Understanding the Impact of Cultivar, Seed Origin, and Substrate on Bacterial Diversity of the Sugar Beet Rhizosphere and Suppression of Soil-Borne Pathogens

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The rhizosphere microbiome is crucial for plant health, especially for preventing roots from being infected by soil-borne pathogens. Microbiota-mediated pathogen response in the soil-root interface may hold the key for microbiome-based control strategies of phytopathogens. We studied the pathosystem sugar beet—late sugar beet root rot caused by Rhizoctonia solani in an integrative design of combining in vitro and in vivo (greenhouse and field) trials. We used five different cultivars originating from two propagation sites (France, Italy) with different degrees of susceptibility towards R. solani (two susceptible, one moderately tolerant and two cultivars with partial resistance).

Analyzing bacterial communities in seeds and roots grown under different conditions by 16S rRNA amplicon sequencing, we found site-, cultivar- and microhabitat-specific amplicon sequences variants (ASV) as well as a seed core microbiome shared between all sugar beet cultivars (121 ASVs representing 80%–91% relative abundance). In general, cultivar-specific differences in the bacterial communities were more pronounced in seeds than in roots. Seeds of Rhizoctonia-tolerant cultivars contain a higher relative abundance of the genera Paenibacillus, Kosakonia, and Enterobacter, while Gaiellales, Rhizobiales, and Kosakonia were enhanced in responsive rhizospheres. These results indicate a correlation between bacterial seed endophytes and Rhizoctonia-tolerant cultivars. Root communities are mainly substrate-derived but also comprise taxa exclusively derived from seeds. Interestingly, the signature of Pseudomonas poae Re*1-1-14, a well-studied sugar-beet specific biocontrol agent, was frequently found and in higher relative abundances in Rhizoctonia-tolerant than in susceptible cultivars. For microbiome management, we introduced microbial inoculants (consortia) and microbiome transplants (vermicompost) in greenhouse and field trials; both can modulate the rhizosphere and mediate tolerance towards late sugar beet root rot. Both, seeds and soil, provide specific beneficial bacteria for rhizosphere assembly and microbiota-
mediated pathogen tolerance. This can be translated into microbiome management strategies for plant and ecosystem health.

Keywords: sugar beet cultivars, seed microbiome, root microbiome, vermicompost, biocontrol, breeding, *Pseudomonas poae* RE*1-1-14

INTRODUCTION

Developing concepts for microbiome-based crop management strategies is challenging due to the multi-fold interactions in these complex systems. It further requires a deep scientific understanding of microbial community dynamics. Microbiome network structure and microbial diversity in the rhizosphere of plants are linked with tolerance towards pathogen invasion (van Elsas et al., 2012; Berg et al., 2017). The rhizosphere is the critical soil-plant interface for resource exchange and interaction between the plant and microbiota (Weller et al., 2002; Philippot et al., 2013). The crucial involvement of rhizosphere-associated microbiota for growth promotion and stress tolerance in crops is known for more than a century (Hiltner, 1904), but was impressively accelerated by the progresses in omics-technologies (Mendes et al., 2012; Raaijmakers and Mazzola, 2016). Due to the importance of microbial rhizosphere assembly for plant health, it was intensively studied in the last decades (Berendsen et al., 2012; Bakker et al., 2013). Diverse factors shaping the microbial rhizosphere community have been identified, with plant genotype and soil traits as the most important determining factors (Berg and Smalla, 2009). However, both plant genotype as well as soil quality, were strongly changed in the period of increasingly intensified agriculture. This resulted in changes in plant-associated microbial communities and reduced indigenous antagonistic potential towards plant pathogens, especially towards soil-borne pathogens (Cardinale et al., 2015; Perez-Jaramillo et al., 2017; Banerjee et al., 2019). Crops often lack clear genetic resistance to soil-borne pathogens. Cook and colleagues postulated already in 1995 that plants could compensate this by recruiting antagonists of pathogens from the soil microbiome. This was now evidenced by omics-technologies (Mendes et al., 2012; Carrión et al., 2019). Although, there are first studies indicating that breeding for resistance towards soil-borne pathogens unintentionally shaped the structure and function of the rhizosphere microbiome, e.g. in the pathosystem common bean (*Phaseolus vulgaris*)—*Fusarium oxysporum* (Mendes et al., 2018a; Mendes et al., 2018b), a generalized relationship between the rhizosphere microbiome structure and plant immunity/tolerance could not be established yet.

Seeds were identified as carrier and ideal target for rhizosphere's microbiome management (Berg and Raaijmakers, 2018). During the last years, the vertical transmission via seed or propagule endophytes has been described for many plants (e.g. Johnston-Monje et al., 2016; Klaedtke et al., 2016; Bergna et al., 2018). However, the role of seed endophytes and their interplay with soil microorganisms in rhizosphere assembly and microbiota-mediated pathogen response is not yet understood (Berg and Raaijmakers, 2018). Furthermore, many basic insights e.g. about the impact of propagation site on seed endophyte assembly are still missing (Bergna et al., 2018). Seed endophytes play an important role in the respective plant-pathogen interactions (Shade et al., 2017). To what extent microbial seed communities are conserved or otherwise shaped by the propagation sites' soil communities may be important for pathogen susceptibility of the next plant generation. Furthermore, only seed endophytes that survive the dynamic process of germination will be represented in the endophytic community of the seedling and its rhizosphere (Shade et al., 2017).

Sugar beet is an interesting model crop for microbiome studies (Zachow et al., 2008; Mendes et al., 2012; Kusstascher et al., 2019a; Kusstascher et al., 2019b) known for their genome and breeding history (Würschum et al., 2013; Dohm et al., 2014). Sugar beet (*Beta vulgaris* L.) is an important root crop and the main source of sucrose in moderate climates. It is grown on approximately $4.8 \times 10^6$ hectares, resulting in $2.7 \times 10^8$ tonnes yield in 2018, with Russia and France being the main producers worldwide (FAOSTAT, 2019). One of the major pathogens in sugar beet is the fungus *Rhizoctonia solani* J.G.KÜHN [teleomorph: *Thanatephorus cucumeris* (A.B.Frank) Donk], which causes a variety of different plant diseases and has a broad host range (Ogoshi, 1996). Especially the late root rot, a disease caused by *R. solani* of the anastomosis group AG2-2IIIB leads locally to high yield losses over 50%, and is estimated to affect 5%–10% of the acreage in Europe and over 24% in the United States (Büttner et al., 2004; Jacobsen, 2006). Although partially *Rhizoctonia*-resistant or -tolerant sugar beet cultivars are commercially available, they usually are less productive or lack resistance/tolerance towards other diseases (Jacobsen, 2006). Therefore, microbiome-based disease management may be an interesting alternative in the future. Microbiomes can be managed either directly by applying microbiome transplants; single or mixed microbes with bioactive properties; or microbiota-active metabolites, or indirectly by changing environmental conditions in a way that microbiomes also shift their structure and function from dysbiosis into a healthy state (Berg, 2009). One possible approach to directly shape rhizosphere communities is using vermicompost as microbiome transplant. Vermicompost is a biofertilizing substrate produced by earthworms, which are one of the key taxa for soil functionality.
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sand:vermiculite 3:1:1 (v/v) mixture, further denoted as

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Rhizoctonia solani

Rhizoctonia-
tolerant and -susceptible cultivars enrich different bacterial taxa

in their rhizosphere, IV) Rhizoctonia-tolerant and -susceptible

cultivars enrich similar taxa from different substrates, and V) 

Rhizoctonia tolerance can be mediated in susceptible cultivars by

seed treatment with bacterial biocontrol agents. Our aim was to

investigate bacterial communities in sugar beet and correlate

them to the variables cultivar, seed origin, Rhizoctonia

susceptibility and growth substrate.

MATERIAL AND METHODS

Experimental Setup

We analysed the naturally composed seed microbiome (“seed”),

the roots of soilless germinated seedlings (“in vitro”) and roots of

seedlings grown in different substrates (“in vivo”). Five sugar beet

cultivars were chosen based on their phenotypic characteristics

regarding the susceptibility towards the fungal phytopathogen

Rhizoctonia solani (J.G.Kühn), the causative agent of late root rot

in sugar beet; two susceptible cultivars [BELLADONNA KWS

(BEL), BERETTA KWS (BER)], one moderately tolerant

[ISABELLA KWS (ISA)] and two cultivars with partial

resistance and tolerance [LAETITIA KWS (LAE)]; MATTEA

(MAT); Rhizoctonia-tolerant cultivars]. Seeds of every

cultivar were provided by KWS SAAT SE & Co. KGaA

(Einbeck, Germany) from two different seed production sites,

Italy (Region Emilia Romagna). All samples were repeated

4 times.

For seed samples, 20 seeds per replicate were washed three times

with sterile distilled water and activated for 4 h in 2 ml water. Sugar

beet seeds were not surface-sterilized to simulate field-like

conditions. For soilless cultivated sugar beet seedlings (“in vitro”),
eight activated seeds were placed in germination pouches (Mega

International, Newport, MN, USA) with two pouches per replicate.

To avoid decoupling the rhizosphere microbiome from plant

selection and to avoid training effects from agricultural field

management including chemical compounds like fertilizer,

pesticides, insecticides, and disseminated gene pools, we used

potting soil instead of field soil for rhizosphere sampling. Nine

activated seeds were placed in 7 cm × 7 cm × 13 cm pots with a soil:

sand:vermiculite 3:1 (v/v) mixture, further denoted as “potting

soil” (for details see Supplementary Table 1). To investigate the

effects of natural-based biofertilizer on rhizosphere diversity, the

setup was repeated for BEL, BER, LAE, and MAT using a natural

product based on earthworm casts (Supplementary Table 1),

further denoted as “vermicompost”. Vermicompost was used

either as sole substrate or as amendment to potting soil. For the

latter approach, ca. 15 g vermicompost was folded in the upper few

centimetres of the potting soil before seeds were planted. Growing

conditions were 23°C, 60% humidity and 16:8 h light/dark

conditions for 2 weeks in all approaches.

DNA Extraction

Activated seeds and “in vitro” roots weighed and grounded with

2 to 5 ml sterile 0.85% NaCl with mortar and pestle under sterile

conditions. Suspensions were centrifuged at 16,500 × g for

20 min at 4°C and resulting pellets were stored at -70°C for

further DNA extractions. DNA was extracted by mechanical

disruption and homogenization of the pellet using and a

FastPrep Instrument for 30 s at 5.0 ms⁻¹ and FastDNA Spin

Kit for Soil (MP Biomedicals, Illkirch, France). DNA was purified

using GeneClean Turbo Kit (MP Biomedicals, Illkirch, France) to

remove humic acids. Extracted DNA was treated with RNase

(0.02 ng μl⁻¹) for 5 min at 65°C to obtain the template for PCR

amplification of 16S rRNA genes from total community DNA.

Isolates From Sugar Beet and Vermicompost

Bacterial strains were isolated from vermicompost and sugar beet

rhizospheres to compare culture-dependent and culture-

independent results for the bacterial communities. For

vermicompost and potting soil, 1 g substrate was diluted in

9 ml sterile 0.85% NaCl, vortexed, centrifuged at 16,500 g and

4°C for 20 min. For rhizosphere, supernatants of the suspensions

used for amplicon sequencing were used. Bacteria were grown on

R2A (Roth, Karlsruhe, Germany) at 30°C for 48 h. Clean CFUs

were randomly chosen, isolated and grown on NA (Sifin GmbH,

Berlin, Germany). Bacterial DNA was extracted using a “quick

and dirty” protocol using a microwave for cell disruption;

bacterial material was transferred to a 1.5 ml Eppendorf and

rayed by a microwave for 3 min with closed lid, 30 μl TE buffer

was added, vortexed and centrifuged at 16,500 x g for 2 min.

Supernatant was used as template for a PCR using the bacterial

universal primer pair 27F/1492R according to Lane et al. (1991).

Amplifications were confirmed by gel electrophoresis in 1x TAE

(0.8% Agarose). PCR-products were purified using Wizard SV

Gel and PCR Clean-Up System (Promega, Madison, WI, USA).

Sequencing was conducted by LGC Genomics (Berlin,

Germany). Resulting sequences were quality-checked using Seq

Scanner 2 (Applied Biosystems) and identified with BLAST

(https://blast.ncbi.nlm.nih.gov/) using refseq_rna and/or

nr/nt database.

Amplicon Sequencing

The hypervariable V4 region of the 16S rRNA gene was amplified

according to the protocol of Caporaso et al. (2011) using the

primer pair 515f and 806r including Illumina cell flow adapters

and sample-specific barcodes. Peptide nucleic acid (PNA) PCR
clamps (PNA Bio, Newbury Park, USA) were used to reduce plastid and mitochondrial 16S contamination (Lundberg et al., 2013). The PCR mix (30 µl) contained 1 × Taq&Go (MP Biomedicals, Illkirch, France), 0.2 mM of each primer, 1.5 µM activated (55°C for 5 min) PNA mix (1:1 mPNA:pPNA) and 1 µl template DNA. PCR conditions were 96°C for 5 min; 30 cycles of 96°C for 1 min, 78°C for 5 s, 54°C for 1 min, 74°C for 1 min; 74°C for 10 min). Amplifications of the resulting 168 samples were confirmed by gel electrophoresis in 1x TAE (0.8% Agarose). PCR-products were purified using Wizard SV Gel and PCR Clean-Up System (Promega, Madison, WI, USA). DNA concentrations were measured using Nanodrop 2000 (Thermo Scientific, Wilmington, USA) and equimolar aliquots of all samples were pooled for amplicon sequencing using an Illumina MiSeq v2 (250 bp paired end) platform conducted by LGC Genomics (Berlin, Germany).

Bioinformatics

Pre-processing and analysis of the sequencing data was performed in QIIME2 v. 2019.10 (Bolyen et al., 2019) and QIIME v. 1.9.1 (Caporaso et al., 2010). Paired sample reads were demultiplexed and primers were trimmed from sequences using cutadapt (Martin, 2011). Since a considerable amount of forward and reverse sequences were flipped, reverse reads were also trimmed to forward primers and vice versa, and the resulting data from both trimming steps was combined. The sequences were denoised using DADA2 (Callahan et al., 2016), aligned with MAFFT (Katoh, 2002) and their phylogeny was constructed with fasttree2 (Price et al., 2010). Taxonomy was assigned with VSEARCH algorithm (Bogner et al., 2016) and Silva 132 99% consensus database (Quast et al., 2013). Mitochondrial, plastid DNA, and taxonomically unassignable sequences were removed from table and representative sequences. Thus, the datasets contained 32,140 amplicon sequences variants (ASV) and read numbers ranging from 254 to 545,913 reads per sample. Seeds and in vitro roots were separately analyzed from rhizosphere and soil. Alpha (Shannon, observed OTUs and evenness) and beta diversity indices (Bray-Curtis dissimilarity, weighted UniFrac) were calculated in QIIME2 and visualized using the q2_emperor plugin (Vázquez-Baeza et al., 2013). Differences in alpha and beta diversity indices were calculated using Kruskal-Wallis and pairwise PERMANOVA implemented in QIIME2. To evaluate taxa that differ significantly regarding cultivar and Rhizoctonia tolerance, samples of seed, in vitro, and in vivo roots were rarefied according to their minimum read number [1,000, 2,400 (losing 2 samples) and 15,000, respectively]. Samples for ISA were removed in Rhizoctonia tolerance testing to perform pairwise comparisons. The remaining samples were collapsed to genus level and compared using the QIIME1 plugin group_significance.py with Kruskal-Wallis test (cultivar, substrate) and nonparametric t-test (Rhizoctonia tolerance, seed origin). Differences between Bray-Curtis distances and weighted UniFrac distances of seeds and roots were tested using the adonis (Anderson, 2001) command of the R script vegan 2.5 (Oksanen et al., 2018) implemented in QIIME2.

To evaluate number proportion of surviving seed endophytes, seed and in vitro root samples were compared on ASV level. Intersection of communities of seed, soil, and rhizosphere were compared on genus-level with mean values of relative sequences.

Field Design

Field trials were conducted in the course of efficacy tests in different years for the sugar beet cultivars BERETTA KWS (Rhizoctonia-susceptible, growing season 2009 and 2010), ISABELLA KWS (moderately tolerant, growing season 2016), and MATTEA KWS (Rhizoctonia-tolerant, growing season 2013). However, all field trials were performed under usual production conditions by local farmers. All sugar beet (Beta vulgaris L.) seed materials were generated and evaluated by KWS SAAT SE & Co. KGaA (Einbeck, Germany). Bacterial treatment was performed with a consortium of Pseudomonas poae RE*1-1-14 obtained from sugar beet (Zachow et al., 2010), Pseudomonas brassicacearum L13-6-12 (Grosch et al., 2005), and Serratia plymuthica 3Re4-18 (Berg et al., 2005) isolated from potato, which all show antagonistic activity towards Rhizoctonia solani. Formulations were prepared by Biotenzz GmbH (Graz, Austria) for integration in the commercial seed coating. Sugar beet seeds of R. solani-susceptible cultivars were routinely coated with the fungicides Thiram® (Cheminova Deutschland GmbH, Germany), Hymexazol® (Mitsui Chemicals, Tokyo, Japan) and the insecticides Imidacloprid (Gaucho®, Bayer CropScience, Leverkusen, Germany), Chlothianidin (Poncho®, Bayer CropScience, Leverkusen, Germany), Thiathemoxaxom (Syngenta Crop Protection, Basel, Switzerland), and Tefluthrin (Syngenta Crop Protection, Basel, Switzerland) respectively. The two test fields (“Kasten”: 48°42’48.6”N 13°04’34.5”E and “Tabershausen”: 48°44’24.5”N 12°52’55.2”E) for the trials were located in the main growing area in Lower Bavaria, Germany. R. solani-infested barley kernels were used for artificial inoculation of the soil (60 kg/ha). Control plants were commercially prepared without microbial inoculants. Field trials were conducted in a randomized block design with four replicates per variant, each with six rows containing 30 plants per row. The 16S rRNA sequence of Pseudomonas poae RE*1-1-14 (NCBI Reference Sequence: NC_020209.1), Pseudomonas brassicacearum L13-6-12 (CP014693.1), and Serratia plymuthica 3Re4-18 (CP012097.1) was extracted from their reference genome and cross-checked with the amplicon dataset to evaluate presence or absence in our amplicon data samples.

RESULTS

Seeds Harbor Cultivar-Specific Bacterial Communities Strongly Influenced by Propagation Sites

Sugar beet seeds of five different cultivars originating from two different seed production sites contained a total of 1001 amplicon
sequences variants (ASVs). For seed samples, read numbers ranged from 1,062 to 152,738 reads per sample (for a summary, see Supplementary Table 2). Seeds of all cultivars share a core of 121 ASVs, which accounts for 80%–91% \( r^2 \) (relative abundance) in the communities, while partially shared ASVs account for 4%–19% (Figure 1A); ASVs that are unique to a cultivar account for less than 4% \( r^2 \) (See also Supplementary Table 3). The seed endophyte community is dominated by *Pseudomonas*, *Pantoea*, and *Paenibacillus* (Figure 1B). Significant differences in alpha diversity indices (Shannon, Pielou’s evenness, observed OTUs) were not observed in seeds, except BER shows higher values for evenness than BEL and ISA (both \( p = 0.027 \)). On the contrary, PERMANOVA results for differences in beta diversity indices (Bray-Curtis dissimilarity, weighted UniFrac distances) revealed to be largely significant (except: weighted UniFrac of BER:LAE; ISA:LAE). Main drivers of cultivar-dependent differences (\( r^2 > 1 \% \)) in seeds were *Pseudomonas* spp., *Paenibacillus* spp. and *Massilia* spp. (\( p < 0.05 \); Bonferroni-corrected).

Seed origin was an important variable for alpha diversity (Figures 1C, 2A–C), pairwise comparison within the same cultivars revealed significant differences (pairwise Kruskal-Wallis) for observed OTUs (all) and Shannon index (except LAE). Additionally, all cultivars differed significantly (\( p < 0.05 \)) due to seed origin in Bray-Curtis and weighted UniFrac distances. Adonis test of Bray-Curtis and weighted UniFrac distances revealed the factors “Cultivar” and “Seed origin” explain more than 50% of the variance (Supplementary Table 4), with “Cultivar” being the more important one (\( R^2 = 0.33 \) and \( R^2 = 0.32 \), respectively; Pr(\( >F \)) = 0.001). Key genera that significantly differ due to seed origin are not the same for the different cultivars (Supplementary Table 5). When merging all seed data, *Sphingomonas*, *Methylobacterium* and unidentified *Sericytachromatia* show higher \( r^2 \)-values in Italian seeds, while *Saccharibacillus*, *Kosakonia*, and *Erwinia* have higher proportion in French seeds; this is also indicated by PCoA-biplot of Bray-Curtis distances (Figure 2B).

**Dominating Seed Endophytes Survive Host Germination**

To assess the proportion of alive bacterial communities within seeds, 20 seeds each cultivar were cultivated under *in vitro* conditions in soilless germination pouches. The *in vitro* rhizosphere contained 397 ASVs, represented by 2,533,778 reads, with read numbers ranging from 254 to 137,909 reads per sample. Depending on cultivar, ASVs accounting for 63%–83% \( r^2 \) found in the seeds could be confirmed to be present in soilless roots (*“in vitro”*, Figure 3A).

Dominant phyla in soilless roots are *Proteobacteria* (range 75%–91% \( r^2 \), depending on cultivar), *Actinobacteria* (1%–18% \( r^2 \)), and *Firmicutes* (2%–22% \( r^2 \)). *Enterobacteriaceae*, mainly represented by *Kosakonia* (19% \( r^2 \)) and *Pantoea* (18% \( r^2 \)) account for 40% \( r^2 \) in soilless roots, although *Methylobacterium* (28% \( r^2 \)) is the most frequent genus.

The number and proportion of ASVs of successfully colonizing bacteria differ between cultivars (\( n = 45–70, \Delta 14%–22\% \) of corresponding seed ASVs), but represent dominating taxa of seed endophytes. Tendentially, a higher proportion of seed

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**FIGURE 1** Overview on microbial community of sugar beet seeds. (A) number (VENN diagram) and corresponding abundance (pie charts) of amplicon sequence variants (ASVs) found in sugar beet seeds. Light green: seed core, found in all cultivars; dark green: cultivar-specific ASVs; orange: partially shared ASVs. (B) OTUs accounting for >1% relative abundance in the sugar beet seed microbial community on phylum and genus level. C: Alpha diversity indices of seed community; * = significantly differing according to Kruskal-Wallis test depending on seed origin (see also Figure 2).
FIGURE 2 | Differences in the bacterial community in sugar beet cultivars’ seeds due to origin. (A) Seed microbial community on genus level. All seeds are dominated to a different extent by Proteobacteria, especially *Pseudomonas* (a) and *Pantoea* (b). (B) PCoA-biplot of Bray-Curtis distance on genus level. Seed communities clearly cluster according to seed origin; spheres: cv. BELLADONNA; diamonds: cv. BERETTA; triangles: cv. ISABELLA; squares: cv. LAETITIA; stars: cv. MATTEA. (C) comparing Shannon diversity in seeds within the same cultivar due to seed origin; all seeds originating from France show a higher alpha diversity except cv. LAETITIA; “∗” = significantly differing according to Kruskal-Wallis test depending on seed origin.
endophytes survive in soilless roots of Rhizoctonia-tolerant cultivars (14%–15% vs. 16%–21% of corresponding seed ASVs; Figure 3A). In all cultivars investigated, in vitro roots display a lower alpha diversity and a lower evenness (Figure 3B) than the corresponding seeds. ASVs representing up to 14% r² (for LAE) in soilless roots were found to be unique and thus of unknown origin. In total, 230 ASVs unique to in vitro roots were found across all cultivars with 12 ASVs accounting for r² >1% per sample. These ASVs belong to the genera Curtobacterium, Bacillus, Pseudomonas, and Methylobacterium, taxa that are frequently present in both seeds and in vitro roots. Cultivar-dependent differences for alpha diversity (Shannon) were not found, and beta diversity indices (Bray-Curtis, weighted UniFrac) were only significantly different between BEL and MAT.

The Ingredients of the Rhizosphere: Soil and Seed Bacteria

The in vivo root community clearly differs from both soil and seed communities and is dominated by Proteobacteria (26% r²), Acidobacteria (16% r²), Actinobacteria (13% r²), Chloroflexi (10% r²), and Planctomycetes (7% r²; Figure 4A). When the dataset was collapsed to the highest taxonomic annotation, the dominant taxa were Acidobacteria Subgroup 6 (7.0% r²), the archaeal family Nitrososphaeraceae (3.9% r²), and the genus Pyrinomonadaceae RB41 (3.9% r²). Differences in root beta diversity due to seed origin were not significant. The bacterial core community of in vivo roots comprised of 3,228 ASVs, representing 82%–88% r² depending on cultivar. Cultivar-specific ASVs account for 2.2%–3.1% of the abundance (Supplementary Table 3).

The bigger part of the root community is originating from soil. Genera accounting for 53.6% are exclusively assembled from soil and another 26.8% of the root community comprises taxa that are found in both seeds and soil (Figure 4B). Only a small proportion of the seed and soil microbiome (5.3% and 7.5%, respectively) is unique to their corresponding habitats, while we find 417 genera accounting for 15.25% of the rhizosphere microbiome to be unique. The key taxa found in seeds appear in lower relative abundances in rhizosphere. From the genera that exclusively originate from seeds, four are found in >50-fold r²-values in root samples; 41 of soil-derived taxa are >50-fold increased (Supplementary Table 6).

Interestingly, Archaea were found in high abundances (up to 14.1% r²) in the root and vermicompost-associated communities, with Nitrososphaeraceae (ad Thaumarchaeota) as dominant taxa (in average representing 91% of all archaeal taxa).

Indicator Bacteria for Rhizoctonia-Tolerant Cultivars

ASVs, that are exclusively found in seeds of Rhizoctonia-tolerant cultivars (n = 20) account for 0.83% r² (LAE) and 0.88% r² (MAT). When combining seeds, potting soil and roots samples, adonis test revealed the factor “habitat” to be the most important variable explaining the variance for Bray-Curtis and weighted UniFrac distances (R² = 0.421 and R² = 0.725, respectively). The factors

![Diagram](image-url)
“seed origin” and “Rhizoctonia-tolerance” explain approximately the same amount of variance in both Bray-Curtis ($R^2 = 0.029$ and $R^2 = 0.027$) and weighted UniFrac distances [$R^2 = 0.014$ and $R^2 = 0.015$, respectively (Supplementary Table 7)]. When exclusively testing root samples (seed origin = Italy) grown in the three different substrates, the factor “Rhizoctonia-tolerance” explains a higher percentage of variance than the factor “Cultivar” for both beta diversity indices (Supplementary Table 8).

Taxa significantly and positively correlated with Rhizoctonia-tolerance in seeds are Halomonas spp., Paenibacillus spp, Enterobacter spp., and Kosakonia spp. (Figure 5A), while Massilia spp. is higher in seeds of susceptible cultivars. In soilless roots, Firmicutes, mainly represented by Paenibacillus (except for LAE: Pullulanibacillus and Bacillus), are higher abundant in Rhizoctonia-tolerant cultivars than in susceptible ones, and vice versa for Actinobacteria (Supplementary Table 9), mainly represented by Curtobacterium. For the root samples, clustering of samples in PCoA plots due to Rhizoctonia-tolerance is less pronounced than in seeds (Figure 5B). A general trend across seeds, in vitro, and in vivo root samples is the higher abundance of Firmicutes in Rhizoctonia-tolerant cultivars.

Root communities of Rhizoctonia-tolerant cultivars show a trend (non-significant) towards higher $r^3$-values of Gaiellales (ad Actinobacteria) and Rhizobiales (ad Proteobacteria). $r^3$-values of Bacteroidetes and Verrucomicrobia are significantly ($p = 0.02$) higher in roots of susceptible.

Genera significantly higher in Rhizoctonia-tolerant cultivars differ according to used substrates (Figure 5C). However, the genus Kosakonia, which is one of the taxa correlated with Rhizoctonia tolerance in seeds, tends to be higher abundant in Rhizoctonia-tolerant root samples across all substrates.

**The Impact of Substrates on Root Microbiota**

Root samples of BEL, BER, LAE, and MAT grown in the three different substrates (potting soil, vermicompost and mixtures) were merged in a data subset. Differences in the overall composition of root microbiomes are already apparent in high taxonomic ranks. Vermicompost addition to potting soil highly increases $r^3$ of Proteobacteria across all cultivars. Firmicutes and Bacillus spp. in particular are found in high abundance in pure vermicompost (Figure 6A). Results for pairwise Kruskal-Wallis test between “substrate” groups were filtered for genera that a) have $r^3 > 1\%$ in at least one of the substrate groups and b) show high $r^3$-values in one of the pure substrates and intermediate in mix of vermicompost-treated potting soil. Six genera matching these prerequisites were thus shown to be gradually enriched with increased use of vermicompost (Figure 6A).

The factor “substrate” explains around 68% of the variance in both Bray-Curtis and weighted UniFrac distances (Supplementary Table 8). In PCoA visualization plots, samples clearly cluster due to substrate (Figure 6B).

**Cultivation-Dependent vs. -Independent Microbiome Assessment**

Bacteria were isolated to find bioactive strains and compare cultivation-dependent to -independent approaches. We successfully isolated and identified 339 bacterial strains belonging to at
FIGURE 5 | Differences in bacterial root communities between Rhizoctonia solani-susceptible (violet), effective (red) and –tolerant (orange) cultivars (A) PCoA-biplot of Bray-Curtis dissimilarities in seeds on genus level; spheres: cv. BELLADONNA; diamonds: cv. BERETTA; triangles: cv. ISABELLA; squares: cv. LAETITIA; stars: cv. MATTEA. Note, that PCoA axes 2 and 4 separate the communities based on Rhizoctonia tolerance, while axes 1 and 2 separate them due to their origin (Figure 2B). For the visualization of the other combinations of PCoA axes, see Supplementary Figure 1. (B) PCoA-biplot of Bray-Curtis dissimilarities in roots on genus level; (C) Taxa significantly enriched in Rhizoctonia tolerance cultivars (genus level) in seeds and roots grown in potting soil, vermicompost-treated potting soil and vermicompost.

FIGURE 6 | Influence of different substrates on bacterial root communities in sugar beet. (A) Composition of bacterial root communities in different substrates on phylum level. Bacterial genera displayed that are significantly different (Bonferroni-corrected p < 0.05) between substrates, have >1% mean relative abundance and have intermediate abundance in VC-PS samples. *: unique for potting soil; #: unique for vermicompost; PS: potting soil; VC-PS: vermicompost-treated potting soil; VC: vermicompost. (B) PCoA-plot of Bray-Curtis dissimilarities of root samples grown in different substrates, only samples with seed origin from Italy; spheres: cv. BELLADONNA; diamonds: cv. BERETTA; squares: cv. LAETITIA; stars: cv. MATTEA.
least 137 different species from pure vermicompost (n = 192; 73 species), roots grown in vermicompost-treated potting soil (n = 95; 41 species) and roots grown in pure vermicompost (n = 112; 56 species, Supplementary Table 10). The most frequently isolated species associated with vermicompost was Agromyces flavus. Additionally, we found 62 strains of at least 20 different Bacillus species, with B. firmus (n = 15) being the most frequently isolated species. Furthermore, the diversity of Microbacterium spp. (n = 19; six species) and Streptomyces spp. (n = 12; 9 species) is worth mentioning.

Results of cultivation-dependent assessment clearly differ from amplicon data. On order level, a third up to half of all orders present in the amplicon data could be cultivated. When comparing r²-values between cultivated strains and amplicon data, the most apparent difference is the high proportion of Micrococcales in the isolate dataset across all three groups. In contrast, sequences of Micrococcales account for r² < 3% to the amplicon dataset. (Figure 7). The dominance of Bacillales in vermicompost and Betaproteobacteriales in roots grown in soil-vermicompost mix is confirmed by both amplicon and cultivation data. Rhizobiales reached higher r²-values in the amplicon dataset across all groups.

**Inducing Rhizoctonia Tolerance Using a Consortium Containing Ps. poae Re*1-1-14**

Sugar beet seeds with standard fungicide and were compared to a combined treatment with both bacterial consortium and fungicide treatment. The percental change in RI, total number of beets and number of healthy beets due to bacterial treatment were compared. Statistical pairwise comparisons did not result in significant differences (p > 0.05), most likely due to the patchy infestations of Rhizoctonia observed in the field and subsequently high standard deviation in the dataset. Nevertheless, trends towards different responses due to cultivar were observed: the application of the bacterial consortia showed strongest effects in the Rhizoctonia-susceptible cultivar, especially at the field with higher soil humidity and thus higher pathogen pressure (“Kasten”). Under these conditions a mean RI decrease of 26% as well as an increase in the number of healthy sugar beets of 65.4% relative to standard treatment could be achieved (Supplementary Figure 2) in susceptible cultivars. For tolerant or moderately tolerant cultivars, positive effects regarding RI and number of healthy beets were less pronounced or not observed. In average, the total number of sugar beets at the time of harvest increased by 5% using bacterial treatment.

**DISCUSSION**

We analyzed microbial communities in seed, roots and corresponding soil to investigate sugar beet microbiota assembly and composition. We found novel aspects, which can be translated to manage the plant microbiome. The bacterial root community of sugar beet derives from both seed and soil communities. Seeds of all sugar beet cultivars were highly colonized by bacteria and carried a selective core of the sugar beet microbiome, which contributed significantly to the rhizosphere assembly. We found relatively high alpha diversity values for the sugar beet seeds, with results comparable to pumpkin seeds (Adam et al., 2018) but significantly lower than in sugar beets at harvest (Kusstatscher et al., 2019). The high proportion of Proteobacteria, Actinobacteria, Firmicutes and Bacteroidetes found in sugar beet seeds is quite typical for seed microbiota (reviewed by Nelson, 2018). However, especially Enterobacteriaceae were identified as important component in seeds. For instance, Enterobacteriaceae and in particular Klebsiella, dominate in pumpkin seeds (Adam et al., 2018). Pantoea, one of the key taxa in sugar beet seeds, was also found to be dominant in Brassica seeds (Barret et al., 2015a).
except for oilseed rape (Rybakova et al., 2017). *Pantoea* comprises versatile lifestyles, including plant pathogens, plant growth promoters as well as strains commercially produced for biocontrol of phytopathogens. Thus, *Pantoea* is a model group for niche-specific adaptations (Walterson and Stavrinides, 2015). Furthermore, abundance of some *Enterobacteriaceae* (Kosakonia and Enterobacter) together with *Paenibacillus* are correlated with *Rhizoctonia* tolerance (Figures 5A, C) in sugar beet seeds. *Paenibacillus* is known to contain species with plant growth-promoting and pathogen-antagonistic properties (reviewed by Rybakova et al., 2016). Similarly, some *Enterobacter* species show antagonistic properties against *Rhizoctonia* (e.g. Abdeljalil et al., 2016). *Kosakonia radicinitrans* is known for both plant growth promotion and indirectly reducing pest pressure in *Arabidopsis thaliana* (Brock et al., 2018). Nevertheless, bacterial seed communities rather cluster due to origin (Figure 2B, displaying PCoA axes 1 and 2) than to *Rhizoctonia* tolerance (Figure 5B, displaying PCoA axes 2 and 4) of the cultivars.

One interesting fact is the higher abundance of different groups of *Firmicutes* (Pullulanibacillus in LAE, Paenibacillus in MAT, see Supplementary Table 9) in seeds of *Rhizoctonia*-tolerant cultivars. This indicates that the function provided by several different groups of bacteria is more important for the plant-microbe interaction than their exact taxonomic position; taxa significantly solely enriched in *Rhizoctonia*-tolerant cultivars account for less than 1% of the total abundance. Although our study suggests which taxa could be correlated with *Rhizoctonia* tolerance, we are aware that five genotypes are not sufficient to suggest a general trend for microbial shifts due to breeding. This also applies for our results focusing on general aspects of the microbial community differences due to seed origin, cultivar and genotype as well as the phenotype (e.g. root architecture) of the host plant—thus the more important factor explaining the seed communities’ variance.

The drop in microbial communities’ diversity during germination (Figure 3B) is usually interpreted as selective force exerted by the seedling, favoring fast-growing r-strategists like *Pseudomonas, Bacillus* or *Trichoderma* (Barret et al., 2015a). In our dataset, *Methylobacterium* dominated soilless roots and were regarded to be typical K-strategists due to their ability to metabolize C1-compounds. Therefore, we suggest the indirect selective force of the germinating host plants rather to favor seed endophytes that are adapted to their hosts’ specific genotype (Berg and Raaijmakers, 2018) or the present cultivation conditions (Bergna et al., 2018). Given that this turnover of the seed core is proportionally more or less stable over generations, there are niches in the seed microbiome that can be colonized with bacteria enriched from soil, shaping the seed microbiome of the next generations’ seeds (Bergna et al., 2018).

The soil microbiome can influence the rhizosphere microbiome composition of the seedlings and thereof influences the host (Nelson, 2018). We confirm soil as the main source of diversity in sugar beet rhizosphere. Nevertheless, a considerable amount of rhizosphere-inhabiting bacteria exclusively originates from seeds, and 26.8% of all bacterial genera could be provided by either bacterial soil or seed endophyte communities. Interestingly, 417 genera accounting for 15.3% of the rhizosphere community could not be traced back to either seed or soil (Figure 4B). The majority of these genera have low values (<0.5%). We regard these members of the root microbiome to be enriched in rhizosphere but under the detection threshold in soil and seed samples. This explains why alpha diversity in roots (Shannon H = 9.61 ± 0.56) is significantly (p = 0.02) higher than in soil samples (Shannon H = 8.94 ± 0.05). The differences in microbial communities due to seed origin, cultivar and *Rhizoctonia* tolerance are more pronounced in seeds than in roots. *Roseiflexaceae* and *Anaerolineaceae*, which are significantly enriched in *Rhizoctonia*-tolerant cultivars, are worldwide distributed taxa (GBIF.org, 2020), found in water as well as soil habitats and mainly comprise yet monotypic genera. The genus *Microvirga* on the other hand is frequently found in root nodules of legumes in temperate climates (e.g. Msadak et al., 2017). Seed origin was an important variable for bacterial seed communities. One should consider that crops are usually planted across large geographical areas and location-dependent bacterial communities were already revealed in maize (Johnston-Monje and Raizada, 2013) and common beans (Kladtke et al., 2016). Although community differences due to the origin of seeds in both alpha and beta seed diversity indices are obvious and apply to several different bacterial taxa (Figures 2A–C), the cultivars—and thus the genotype as well as the phenotype (e.g. root architecture) of the host plant—is the more important factor explaining the seed communities’ variance.

**Figure 2A** shows the highest seed alpha diversity across all tested cultivars. Although we know that bacterial alpha diversity is usually correlated with pathogen tolerance (van Elsas et al., 2012), a high diversity may not be necessary if members of the microbial community provide all important functions to the plant-microbe-pathogen interaction. We hypothesize this to be the case in cv. ISABELLA and cv. MATTEA since the majority of seed endo- and epiphytes survive in the germinated seedlings.
Considering earthworms, their ecosystem services and their microbes in agricultural practices holds a big potential for the agriculture of the future (Singh et al., 2020). The microbial community of vermicompost, as a product obtained mainly from organic litter earthworm casts, clearly differs from potting soil, with especially higher proportions of Bacillus (5.22 vs. 0.02%). High abundance of Bacillus spp. relates to the processing of organic matter in the gut of earthworms. In the course of digestion, the microbial community is exposed to several changes in conditions, including pH neutralization, higher water content, complete anoxia, secretion of digestive enzymes and enrichment of organic compounds including fermentation products from other microbes (reviewed by Drake and Horn, 2007). These conditions on the one hand favor facultative anaerobic or aerobic bacteria that are able to form endospores (Drake and Horn, 2007), and on the other hand activate endospores of Bacillus (Fischer et al., 1997). The genus Bacillus contains several species that are known for direct as well as indirect phytopathogen antagonism and plant growth-promoting effects (e.g. Pérez-Garcia et al., 2011). We observed higher abundance of Firmicutes—in particular the genus Bacillus—in roots of Rhizoctonia-tolerant cultivars grown in untreated and vermicompost-treated potting soil. When using pure vermicompost as substrate, abundance of Bacillus was comparably high (4.1%–6.2% 19) in all cultivars. We interpret this as specific enrichment of pathogen-antagonistic bacteria by Rhizoctonia-tolerant sugar beet cultivars from the substrate. Since the abundance of Firmicutes is generally higher in vermicompost than in potting soil, enrichment in the rhizosphere may not be necessary to achieve pathogen suppression. Some basic information on the microbial community when using vermicompost as substrate are concordant in both cultivation-dependent and –independent approaches. This includes the dominance of Bacillales in vermicompost-associated samples and the dominance of Betaproteobacteria in vermicompost-treated potting soil samples (Figure 7).

The effect of bacterial control agents in field trials was strongest for the Rhizoctonia-susceptible cultivar BERITTA, indicating a strong influence of the microbial community in the early stage of rhizosphere establishment. Applying bacterial control agents to Rhizoctonia-tolerant cultivars did not result in further increased, but partially showed slightly decreased health parameters. This may indicate interference of the applied control agents with native root bacteria. The bacterial treatment consisted of three strains, of which Pseudomonas poae RE′1-1-14 was originally isolated from sugar beet. This strain shows an interesting pattern across our dataset were found in sugar beet endosphere before (Zachow et al., 2008; Zachow et al., 2010). RE′1-1-14 was initially selected in a screening for antagonistic strains against several different sugar beet pathogens (Phoma betae, R. solani AG2-2IIIIB, R. solani AG4 and Sclerotium rolfsii; Zachow et al., 2010). Despite the specific ASV was not exclusively found in Rhizoctonia-tolerant cultivars, Ps. poae seems to be an integral part of Rhizoctonia-tolerant cultivars and provide important plant-protecting functions to the seedling. Interestingly, Ps. poae is more frequently found in sugar beets originated from France, where sugar beets are grown on much larger geographical areas (EUROSTAT, 2019) and historically for a longer time span compared to Italian acreages. These factors are known to favor the development of soil communities suppressive to certain pathogens (Mazzola, 2002; Eberlein et al., 2020). Continuous cropping of sugar beet is known to influence the field soil community, accelerating the abundance of different taxa in the microbial soil community over time (Huang et al., 2020). We found partially cultivar-specific differences between root communities. Thus, continuous cropping of self-reproduced genotypes may shape both bacterial soil and seed endophyte communities, potentially leading to a cultivar-dependent enrichment of taxa that are beneficial and/or usually uncommon in seeds.

CONCLUSIONS

Overall, found differences in the microbiomes of Rhizoctonia-susceptible and -tolerant sugar beet cultivars across various life stages. In seeds, the genus Kosakonia may play a role in pathogen tolerance. Seed communities in sugar beet differ due to seed origin and cultivar; differences are more pronounced than in seedling roots and rhizospheres, although substrate heavily determine root community structure. Further investigation with a higher number of different Rhizoctonia-susceptible and tolerant cultivars are needed for confirming general trends in sugar beet-associated bacterial communities with Rhizoctonia-tolerance. This investigation can be explored in more detail by implementing crop breeding strategies that include the trait ”plant microbiome”. Future strategies for sustainable agriculture might be able to include microbiome management.: 1) Selection of seed production sites could include soil microbiome analysis as seeds take up sets of microbes differently involved in stress response in the next generation. 2) The crop/plant microbiome can be adjusted via seed treatments that add further microorganisms to the soil 3). The entire soil microbiome can be managed by microbiome transplants.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://www.ebi.ac.uk/ena, PRJNA299370 https://www.ebi.ac.uk/ena, PRJEB37140.

AUTHOR CONTRIBUTIONS

AW, GB, CZ, and HM drafted the manuscript. CZ, AW, HM, AG, NT, RT, and GB planned the experimental setup. AW, CZ,
and HM performed molecular, microbial, and ad planta work. AW conducted bioinformatic analyses. AW, CZ, HM, GB, NT, and RT either planned and/or performed the field trials. All authors contributed to the article and approved the submitted version.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2020.560869/full#supplementary-material

**SUPPLEMENTARY FIGURE 1** | PCoA-biplot of sugar beet seed endophyte communities on genus level displaying Bray Curtis dissimilarities using PCoA axes 1 and 3 (A), 1 and 4 (B) and 2 and 3 (C).

**SUPPLEMENTARY FIGURE 2** | Results of field trials for relative in-/decrease of Rhizoctonia disease index (FI, according to Böttner et al., 2004), number of sugar beets and number of healthy sugar beets for the cultivars BER, ISA and MAT using a consortium of three bacterial antagonists additionally to standard fungicide treatment. Results are relative to sugar beet seeds with standard fungicide treatment. Comparisons did not result in significant differences but trends towards stronger positive effects in the Rhizoctonia-susceptible cultivar.
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The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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