Bacteria Associated to Plants Naturally Selected in a Historical PCB Polluted Soil Show Potential to Sustain Natural Attenuation

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The exploitation of the association between plants and microorganisms is a promising approach able to boost natural attenuation processes for soil clean-up in vast polluted areas characterized by mixed chemical contamination. We aimed to explore the selection of root-associated bacterial communities driven by different plant species spontaneously established in abandoned agricultural soils within a historical polluted site in north Italy. The site is highly contaminated by chlorinated persistent organic pollutants, mainly constituted by polychlorobiphenyls (PCBs), together with heavy metals and metalloids, in variable concentrations and uneven distribution. The overall structure of the non-vegetated and root-associated soil fractions bacterial communities was described by high-throughput sequencing of the 16S rRNA gene, and a collection of 165 rhizobacterial isolates able to use biphenyl as unique carbon source was assayed for plant growth promotion (PGP) traits and bioremediation potential. The results showed that the recruitment of specific bacterial communities in the root-associated soil fractions was driven by both soil fractions and plant species, explaining 21 and 18% of the total bacterial microbiome variation, respectively. PCR-based detection in the soil metagenome of bacterial \textit{bphA} gene, encoding for the biphenyl dioxygenase \(\alpha\) subunit, indicated that the soil in the site possesses metabolic traits linked to PCB degradation. Biphenyl-utilizing bacteria isolated from the rhizosphere of the three different plant species showed low phylogenetic diversity and well represented functional traits, in terms of PGP and bioremediation potential. On average, 72% of the strains harbored the \textit{bphA} gene and/or displayed catechol 2,3-dioxygenase activity, involved in aromatic ring cleavage. PGP traits, including 1-aminocyclopropane-1-carboxylic acid deaminase activity potentially associated to plant stress tolerance induction, were widely distributed among the isolates according to \textit{in vitro} assays. PGP tested \textit{in vivo} on tomato plants using eleven selected bacterial isolates, confirmed the promotion and protection potential of the rhizosphere bacteria. Different spontaneous plant species naturally
INTRODUCTION

Polychlorinated biphenyls (PCBs) are highly stable, hydrophobic and persistent organic pollutants. Due to their lipophilic nature, they bioaccumulate and biomagnify through the food web and can have a broad range of toxic effects on humans (Turri-Baldassarri et al., 2009; Letcher et al., 2010; Quinete et al., 2014; IARC, 2015). Physico-chemical remediation techniques of PCB polluted soils are not sustainable environmentally and economically for extended contaminations and the properties of these molecules make them recalcitrant to biodegradation, impairing the efficacy of bioremediation technologies. However, PCB pollution is a worldwide problem associated to their past production and utilization in industrial facilities (Passatore et al., 2014). PCBs strongly bind to the soil organic matter, resulting into low bioavailability both for plant uptake and microorganism metabolism (Sinkkonen and Paasivirta, 2000). Nonetheless, several studies showed that PCB-polluted soils host bacterial communities endowed with aerobic degradation abilities and harboring the biphenyl dioxygenase (bph) operon, the most studied and widespread pathway sustaining the aerobic biodegradation ability (Abraham et al., 2002; Leigh et al., 2006; Leigh et al., 2007; Kurzzawova et al., 2012). Plants establish mutual beneficial interactions with selected bacterial populations, which promote nutrient uptake and enhance stress tolerance against pollutants, such as PCBs, in turn decreasing their phytotoxicity (Ma et al., 2011; Abhilash et al., 2016). Root exudates, besides creating a selective ecological niche for specific bacteria, may contain molecules that can induce the expression of genes involved in the PCB degradation pathways, such as the bph operon, in turn stimulating PCB degradation in the rhizosphere (Toussaint et al., 2012; Pham et al., 2015). Rhizoremediation, the exploitation of such positive interactions between plants and bacteria for reclamation of polluted soils, represents a sustainable alternative for clean-up of recalcitrant organic pollutants in extended areas (Vergani et al., 2017). Soil natural attenuation potential therefore relies on the presence, diversity and activity of the resident microbiota, driven by the selecting activity of autochthonous plants.

In this work, we studied the phylogenetic and functional diversity of the microbiota associated to the root-soil fractions of the three spontaneous plant species Medicago sativa, Centaurea nigrescens, and Dactylis glomerata, naturally established in a former agricultural field located within the heavily polluted SIN (National Priority Sites) site Caffaro in north Italy. This is a large area polluted by the activity of the former Caffaro chemical factory and includes more than 100 ha of former agricultural areas (Di Guardo et al., 2017). It presents a mixed and uneven contamination of chlorinated persistent organic pollutants, mainly PCBs, heavy metals and metalloids and several other contaminants including dioxins, furans, tetra-chloromethane (Turri-Baldassarri et al., 2007; Di Guardo et al., 2017). PCBs were produced by Caffaro for 46 years and banned in Italy in 1984. To avoid pollutant exposure to humans, economic activities, including agriculture, have been banned in 2002 in the SIN Caffaro area. In the last 12 years only natural attenuation processes occurred, with the establishment of spontaneous plants resisting phytotoxic effects. Remediation strategies for such a large and complex site represent a challenge raising the interest for the soil self-depuration potential, which could be exploited and boosted by phyto-rhizoremediation approaches. A basic question is whether PCB degradation potential is present in the bacterial communities enriched by plants able to grow in the polluted soil, mediated by the selection force determined by the rhizosphere effect. We compared the bacterial communities in the root-associated soil fractions with that of the non-vegetated soil, through 16S rRNA gene phylogenomics and the amplification of the bphA gene, encoding for the α sub-unit of the biphenyl dioxygenase enzyme, a proxy for the PCB degradation ability (Iwai et al., 2010; Sylvestre, 2013; Vergani et al., 2017). By looking to the culturable bacterial fraction, we assessed whether bacteria represent a potentially exploitable resource for rhizoremediation purposes, by assessing their PGP capacity and PCB biodegradation potential.

MATERIALS AND METHODS

Site Description, Plants and Soil Sampling

Non-vegetated and root-associated soil fractions were collected in the SIN Caffaro, a site located in Northern Italy (Brescia municipality), within the sampling area A, a former agricultural grassland field that was previously chemically characterized and contained PCBs and other chlorinated pollutants in concentrations often exceeding the safety limits (Di Guardo et al., 2017). The root system of three spontaneous plant species was collected from triplicate specimens for each plant species. Plants were identified basing on their morphological traits as M. sativa L. (MS), C. nigrescens Willd. (CN), and D. glomerata L. (DG). These plant species were among the most widely widespread in the meadows present in the agricultural areas of the SIN Caffaro, according to a recent survey (Armiraglio et al., 2009). Non-vegetated bulk soil (B) was collected in triplicate in the same sampling station. Samples were transferred to the laboratories and the different soil fractions selected in a historical chronically polluted site showed to determine the enrichment of peculiar bacterial communities in the soil fractions associated to the roots. All the rhizosphere communities, nevertheless, hosted bacteria with degradation/detoxification and PGP potential, putatively sustaining the natural attenuation process.

Keywords: soil pollution, natural attenuation, bacterial diversity, rhizosphere, polychlorobiphenyls, plant growth promotion, bphA gene
(as defined below) were separated within 4 h from collection as described in Marasco et al. (2012). Soil fractions were stored at −20°C for molecular analyses and at 4°C for bacterial isolation.

**DNA Extraction and Analyses of the Bacterial Community Structure**

Total DNA extraction was performed from 0.25 g of each of the three non-vegetated bulk soil (B), root surrounding soil (S) and rhizosphere soil (R) replicates, using the PowerSoil DNA kit (MoBio) according to the manufacturer's protocol. Illumina tag screening of the V3–V4 hypervariable regions of the 16S rRNA gene was applied on DNA of R, S, and B triplicate samples using the primers 341F and 785R (Klindworth et al., 2013). The obtained sequences were analyzed using a combination of the UPARSE v8 (Edgar, 2013) and the QIIME v1.8 (Caporaso et al., 2010b) softwares. Briefly, raw forward and reverse reads for each sample were assembled into paired-end reads considering a minimum overlapping of 50 nucleotides and a maximum of one mismatch within the region using the fastq-join algorithm. The paired reads were then quality filtered, the primer sequences were removed and the individual sample files were merged in a single fasta file. This file was imported in UPARSE where operational taxonomic units (OTUs) of 97% sequence similarity were formed and chimeras were removed using both de novo and reference-based detection. For reference chimera detection, the “Gold” database containing the chimera-checked reference database in the Broad Microbiome Utilities was used. Taxonomy was assigned to the representative sequences of the OTUs in QIIME using UClust (Edgar, 2010) and searching against the latest version of the Greengenes database (McDonald et al., 2012). After the removal of reads affiliated to chloroplast, Archaea and unassigned sequences, a total of 1319653 high-quality merged paired-end reads with an average length of 450 bp were obtained. All the samples analyzed presented Good's coverage values ranging from 90 to 99 capturing sufficient diversity with an adequate sequencing depth (Supplementary Table 1).

The OTU table, composed by 3587 OTUs, and the phylogenetic tree were calculated with FastTree2 (Price et al., 2010) using default parameters and the PyNast-aligned (Caporaso et al., 2010a) representative sequences as an input. The OTU table and the phylogenetic tree were used as inputs for the subsequent analyses of alpha- and beta-diversity. Bray–Curtis distance matrix on the log transformed OTU table was used to perform a Principal Coordinates Analysis, Canonical Analysis of Principal coordinates (CAP) and to conduct a permutational multivariate analyses of variance (PERMANOVA). Statistical analyses were conducted in PRIMER v. 6.1, PERMANOVA++ for PRIMER routines (Anderson et al., 2008) to test differences in bacterial community composition among the three soil fractions and among the three plant species. A first analysis was conducted in a one-way anova to explore difference among the fraction considering as a factor “Fraction” as fixed and orthogonal (three levels: rhizosphere/root surrounding soil/non-vegetated bulk). A second analysis was conducted exploring difference of the microbial assemblage among plants and fractions using the factor “Plant” (three levels: *M. sativa*, *C. nigrescens* and *D. glomerata*) and the factor “Fraction” (two levels: rhizosphere and root surrounding soil) both as a fixed and orthogonal, and their interaction (Fraction × Plant). Prior to perform the statistical analysis we verify that the data were not over-dispersed using PERMDISP for the factor "Fraction" ($F_{1,18} = 3.61, p = 0.21$) and the factor "Plant" ($F_{2,15} = 9.36, p = 0.06$). The shared OTUs among different fractions and plant species have been defined by Venn-diagram analysis using the software available at http://bioinformatics.psb.ugent.be. Diversity indexes were calculated using PAST (Hammer et al., 2001) and their statistical difference was evaluated with the analysis of variance considering the index as response variable and “Fraction” and “Plant” as explanatory categorical variable (see above for details). Raw sequences have been deposited at the ENA European Read Archive under accession number from SAMN07167786 to SAMN07167806, BioProject PRJNA388028, Submission ID SUB2728610.

16S rRNA gene quantitative PCR (qPCR) was performed in a reaction volume of 15 µl with 1 µl of template using universal primers 27F and 1492R and the SsoAdvanced™ Universal SYBR® Green Supermix (Bio-Rad). Thermal protocol was set up as follow: 98°C (3 min), then 40 cycles at 98°C (15 s), 58°C (30 s) and 72°C (1 min). Starting DNA concentration (ng/µl) was measured using a PowerWave HT Microplate Spectrophotometer (BioTek) and the number of 16S rRNA copies obtained with qPCR was normalized with the DNA concentration.

**Bacteria Isolation and Identification**

For bacterial isolation, the R samples obtained from the triplicate plants of the same species were pooled and homogenized. One gram of the resulting soil was suspended in 9 ml of physiological solution (0.9% NaCl), diluted in 10-fold series and plated onto mineral medium (Uhlík et al., 2011), adding biphenyl crystals on the plate lid as unique carbon source. Colonies were randomly picked after 1 week of incubation at 30°C, after the appearance of a stable number of colonies, and were spread three times on the same medium to obtain pure bacterial cultures. A collection of 165 biphenyl-utilizing rhizobacterial strains was established (52 isolates from MS, 56 from DG and 57 from CN) and cryopreserved in 25% glycerol at −80°C. Strain code includes a different number according to the plant of origin (1: MS, 2: CN, 3: DG).

The genomic DNA of each isolate was extracted through boiling cell lysis (Ferjani et al., 2015). Bacterial strains were identified through 16S rRNA gene amplification and partial sequencing (Macrogen, Rep. of South Korea) as described by Mapelli et al. (2013). 16S rRNA nucleotide sequences were subjected to BLAST search (using the blastn suite) and were deposited in the ENA database under accession numbers LT838007–LT838169.

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1https://code.google.com/p/ea-utils/wiki/FastqJoin
2http://microbiomeutil.sourceforge.net/
**bphA Gene Detection, Quantification, and Sequencing in Soil Metagenomes and Strain Genomes**

The presence of the genes encoding for biphenyl dioxygenase α subunit (bphA) was assessed in the metagenome of soil samples (B, S, and R) through PCR as described by Iwai et al. (2010), using the primers BPHD-F3/R1, and further confirmed using 512F and 674R primer set (Leewis et al., 2016b). The latter primer set was also used for a bphA gene qPCR assay as described by Leewis et al. (2016b) and the relative abundance of bphA gene copies was expressed as a ratio over the total community 16S rRNA gene copy number. Bacterial isolates were grown overnight in Tryptic Soy Broth medium and then subjected to CTAB – phenol chloroform DNA extraction (Chouaia et al., 2010). Bacterial isolates were grown overnight in Tryptic Soy Broth medium and then subjected to CTAB – phenol chloroform DNA extraction (Chouaia et al., 2010). BphA gene amplification was performed with 2 µl of DNA template in a final volume of 30 µl with primers 463F/674R (Petric et al., 2011), at the following conditions: buffer 1X, MgCl₂ 1.8 mM, dNTPs 0.2 mM, primers 1 µM, Taq 1.5 U per reaction. PCR thermal protocol was set up as follows: 10 min at 95°C, 30 cycles of 95°C (15 s), 65°C (1 min), 72°C (2 min) and a final elongation step of 10 min at 72°C. All PCR reactions were performed utilizing FastStart™ High Fidelity PCR System (Roche). Genomic DNA of the model PCB-degrading strain *Paraburkholderia xenovorans* LB400 (DSMZ, Germany), was used as positive control for all bphA PCR reactions. PCR results were visualized on 1.2% agarose gel and PCR products that did not show the presence of aspecific bands were sequenced at Eurofins Genomics (Italy). Sequences were then identified using the BLASTn suite of the NCBI website¹ and a Neighbor-Joining phylogenetic tree was then built using MEGA 5.1 (Tamura et al., 2011), computing the evolutionary distances using the Jukes–Cantor method. Nucleotide sequences were deposited in the ENA database under accession numbers LT840193–LT840239.

**In Vivo Assessment of Plant Growth Promotion of Tomato**

Tomato seeds were sown in commercial non-sterile soil placed in trays and, after 1 week, uniform-sized seedlings were selected, each transplanted in separated 0.3 kg commercial soil pots and maintained under greenhouse conditions (≈110 photons m⁻² s of light for 12 h during the day and average temperature of 25°C). A subset of 11 rhizobacteria strains, selected based on a cluster analysis performed using MVSP (Kovach, 1999) combining the PGP activities and the abiotic stress tolerance traits, were inoculated separately on tomato plants. One week after transplantation, the tomato plants (n = 5 for each treatment) were fertilized once with a bacterial suspension of the selected strains at a final concentration of 10⁶ cells g⁻¹ of soil (fresh weight). The inocula have been prepared according to Rolli et al. (2015) resuspending the bacterial cells in sterile tap water. Non-inoculated plants (n = 5) were irrigated with the same amounts of sterile tap water and used as control. After 30 days, plants were harvested for the measurement of shoot/root length and weight. Statistical analysis was performed by a pairwise comparison between each bacterial strain inoculated with the negative control, using Student’s *t*-test.

**RESULTS**

**Diversity and Degradation Potential of Bacterial Communities Associated to Non-vegetated and Root-Associated Soil Fractions in the SIN Caffaro**

The β-diversity of bacterial communities colonizing non-vegetated bulk soil (B) and root-associated soil fractions (rhizosphere, R, and root surrounding soil, S) in the SIN Caffaro was significantly different (Figure 1A, PERMANOVA, F₂,₁₈=3.48, p = 0.0007, Supplementary Table 2A). Considering only the R and S soil fractions associated to the three plant species *M. sativa* (MS), *C. nigrescens* (CN) and *D. glomerata* (DG), the bacterial population assemblage was significantly driven by both plant species and soil fraction (Figure 1B, PERMANOVA, F₁,₁₂ = 4.72, p < 0.001 and PERMANOVA, F₂,₁₂ = 3.21, p < 0.05, respectively, Supplementary Table 2B) but not by their interaction (PERMANOVA, F₂,₁₇ = 1.67, p > 0.05, Supplementary Table 2B). Quantification of the individual factors’ contributions to the observed bacterial community variations, determined by PERMANOVA of Bray–Curtis, showed that the two factors “Fraction” and “Plant species” equally contributed to determine the observed bacterial microbiome variation, explaining 21 and 18% of the variation, respectively (Supplementary Table 3), followed by the interaction of these two factors (11%). For the three fractions, not significant differences have been retrieved considering the OTU richness, but among the rhizosphere the MS and CN plants presented a lower number of OTUs (Figure 1C). OTU evenness was instead significantly different among fractions (ANOVA, F₂,₂₀ = 21.63, p < 0.001) but not between plants (Figure 1D). OTU diversity, calculated as inverse Simpson diversity index, was significantly

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¹www.ncbi.nlm.nih.gov
lower in R compared to B and S (ANOVA, $F_{2,20} = 4.7, p < 0.05$). While not significant differences have been observed among the different plants in S fraction, plant species determined a strong effect in R fraction where DG hosted the significantly highest diversity, followed by CN and MS (Figure 1E).

The three soil fractions collected in the SIN Caffaro showed (i) low number of specific OTUs typical of each fraction, and (ii) a consistent group of shared OTUs (2831/3587) accounting the 97% of the 16S rRNA sequences (Figure 2A and Supplementary Table 4A). The rhizosphere and root surrounding soil associated to the three plant species shared a total of 1922/3436 and 2558/3436 OTUs, respectively, accounting up to 98% of relative abundance in S (Figures 2B,C and Supplementary Tables 4B,C).

The high number of shared OTUs in non-vegetated and plant-associated soils indicated that all the fractions hosted bacterial communities with similar phylogenetic composition and different in the structure, as demonstrated by the analysis of Beta and Alpha-diversity (Figure 1).

The taxonomic affiliation of OTUs (Figure 2D and Supplementary Tables 5, 6) revealed that the soil fractions hosted 37 bacterial phyla, 105 classes (99.3% sequences classified), 153 orders (89% classified), 167 families (67% sequences classified) and 162 genera (23% sequences classified). The most represented phyla were present in all the soil fractions (R, S, and B): Proteobacteria 34% (among these Gammaproteobacteria 16% and Alphaproteobacteria 10%), Actinobacteria 30%, Chloroflexi (8%), and Acidobacteria (8%). The three soil fractions were nevertheless dominated by different phyla/classes. In the non-vegetated soils Proteobacteria (on average 35%, mainly represented by Alphaproteobacteria and Betaproteobacteria) was the prevalent phylum followed by Acidobacteria (22%) and Bacteroidetes (12%) (Figure 2D). Proteobacteria dominated also the rhizospheres with Gammaproteobacteria (26%) and Alphaproteobacteria (9%) as the main Proteobacteria-classes, while in S fraction the Actinobacteria was the dominant phylum (30%).

Moving from the root-associated fractions (R and S) to the non-vegetated bulk soil, a strong reduction of the Actinobacteria phylum was observed in the bacterial communities, from an average of 33% in R to an average of 6% in B (Supplementary Table 5). An opposite trend has been observed for Acidobacteria, Bacteroidetes, Verrucomicrobia, and Gemmatimonadetes phyla that were enriched in the non-vegetated bulk soil and strongly limited in R and S fractions (Supplementary Table 5). Finally, while Firmicutes bacteria were enriched in R (7%) respect to the other two fractions (1%), Chloroflexi were more prevalent in S (10%) respect to R (7%) and B (4%). Difference in taxa enrichment and selection were observed also according to the plant species in both R and S fractions (Supplementary Table 7). The MS and CN bacterial communities in R fractions were dominated by Gammaproteobacteria (52%) and Actinobacteria (50%) respectively, determining a higher dominance index value (0.32 and 0.28) in these communities respect to the DG one (0.15), while DG hosted a more equally distributed bacterial
community with Evenness index of 0.62. An opposite trend has been observed in the S fraction, where DG was dominated by Actinobacteria (37%) while MS and CN presented several taxa equally distributed within the bacterial communities (Evenness 0.72 and 0.69, respectively, Supplementary Table 7).

The \textit{bphA} gene was detected, by means of qualitative PCR using different primer sets, in the metagenomes of all B, S and R samples. Despite that, the quantification of \textit{bphA} gene with primers 512F and 674R was possible only for two replica of the CN rhizosphere (\textit{bphA}/16S rRNA ratio, CN R2: 0.56 ± 0.03; CN R3: 0.50 ± 0.08) and one replica of the DG S (\textit{bphA}/16S rRNA ratio, DG S2: 0.08 ± 0.004), showing the highest \textit{bphA}/16S rRNA gene ratio in the CN rhizosphere.

**Identification of Rhizosphere Biphenyl-Utilizing Bacterial Isolates**

One-hundred sixty-five bacterial strains have been isolated from the rhizosphere of the three plant species MS, CN, DG in mineral medium supplemented by biphenyl as unique carbon source, hence considered as potential biphenyl degraders. The strains belonged only to 3 phyla, \textit{Actinobacteria} (75, 51, and 52% in MS, CN, and DG collections, respectively), \textit{Proteobacteria} (17, 30, and 32% in MS, CN, and DG collections, respectively) and \textit{Bacilli} (8, 11, and 13%, in MS, CN, and DG collections, respectively). All the isolates belonged to phyla that were present in all the metagenome 16S rRNA high-throughput sequencing libraries, even though with a different relative abundance. All the isolates exhibited sequence identity with the closest described species in NCBI database higher than 97% (Supplementary Table 8) and their phylogenetic affiliation at the genus level is represented in Supplementary Figure 1. Among \textit{Gammaproteobacteria}, \textit{Pseudomonas} species were widespread in the three sub-collections, and constituted 15, 23 and 13% of MS, CN, DG rhizosphere, respectively. \textit{Acinetobacter} was isolated only from CN and DG rhizospheres (5 and 16%, respectively). Among the phylum \textit{Bacilli}, the genus \textit{Bacillus} was the most represented, isolated from the rhizosphere of all plant species (7, 13, and 6% in CN, DG, and MS, respectively). Seventy-three percent of all the isolates from the rhizosphere of the three plant species belonged to three genera: \textit{Arthrobacter} (18, 14, 10 strains from DG, MS, CN rhizospheres, respectively), \textit{Microbacterium} (7, 24, 11 strains from DG, MS, CN rhizospheres,
respectively), *Pseudomonas* (7, 8, 13 strains from DG, MS, CN rhizospheres, respectively). MS rhizosphere showed the lowest number (6) of cultivable genera within the biphenyl-utilizing bacteria, widely dominated by *Microbacterium* (44% of the isolates) and *Arthrobacter* (29% of the isolates), compared with CN and DG rhizosphere that included 11 genera.

**Functional Characterization of the Cultivable Rhizosphere Biphenyl-Utilizing Bacteria**

The isolate collection has been screened in vitro for PGP activities, bioremediation potential and abiotic stress resistance (results are detailed for each isolate in Supplementary Table 8). Production of IAA and proteases were the PGP related activities most represented among the collection, present in more than 50% of the strains, in similar percentages among sub-collections isolated from each plant species (Figure 3). A large fraction of the collection produced ammonia (21, 33, and 39% of the MS, CN, and DG strains, respectively), EPS (13, 25, and 14%, respectively) and siderophores (10, 24, and 24%, respectively). ACC deaminase activity was detected in higher percentage among CN (65%) than MS (37%) and DG (36%) sub-collections. Phosphate solubilization activity was present in a small proportion among CN (4%) and DG (11%) isolates, while it was absent in MS sub-collection. Concerning the biodegradation potential, catechol 2,3-dioxygenase activity was widespread in the entire collection with 83, 70, and 63% of the MS, CN, and DG active isolates, respectively. Biphenyl dioxygenase α subunit gene (*bphA*) was detected in the genomic DNA of 65, 81, and 68% of the MS, CN, and DG isolates, respectively. PCR products presenting one single band of the expected size (211 bp) in agarose gel electrophoresis were sequenced, obtaining 45 partial *bphA* gene sequences (9, 19, and 17 from MS, CN, and DG isolates, respectively). Eighty percent of the sequences showed high nucleotide identity (>99%) with *Pseudomonas pseudoalcaligenes* KF707 *bphA*, and the remaining ones showed >99% nucleotide identity with *Rhodococcus w rattisaviensis* strain P13 *bphA1* and *R. opacus* *bphA1* genes (Supplementary Table 9). The *bphA* sequences were clustered in a phylogenetic tree with *bphA* sequences of reference strains having demonstrated PCB degradation ability. The sequences were clustered in two groups, both including known PCB-degraders (Supplementary Figure 2): *Pseudomonas* KF707-like sequences clustered together with *Paraburkholderia xenovorans* LB400, while *Rhodococcus*-like sequences grouped separately.

All strains were able to cope with the osmotic stress induced by the addition of 20% of PEG in the growth medium (Supplementary Table 2). The majority of the strains (71% on average) from each plant species proved to grow in presence of salt in the growth medium, while a percentage comprised between 58 and 71% tolerated high (42°C) and low (4°C) temperatures for growth (Figure 3 and Supplementary Table 8).

A total of 11 strains has been selected basing on PGP activities and abiotic stress tolerance traits (Supplementary Figure 3) and have been tested in vivo on tomato as a model plant. The strains were affiliated to different species belonging to the genera *Pseudomonas, Acinetobacter, Arthrobacter, and Curtobacterium*, and presented an array of different in vitro PGP traits, abiotic stress tolerance and bioremediation potential (Supplementary Table 10).

Despite showing only 1 and 2 in vitro PGP traits, respectively, strains 2–30 and 2–50, both isolated from CN plant and affiliated to different species of the genus *Arthrobacter* (Supplementary Table 10), significantly promoted plant growth compared to the non-inoculated control (Figure 4). Both strains mainly affected shoot development, significantly increasing shoot length (*p* < 0.01) and fresh weight (*p* < 0.001). While 2–30 did not show effect on the root development, 2–50 significantly promote also the root length (*p* < 0.05). Strain 2–50 possesses, moreover, both the tested bioremediation potential traits (Supplementary Table 10).

**DISCUSSION**

**Plant Influence on Soil Bacterial Community Structure and Degradation Potential**

In vegetated soils physico-chemical properties, together with plant root exudation and turnover, shape the structure of the microbial community. Resulting in the so-called “rhizosphere effect,” every plant species presents a specific pattern of root exudation and interacts with the soil community selecting a specific microbiota recruited from the surrounding soil (Haichar et al., 2008; Berg and Smalla, 2009). Here we compare the bacterial communities inhabiting the rhizosphere and root surrounding soils of three spontaneous plant species with a non-vegetated bulk soil from a site heavily contaminated with chlorinated persistent organic pollutants and heavy metals, demonstrating that both the soil fraction and the plant species cooperated in the selection of a specific bacterial assemblage in the root-associated soils. The bacterial communities inhabiting all soil fractions hosted the typical phyla that dominate soil ecosystems, i.e., *Acidobacteria*, *Actinobacteria*, *Bacteroidetes*, *Chloroflexi*, and *Proteobacteria* (Bulgarelli et al., 2013). R and S fractions, that are directly under plant root influence, showed an increase of *Actinobacteria* and *Proteobacteria*. Within the *Actinobacteria*, the order *Gaiellales* was retrieved at higher percentage in all the plant-associated fractions and in particular in the CN rhizosphere. This bacterial order was previously detected in the root system of rice (Hernández et al., 2015) and maize planted on heavy metals polluted soil (Touceda-Gonzalez et al., 2015), however, it is still poorly studied and includes only one described species (Albuquerque et al., 2011). Plant exudates enriched also the orders *Sphingomonadales* and *Pseudomonadales* in the DG and MS rhizospheres, respectively. Members of these orders have already been retrieved in PCB polluted soils, involved in biphenyl degradation and for some genera also in PCB degradation (Leigh et al., 2007; Hu et al., 2015; Jayanna and Gayathri, 2015; Leewis et al., 2016b). The phylum *Chloroflexi*, comprising a well known group of anaerobic PCB degraders (Jugder et al., 2015), was retrieved at relative
abundance higher than 1% exclusively in the S fractions, being particularly enriched in MS1. It is possible that an anoxic microniche occurred in this sample, allowing the selection of Chloroflexi members, in agreement with previous finding of Chloroflexi enrichment in the rhizosphere of M. sativa during a PCB polluted soil phytoremediation trial (Tu et al., 2011) and of Sparganium sp. during biostimulation of the autochthonous microbiota in historically PCB polluted sediments (Di Gregorio et al., 2013).

Studies on a hydrocarbon-contaminated soil proved that the pollutant concentration rather than rhizosphere effect of planted willows had a major role in shaping the bacterial community (Bell et al., 2014; Yergeau et al., 2014). In a previous work (Di Guardo et al., 2017) we suggested that soil pollutant profiles of different former agricultural fields in the SIN Caffaro acted as drivers of the bacterial community composition. Here we demonstrated that in one of the previously analyzed fields, the rhizosphere effect of spontaneous plants adapted to the occurring soil contamination significantly influenced bacterial assemblages in the root-associated soil fraction, R and S. We speculate that in the rhizosphere an enhanced pollutant degradation, according to the higher abundance of taxa previously associated to PCB metabolism, contributed to shape bacterial diversity (Leewis et al., 2016a; Ridl et al., 2016). Our results indicated that autochthonous vegetation, not specifically selected for rhizoremediation efficiency, but rather naturally adapted to counteract phytotoxicity and the specific polluted soil conditions, can establish a strong relationship with the soil bacterial community, potentially sustaining PCB detoxification and in turn the natural attenuation process. The diversity of R bacterial communities was significantly influenced by the three plant species, which selected a root-associated microbiome having a peculiar structure. Despite this, the occurrence of a high percentage of OTUs shared among the different soil fractions, may suggests a strong selection force toward specific taxa constituted by the high pollution level, and possibly related to contaminant detoxification.

Different plant species, including MS, were previously demonstrated to enrich and stimulate the PCB-degrading bacterial communities, inducing an increase of bphA gene copy number and its expression levels in the rhizosphere compared to the non-vegetated bulk soil (Tu et al., 2011; Li et al., 2013; Pagé et al., 2015). In this work, the bphA gene was detected in the metagenome of all the soil fractions and could be quantified in CN rhizosphere and DG soil surrounding roots. The copy number ratio between this gene and the 16S rRNA bacterial gene was higher in R than in S samples, leading to speculate that bacterial populations harboring degrading potential were enriched in the rhizosphere. However, the spread of bphA gene in the bacterial microbiota of the SIN Caffaro soil indicated an intrinsic PCB attenuation potential in this site, regardless of soil fraction or plant species, suggesting a stronger role of edaphic conditions rather than vegetation.

**Isolates Potential for Rhizoremediation**

The root system of plant growing on other PCB polluted sites, was shown to host PCB metabolizing bacteria (Leigh et al., 2006; Ionescu et al., 2009). Hence, we applied a cultivation approach to identify indigenous bacteria in the rhizosphere of plants naturally able to cope with the high level of pollution in the SIN Caffaro, having PGP and PCB degrading potentials. The synergistic effect of these traits has indeed the potential to sustain rhizoremediation approaches, in which both plants and bacteria are involved in remediation (Vergani et al., 2017). The cultivation approach applied to MS, CN, DG rhizospheres, due to the selective conditions determined by biphenyl as unique carbon source, led to isolate only bacteria species belonging to three of the thirty-seven phyla identified in the soil metagenome by 16S rRNA high-throughput sequencing (Actinobacteria, Proteobacteria, and Firmicutes). The high taxonomic similarity
shown by the three sub-collections, in which four species over the 21 detected accounted for the 78% of the whole collection, suggests that the cultivable biphenyl-utilizing bacteria likely were mainly selected by the soil characteristics occurring at the SIN Caffaro site rather than the plant species. The detection of *bphA* gene in 72% of the isolates indicates that these strains harbor the genetic information to initiate the upper pathway of PCB degradation. Moreover, the large majority of the isolates belonged to the genera *Arthrobacter*, *Microbacterium*, and *Pseudomonas*, taxa previously isolated from the rhizosphere.
of plants growing in PCB contaminated soils and known for their capacity to grow on biphenyl and, for some strains, to degrade PCB (Gilbert and Crowley, 1997; Leigh et al., 2007; Uhlik et al., 2011; Kurzawova et al., 2012). Most of the bphA sequences amplified from the rhizobacteria collection showed higher similarity with the biphenyl dioxygenase α subunit of *Pseudomonas pseudoalcaligenes* KF707, a model strain studied for its ability to metabolize PCBs through 2,3-dioxygenation (Furukawa, 1994). This large group of sequences clusters together with the *bphA* sequence of the reference strain *Paraburkholderia xenovorans* LB400. The functional diversity between *Paraburkholderia xenovorans* LB400 and *Pseudomonas pseudoalcaligenes* KF707 is determined by minor differences in the gene sequence (Furukawa and Fujihara, 2008), whose detection was not possible in our analysis due to the short sequence coverage of the used primer set. Functional redundancy for PCB degradation in the SIN Caffaro soil is supported by *bphA* sequences displaying sequence divergence. A second cluster of sequences was affiliated with a gene previously sequenced in *Rhodococcus* spp., including the reference strain *R. jostii* RHA1, which shows a different range of substrates and a low degree of homology compared with the *bphA* proteins produced by *Pseudomonas* KF707 and *Paraburkholderia* LB400 (Masai et al., 1995). The *bphA* sequences of the SIN Caffaro isolates were not related to the phylogenetic identification of the isolates, reflecting that *bph* genes are frequently associated to mobile elements and can be spread through horizontal gene transfer (Pieper and Seeger, 2005). Catechol 2,3-dioxygenase activity was widespread throughout the collection and all the isolated strains harboring *bphA* gene also presented this activity. Biphenyl dioxygenase and catechol 2,3-dioxygenase are enzymes involved in the metabolism of several root-derived and xenobiotic aromatic compounds, so these results support the hypothesis that plants foster organic contaminant degradation by their root-associated bacteria (Fuchs et al., 2011). In particular, since catechol is a toxic metabolite produced by the degradation of biphenyl in the benzoate lower pathway, its degradation capability is essential for a complete mineralization of PCBs that can occur through co-metabolism by bacteria harboring the upper and/or lower degradation pathways (Leevis et al., 2016b).

Plant growth promotion activities have been frequently reported in bacteria from polluted soils (Croes et al., 2013; Thijs et al., 2014; Franchi et al., 2016), according to the ability of plants to select beneficial bacteria when growing under phytotoxic and stress conditions. Likewise, IAA production and ACC deaminase activity are PGP traits well represented in the cultivable biphenyl-utilizing rhizobacteria hosted by all the three plant species. Bacterial IAA production can influence root proliferation and elongation, thereby affecting nutrient and water uptake by plants (Lambrecht et al., 2000) besides phytoextraction and phytostabilization in soils contaminated by heavy metals (Ma et al., 2011). Moreover, indole and its derivatives are considered inter-kingdom signal molecules and play a role as biofilm regulators, a further feature in PGP bacteria-plant interactions (Lee et al., 2015). ACC deaminase activity is known to lower ACC level in plant cells interfering with ethylene biosynthesis and thereby decreasing plant stress response potentially deriving from chemical phytotoxicity (Glick, 2010). Therefore, the isolated strains showed a notable potential in supporting plant adaptation and growth in the highly polluted soil of the SIN Caffaro. Moreover, a significant fraction of isolates tolerated moderate saline, osmotic and temperature stresses. These phenotypes are not directly related to PGP or biodegradation activity, but could confer to the strains a higher fitness in soils with complex and uneven pollutant fingerprints and subjected to seasonal changes (Di Guardo et al., 2017).

Two strains belonging to the genus *Arthrobacter*, one of the most abundant in our collection, also proved to promote the growth of a model plant (tomato) under greenhouse conditions. Since in vitro screening alone is not always sufficient to evaluate the actual PGP potential of bacterial strains (Cardinale et al., 2015), this result is encouraging for further in vivo tests with plant species selected for rhizoremediation trials, considering that these bacteria are also well adapted to the heavy pollution of the SIN Caffaro soil.

**CONCLUSION**

In this work we reported that three spontaneous plant species selected in the strongly PCB-polluted soil of a historical contaminated site, differently affected the composition of the bacterial community in the root-associated soil fractions. The selective pressure imposed by the persistent chlorinated organic pollutants, heavy metals and metalloids in the soil, putatively affected the soil shared microbiome and determined an intrinsic functional potential for natural attenuation in the root-associated soil fractions. Rhizosphere bacterial strains harboring *bphA* gene and displaying catechol dioxygenase and PGP abilities, such as IAA production and ACC deaminase activity, are a potential resource for the improvement of plant growth and the detoxification in the heavily contaminated soils of SIN Caffaro, possibly exploitable for future rhizoremediation interventions.

**AUTHOR CONTRIBUTIONS**

Conceived and designed the experiments: LV, FM, and SB. Selected the sampling site and collected the samples: RM, SA, AD, and SB. Performed the experiments: LV, RM, FM, EC, and MF. Analyzed the data: LV, RM, MF, and FM. Contributed reagents/materials/analysis tools: DD and SB. Wrote the paper: LV, FM, RM, SB, and DD. All authors critically revised the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2017.01385/full#supplementary-material

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

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