strains showed consistent growth-promoting effects over the different repeats: RHG5 (Bradyrhizobiaceae – Bosea sp.) and RHG12 (Oxalobacteraceae – Pseudoduganella sp.) with an average fresh plant weight increase of 18% and 33%, respectively (Fig. 5; Additional file 1: Table S9 and Fig. S4). The Oxalobacteraceae were identified as a main microbiome family, whereas the Bradyrhizobiaceae were enriched in the root endosphere, albeit at a low relative abundance, but neither were identified as chilling-responsive families.

Additionally, to investigate whether the two identified PGPR strains specifically promoted maize growth under chilling temperatures or whether they were robust growth strains, we tested their effect under control temperature conditions. Neither RHG5 nor RHG12 consistently affected maize growth under these control temperatures (Additional file 1: Fig. S5).

Furthermore, to detect whether the PGPR selection would be improved at the sequence level, we compared the identified PGPR strains of experiments IV and V at sequence level. RHG12 had a 100% similarity with ASV ID_10, that was highly abundant in the chilling and control root endosphere of both experiments IV and V and was strongly enriched compared to the bulk soil (Additional file 1: Table S10). In contrast, for RHG5, we could not detect an ASV with a 100% similarity in the filtered count table.

Thus, selection of bacterial strains belonging to chilling-responsive families is not enough to detect PGPR; nevertheless, families enriched in the root endosphere might be a good criterium to pick strains for large screening assays.

Discussion
Chilling temperatures are one of the major abiotic stress factors affecting maize growth in the Northern hemisphere, with yield decrease as a consequence [3]. The use of growth-promoting bacteria for maize could boost maize growth under chilling conditions, but are, to our knowledge, not identified yet, despite other PGPR strains for maize had been detected previously [35–37]. We explored the maize root endosphere microbiome and the effect of chilling temperatures on the root endosphere bacterial communities and assessed whether these results could facilitate the selection of PGPR strains to promote maize growth under chilling conditions.

By means of two different experimental setups, we identified 12 root endosphere families as the main microbiome of maize grown in field soil. We consider these families as robust colonizers over different experiments for the same soil sampled at various time points, confirming previous results in other crop systems [11–13]. Previously, a core maize microbiome has been determined at ASV level [38]. In total, eight of the 12 families we identified in our setup, had been described as core microbiome members [38]. Hence, these families, Caulobacteraceae, Comamonadaceae, Cytophagaceae, Hymenobacteraceae, Oxalobacteraceae, Rhizobacteraceae, Streptomycesaceae, and Xanthomonadales can be considered as true core microbiome families of maize. The differences however confirm that the root endosphere community is also highly influenced by factors, such as soil, genotype, experimental set-up, and climatic conditions [12, 13, 39, 40].

Bacterial inoculants currently on the market are applied via seed coatings indicating that the bacterial inoculant should be able to colonize the maize roots from the surroundings [41, 42]. Hence, we hypothesized that seed-derived root endophytes would have a lower efficiency in root colonization than the bacteria that are recruited from the soil and thus are of less interest for application. Correspondingly, we did not screen these bacterial families for growth promotion. We revealed that the soil-grown root endosphere is only colonized to a minor extent by seed endophytes [43, 44]. Firstly, we validated that the family Moraxellaceae is recruited from the seed tissues because of its absence in the bulk soil samples and its abundance in the root endosphere of
both soil- and in vitro-grown maize. Due to their absence in the bulk soil, we can speculate that members of the \textit{Moraxellaceae} family are true endophytes that can be transmitted from the seeds to the root endosphere. This family has been previously discovered in root environments and several genera of the \textit{Moraxellaceae} are known to promote plant growth \cite{45-47}. Secondly, the family \textit{Flavobacteriaceae} was detected in the root endosphere of both soil- and in vitro-grown maize and additionally, also in the bulk soil samples. Accordingly, we hypothesize that this family can colonize the root endosphere both through seed transmission and via the surrounding soil. Together, most of the detected root endophytes are recruited from the soil and can thus be used in seed coating strategies. Whether or not seed-transmitted bacteria, such as those belonging to the \textit{Moraxellaceae}, perform worse in seed coating treatments than soil-recruited strains has to be tested in the future. Additionally, it would also be interesting to analyze whether recruitment of bacterial families from the seed into the root is influenced by environmental factors, such as chilling temperatures.

Abiotic stresses affect bacterial communities in root and soil \cite{13, 18, 39}. A shift in the bacterial community was
detected upon chilling conditions that, as under drought conditions [39], is more pronounced in the root endosphere than in the bulk soil. Hence, it is tempting to speculate that plants mediate the selection of the bacterial communities in the root endosphere under stress conditions. As changes in root exudates are known to affect the microbiome [18, 43, 48] and abiotic stress conditions to alter the root exudate composition [49, 50], the chilling effect on the root exudates might have an impact on the root endosphere microbiome. Therefore, it would be worthwhile to study whether the detected shifts are caused by the influence of temperature on the bacterial life cycle or on the entire bacterial population, by plant physiological alterations, or by a combination of these factors.

Based on two independent experiments under chilling stress, 12 chilling-responsive families were detected that were repeatedly enriched in the root endosphere upon chilling conditions. Additionally, three other chilling-responsive were recurrently depleted upon chilling temperatures of which the actinobacterial family *Streptomycetaceae* was the most striking one. This family has previously been shown to be enriched in endo- and rhizospheres of several grass species upon drought, indicating that not every abiotic stress provokes the same types of changes in the composition [19, 20]. Further experiments are needed to determine whether the decrease in *Streptomycetaceae* in our study is caused by the actinobacterial life cycle and whether their susceptibility to chilling conditions is indirectly affected by chilling on other bacterial families or by the alternate plant-mediated attraction of microbes during chilling periods.

We hypothesized that enriched chilling-responsive families might promote growth of maize under chilling conditions. To investigate whether the microbiome results could be used as a tool to facilitate PGPR selection, we screened a selection of maize endophytes from an established bacterial collection for their growth-promoting capacities under chilling conditions. RHG12 (*Oxalobacteraceae–Pseudoduganella* sp.) and RHG5 (*Bradyrhizobiaceae–Bosea* sp.) were identified as growth-promoting strains under chilling temperatures, whereas they did not under normal temperature conditions. Both belong to families that are, although known to be endophytic, not studied in-depth for these abilities. We demonstrated that *Oxalobacteraceae* are part of the main endosphere microbiome but not chilling responsive. As *Oxalobacteraceae* were not detected in the endosphere of in vitro-grown plants (experiment III), we hypothesize that this family colonized the root system from the soil environment. Further research on the maize–*Oxalobacteraceae* interaction and colonization will help to unravel questions regarding the underlying growth-promoting pathways and whether the plant has an active role in the recruitment of this bacterial family.

The *Bradyrhizobiaceae* family, to which RHG5 belongs, was enriched, however low abundant in the root endosphere. Similarly as for the *Oxalobacteraceae*, this family was not demonstrated to be chilling-responsive and not detected in the root endosphere of in vitro-grown plants. The family contains known rhizobia species that uniquely interact with legumes, such as *Bradyrhizobium japonicum* that nodulates soybean (*Glycine max*) [51]. Based on the genome, the family has been linked with nitrogen fixation [52], but the link with the detected growth promotion should however be further determined.

Thus, both identified PGPR strains belong to endosphere-enriched families of which neither were shown to be chilling-responsive, implying that strains do not necessarily have to be enriched in the endosphere upon chilling conditions to enhance plant growth under these conditions. At the sequence level, we only identified a matching ASV for RHG12, ID_10, which was highly abundant and strongly enriched in the root endosphere, whereas it did not shift under chilling conditions. Hence, enrichment and high abundance in the endosphere are not always required to promote plant growth. Robust root colonization on the contrary, typical for endophytes, might be required to maintain the growth-promoting effects on the plant, because it can indeed be assumed as a competitive advantage in the complex root-surrounding bacterial community [51, 53, 54]. Nevertheless, the effect of the identified strains has only been tested in the semi-sterile sand-perlite system and growth-promoting strains could have been missed because of the different environmental conditions. Ideally, validation should be done within field soil environments and preferentially in the field. Because such experiments are not feasible for large-scale screening assays, the effects of only the two identified PGPR should definitely be validated in the field in the future. Additionally, to better understand the link between root endosphere enrichment and PGPR, a larger screening with higher throughput should be pursued. More in-depth studies on the colonization of the different growth-promoting strains and their functional traits both in reductionist and complex approaches will further help to address this question.

**Conclusions**

Plants create a stable root endosphere microbiome containing mostly bacteria that are attracted from the surrounding soil environment into the root endosphere, whereas only a few are derived from the seed. When grown under chilling conditions, the shift detected in the bacterial communities of the roots was more pronounced than that upon chilling in the bulk soil samples. Our screening attempts demonstrated that strains do not need to belong to families that respond to chilling temperatures to provoke growth promotion under these conditions. However, because both strains belong to root endosphere-enriched families and are expected to be
good root colonizers, coating seeds with these strains might definitely help the growth of juvenile maize in chilling temperatures. Hence, the two PGPR strains identified here will be studied further in detail to explore their potential for agricultural applications.

**Supplementary information**

**Supplementary information** accompanies this paper at https://doi.org/10.1186/s40168-020-00833-w.

**Additional file 1: Figure S1.** Difference in the relative abundances between experiments I and II of the phylum Actinobacteria and the Streptomyces coeruleus family. **Figure S2.** Shannon diversity indices (effective number of species) accounting for the within-sample diversity of the different sample types (all experiments) and between temperature conditions (experiments IV and V). **Figure S3.** PCAo plots of the endosphere and bulk soil samples of experiment IV and V. **Figure S4.** The effect of all the screened bacterial strains on the total fresh weight of maize grown under chilling conditions. **Figure S5.** The effect of RHG5 and RHG12 on maize grown under control temperature conditions. **Table S1.** Soil characteristics of the field soil used in the different experiments (determined by ILVO). **Table S2.** Composition of bacterial community media used for isolation approaches. **Table S3.** Read counts of the different samples (before and after filtering) in each microbiome experiment. **Table S4.** Bacterial communities of bulk soil and endosphere in field and pot experiments I and II. **Table S5.** Bacterial seed endophytes (experiment III). **Table S6.** Results of the PERMANOVA analysis for experiments IV and V. **Table S7.** Relative abundance of bacterial communities of the temperature experiments IV and V. **Table S9.** Effect of the screened bacterial isolates on the root, shoot and total fresh weight. **Table S10.** Relative abundance of ASV ID_10 in experiment IV and V.

**Additional file 2: Table S8. A.** Summary of the phyla and families represented by the bacterial isolates in the bacterial collection. **Table S8 B.** Bacterial isolates of the collection with the assigned taxonomy. **Table S8 C.** Bacterial isolates of the collection with the 16S rRNA DNA sequence.

**Acknowledgments**

The authors thank Veronique Storme for help with statistical analyses, Kelly Hamonts for input on the manuscript, and Martine De Cock for help in preparing it.

**Authors’ contributions**

S.B., T.V., S.V., and S.G designed the experiments. S.B and T.V. collected microbiome samples. S.B., A.H., R.T, and J.R. performed raw data analysis. S.B., T.V., S.V., and S.G interpreted the data and wrote the manuscript. All authors read and approved the final manuscript.

**Funding**

This research was supported by grants from the Research Foundation–Flanders (project no.1259418N to C.D.T) and the VIB Tech Transfer projects. S.B. is a postdoctoral fellow of the Research Foundation–Flanders – Strategic Basic Research (project no.151553).

**Availability of data and materials**

The raw sequence reads were deposited and are available for download at the NCBI sequence reads archive (SRA) with project number PRJNA523518 (experiments I, II, and III) and PRJNA524079 (experiments IV and V).

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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**Received: 6 December 2019 Accepted: 19 March 2020**

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