Characterization of the belowground microbial community in a poplar-assisted bioremediation strategy for restoring a multi-contaminated soil

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Research

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Abstract

**Background:** Due to their widespread use in industrial applications in recent decades, Polychlorobiphenyls (PCBs) and heavy metals (HMs) are the most common soil contaminants worldwide, posing a risk for both ecosystems and human health.

**Results:** A poplar-assisted bioremediation strategy has been applied for more than four years to a historically contaminated area (PCBs and HMs) in Southern Italy using the Monviso poplar clone. This clone was effective in promoting a decrease in all contaminants and an increase in soil quality in terms of organic carbon and microbial abundance. Moreover, a significant shift in the structure and functioning of the belowground microbial community was also observed when analysing both DNA and cDNA sequencing. In fact, an increase in bacterial genera belonging to *Proteobacteria* and able to degrade PCBs and resist HMs was observed. Moreover, the functional profiling of the microbial community predicted by PICRUSt2 made it possible to identify several genes associated with PCB transformation (e.g. *bphAa*, *bphAb*, *bphB*, *bphC*), response to HM oxidative stress (e.g. catalase, superoxide reductase, peroxidase) and HM uptake and expulsion (e.g. ABC transporters).

**Conclusions:** This work demonstrated the effectiveness of the poplar clone used in stimulating the natural belowground microbial community to remove contaminants (phyto-assisted bioremediation) and improve the overall soil quality. It is a practical example of a nature-based solution involving synergic interactions between plants and the belowground microbial community.

**Background**

Polychlorinated biphenyls (PCBs) are highly stable, hydrophobic and persistent organic pollutants, which have been used for a wide range of industrial purposes, but also as pesticide additives. Due to their recalcitrance to degradation and their lipophilic nature, they bioaccumulate and biomagnify through the food chain and have a broad range of toxic effects on humans, including as probable carcinogens and endocrine disruptors [1,2,3]. Although their production has been banned since the 1970s, their extensive use and high persistence have caused widespread environmental contamination. Consequently, their removal from contaminated ecosystems remains a challenge.

Heavy metals (HMs) are “trace elements” in pristine ecosystems, whereas their concentrations increase when there are anthropogenic activities (steel industry, waste disposal, agriculture, mining). HMs pose a risk owing to their toxicity, bioaccumulation and biomagnification [4]; they cause cell oxidative stress through formation of free radicals and are able to replace essential metals in pigments or enzymes, with consequent disruption of their functions [5,6]. PCBs and HMs are frequently found as soil co-contaminants [7,8], posing a risk for both ecosystems and human health. Widespread and chronic environmental contamination by PCBs and HMs requires remediation solutions in line with environmental sustainability. The European Commission recently proposed a new priority and environmental friendly approach termed “Nature-Based Solutions, NbS” [9], relying on increasing and exploiting ecosystem...
homeostatic natural capacities to recover from environmental impacts and providing at the same time social and economic benefits. In this context, natural microbial communities have, thanks to their wide metabolic versatility and adaptation capacity, a key role in ecosystem regulation services, ensuring geochemical cycles and removing chemicals from contaminated environments [10]. The presence of an abundant and diverse microbial community with the ability to remove contaminants is a necessary prerequisite for an immediate and effective response to the various contaminants continuously affecting ecosystems [11]. The changes that can be observed in a microbial community can be analyzed from a global and multivariate perspective, to understand and assess the impact of a complex chemical mixture on both sensitive microbial populations and resistant and/or resilient ones [12]. Although the metabolic potential of microbial communities is unlimited, some recalcitrant compounds such as PCBs require various abiotic conditions (e.g. both aerobic and anaerobic conditions, and electron donors) for biodegradation, and these are difficult to achieve in one and the same environment [13]. To improve natural biodegradation of chemicals, Plant-Assisted Bioremediation (PABR) has been proposed in recent years for the restoration of contaminated soils [14,15,16]. PABR is an in situ treatment where plants are used for stimulating the bacterial degradation of persistent organic contaminants such as polychlorinated biphenyls (PCBs) and phyto-stabilization of inorganic ones (e.g. heavy metals), [8,17].

The complex and synergistic actions established in the rhizosphere between the tree root system and natural belowground microbiota make it possible to remove, convert or contain toxic substances in soils [18,19,20]. Our recent research has shown the ability of the hybrid poplar genotype Monviso (Populus generosa x Populus nigra) to grow in a nutrient poor soil, resisting high metal concentrations and promoting PCB biodegradation in the rhizosphere in a chronically polluted area [8,21]. The Monviso clone was effective in promoting both an overall contaminant decrease (PCB degradation and HM phyto-stabilization) and an increase in soil microbial activity, a little more than one year after the tree planting. This bioremediation strategy has continued until now in the same area. In this regard, the results of subsequent chemical (PCBs and HMs) and microbiological analyses at 55 months of bulk and rhizosphere soil samples are reported in the present study. In particular, the microbial community was analysed in detail by determining the total microbial abundance, the cell viability and the composition of the bacterial groups using NGS. Both environmental DNA and RNA were used as templates for sequencing the hypervariable V4-V5 regions of 16S rRNA using MiSeq Illumina. The microbiological analyses were performed in the planted plots (Rhizosphere and Bulk Soil samples) and at control points (un-planted soil). The shifts in the structure and function (functional profiling predicted by PICRUSt2) of the belowground microbial community of the planted plots (poplar Rhizosphere and Bulk Soil immediately close to it) are reported.

**Results**

**Polychlorinated Biphenyls**

The concentrations of six PCB markers (28, 52, 101, 153, 138 and 180) and two dioxin-like congeners (105, 118) are reported in Table S1 (Additional File 1) for unplanted (Control) and planted plots before the poplar plantation (t=0) and at 55 months. In particular, for the planted plots, the analyses were performed
in the Bulk Soil and in the Rhizosphere. Only residual concentrations of PCBs, below the national legal limit of 60 ng/g soil (Italian Decree 152/06), were found at 55 months in the planted plots (both in the Bulk Soil and the Rhizosphere samples). On the other hand, in the Control soil, outside the planted area, a high PCB concentration (~1400 ng/g soil) still persisted (Table S1; Figure S1, Additional File 1).

**Heavy Metals**

Metal concentrations in soil samples from the planted plots (Rhizosphere and Bulk Soil) decreased if compared with the same soil before the planting (Control t=0); all HMs were below the Italian legal limits, excepted for cobalt (Co) in the Bulk Soil (Table S2, Additional File 1). On the contrary, clear contamination still persisted in the un-planted soil (Control t=55 months), where the highest metal concentration was found for Zn (549.73 ± 242.59 mg/kg, Table S2; Figure S2, Additional File 1).

**Microbiological abundance, Organic carbon and Nitrogen contents**

The microbial abundance (N. cells/g soil) and cell viability (% live cells/live + dead) were significantly (test t, p < 0.01) higher in the planted plots (Rhizosphere and Bulk Soil) than in the Control. The highest values were found in the Rhizosphere (Figure 1A and 1B). Similarly, cell viability was higher (t test, p < 0.01) in the Rhizosphere and Bulk Soil than in the Control (Figure 1B). In line with these results, the percentages of organic carbon were higher in the Rhizosphere (1.9%±0.11) and Bulk Soils (2.0±0.10) than in the Control (1.5±0.1).

Finally, the highest total nitrogen value (0.16%±0.001) was detected in the Rhizosphere, while lower values were found in the Bulk (0.12±0.001) and Control soils (0.09±0.001).

**DNA sequencing results**

A total of 5,745,690 raw reads were obtained from the DNA (V4-V5 regions) deep-sequencing. After a trim and quality edit, 2,485,475 reads were retained: 21% of the total reads belonged to the Control, 41% to the Bulk Soil and 38% to the Rhizosphere. 99.9% of the total OTUs were classified at the Phylum level, 78.3% at the Class level, 66.6% at the Order level, 57.7% at the Family level and 39.0% at the Genus level, respectively.

*Proteobacteria* were the dominant Phylum both in the planted plots (62% in the Rhizosphere and 55% in Bulk Soil, respectively) and in the Control (32%); the latter percentage was significantly (t test, p <0.01) lower if compared to the Rhizosphere and Bulk Soil. *Acidobacteria, Firmicutes* and *Actinobacteria* were the second most abundant group in the Rhizosphere (14.39%), Bulk Soil (13.70%) and Control (26%), respectively (Figure 2A). In the latter, percentages of 5% of *Chloroflexi* and 6% of *Thaumarchaeota* (Archaea) were also present.

At the class level, the *Gammaproteobacteria* group was the most abundant one, both in the Rhizosphere and Bulk Soil (36% and 38% respectively), while in the Control its percentage was significantly lower (8%), (p < 0.001). *Alphaproteobacteria* were present with a percentage of 18% in the Rhizosphere, and 13.5%
and 15% respectively in the Bulk and Control soils. *Betaproteobacteria* were 8% in Rhizosphere, and 4% and 5% respectively in the Bulk and Control soils. *Acidobacteria* were 12% in the Rhizosphere, 9% in the Bulk soil and significantly lower in the Control (4%, p <0.05). Finally, the percentage of *Actinobacteria* was higher in the Control (26%) than in the other conditions (6% in both Rhizosphere and Bulk Soil; t test, p<0.0001), (Figure 2B).

The most abundant genera found are reported in Figure 2C. A higher number of genera, most of them belonging to *Proteobacteria* (*Gammaproteobacteria*: *Aeromonas, Pseudomonas, Stenotrophomonas, Steroidobacter, Alphaproteobacteria*: *Mesorhizobium, Rhizobium*), were identified in the planted plots (Rhizosphere and Bulk Soil); however higher percentages of Gp6 (*Acidobacteria*) and of *Nitrososphaera* (*Nitrososphaerales - Archaea*) were also found.

Some genera, such as *Nocardiodes* (*Actinobacteria*), *Skermanella* (*Alphaproteobacteria*) and *Steroidobacter* (*Gammaproteobacteria*), were found in higher percentages in the Control than in the planted plots. Finally, several genera were found in percentages lower than 1%. Interestingly, significant differences were found between the Control and planted soils, not least for the less abundant genera.

The community diversity of the planted plots (Rhizosphere and Bulk Soil) and Control soil were evaluated considering both the alpha (Table 1 for DNA and Table 2 for cDNA) and beta diversity (Figure S3A and S3B, Additional File 1).

The Kruskal-Wallis test for the Chao1 index (Table S3, Additional File 1) showed significant differences between Control and Bulk Soil communities (p <0.05); no differences were found between Rhizosphere and Bulk Soil and Rhizosphere and Control. The test performed for the Shannon Index found significant differences between Control and planted plots (Rhizosphere and Bulk Soil), (p < 0.02 both the samples). Finally, the Evenness index showed Control with a more homogenous community than Rhizosphere and Bulk Soil (p < 0.02).

**Table 1** Average values of reads obtained from DNA and values of Alpha diversity expressed as Chao1, Shannon and Evenness indices in planted plots (Rhizosphere and Bulk Soil) and Control; standard errors are in italics.

|        | DNA  | Reads Filtered | Chao1 | Shannon Index | Evenness |
|--------|------|----------------|-------|---------------|----------|
| Rhizosphere | 15,4645 ± 16.974 | 2,180 ± 183.79 | 9.68 ± 0.22 | 0.83 ± 0.01 |
| Bulk Soil    | 173,528 ± 12.710  | 2,063 ± 102.70 | 9.10 ± 0.16  | 0.88 ± 0.01 |
| Control      | 129,109 ± 6.442   | 2,855 ± 88.65  | 10.70 ± 0.08 | 0.93 ± 0.00 |

The Bray-Curtis PCoA confirmed significant differences in the community structure under the different conditions: Control was significantly dissimilar from both Bulk Soil and Rhizosphere (PERMANOVA, p < 0.001). Moreover, a significant dissimilarity (Bray-Curtis PCoA, PERMANOVA, p < 0.002) was also found inside each plot between the Rhizosphere and Bulk Soil samples (Figure S3A, Additional File 1).
cDNA sequencing results

The microbial community was analysed through amplicon sequencing of the 16S rRNA gene and the 16S rRNA transcripts to directly compare the total and active microbial communities [22,23,24] and to better identify the microorganisms potentially involved in the soil decontamination processes. In total 3,995,841 raw reads were obtained from cDNA sequencing and after a trim and quality edit, 1,801,772 were retained: 48% of them derived from the Bulk Soil and 52% the Rhizosphere, respectively. The average trimmed read values for each Rhizosphere and Bulk Soil are reported in Table 2.

The Kruskal-Wallis test on the alpha diversities indices showed that there were no significant differences for the Chao1 index, whereas species diversity and evenness were significantly higher in the Rhizosphere than Bulk Soil (respectively p <0.02 and p <0.01, Table S3).

| cDNA | Filtered Reads | Chao1 | Shannon Index | Evenness |
|------|----------------|-------|---------------|----------|
| Rhizosphere | 149,644 ± 15,393.98 | 2,045 ± 198.33 | 9.31 ± 0.19 | 0.85 ± 0.01 |
| Bulk Soil | 150,652 ± 15,641.73 | 1,334 ± 250.89 | 6.94 ± 0.77 | 0.67 ± 0.06 |

The PCoA analysis confirmed significant differences between the overall OTU data from DNA and cDNA and between the Rhizosphere and Bulk Soil cDNA (PERMANOVA, p<0.001 Figure S3B, Additional File 1).

In cDNA the dominance of Proteobacteria was even more evident (75% of the overall Prokaryotic community in Rhizosphere and 78% in Bulk Soil) than in DNA and quite low percentages of other phyla (such as Acidobacteria, Actinobacteria, Firmicutes and Planctomycetes) were found, in both the Rhizosphere and Bulk Soil samples (Figure 3, A-B-C). The genera identified in the Rhizosphere (119) and in the Bulk Soil (118) are reported in Additional File 2. Several genera were present in different percentages when comparing the Rhizosphere and Bulk Soil (e.g. Pseudomonas 14% Rhizosphere and 12% Bulk Soil, Stenotrophomonas 11% Rhizosphere and 13% Bulk Soil, GP6 6% Rhizosphere and 8% Bulk Soil and Aeromonas 5% Rhizosphere and less than 1% Bulk Soil).

Functional profiling of the microbial community

The functions of the bacterial communities in the samples were predicted by PICRUSt2, based on KEGG pathways using both DNA and cDNA sequences.

A total of 7,718 KOs and 440 pathways were identified (Additional File 3 and Additional File 4). The PCA of the predicted functional profiling of the soil microbial community based on the DNA samples showed notable differences in the basic cellular processes between the various conditions (Rhizosphere, Bulk Soil and Control, Figure S4A, Additional File 1). This result demonstrated that the poplars influenced not only the structure (i.e. bacterial diversity), but also the functioning of the soil bacterial community.
In any case, various genes related to PCB degradation, HM and oxidative stress, membrane transporters, germination and sporulation were identified in both the planted plots and Controls (Figure 4). For example, four gene families encoding enzymes involved in biphenyl degradation were found: \textit{bphAa}, \textit{bphAb} (biphenyl 2,3-dioxigenase), \textit{bphB} (cis-2,3-dihydro-2,3-dihydroxybiphenyl dehydrogenase) and \textit{bphC} (2,3-dihydroxybiphenyl 1,2-dioxigenase), (K08689, K08690, K015750, K018087, respectively). However, \textit{bphB} was found with the highest values in the planted plots and particularly in the Bulk Soil.

Moreover, genes potentially involved in aerobic PCB degradation with the meta cleavage pathway of aromatic compounds (i.e. degradation to pyruvate and acetyl-Coa), were also identified and were most abundant in the planted plots (Bulk Soil and Rhizosphere).

As regards heavy metals, several genes associated with their presence were also identified. In fact, functional genes associated with the oxidative stress caused by HMs, such as catalase (CAT) superoxide reductase (SOR), peroxidase, glutathione reductase (GR) and thioredoxin, were identified with a higher relative abundance in the planted plots than in the Controls. Similarly, the highest relative abundance of germination and sporulation genes was found in the Bulk Soil. Finally, ABC transporters, which make cellular HM uptake and expulsion possible, were higher in the planted plots than in the Controls.

Interestingly, the PCA performed on cDNA (Figure S4B, Additional File 1) showed that the active microorganisms inside the planted plots were metabolically similar. Significant differences between the Rhizosphere and Bulk Soil were still found in the meta cleavage pathways of aromatic compounds and in \textit{bphB}, which were more abundant in the Bulk Soil than in Rhizosphere (t test p<0.05) (Figure 5A).

**Discussion**

Due to their low bioavailability and complex pathways, the degradation of PCBs in the soil is normally limited [25]. For this reason, the poplar-assisted bioremediation (PABR) strategy was applied in this contaminated area. This paper, to our knowledge, is one of the first to report the characterization of the belowground microbial community in a soil decontaminated through poplar-assisted bioremediation. The overall data showed an overall improvement in soil quality not only in terms of decontamination and increase in nutrients, but also in the structure and functions of the microbial community. The comparison of the chemical (PCBs and HMs) and microbiological analyses between control and planted soils showed the effectiveness of the PABR in recovering the contaminated area, considering both the soil immediately surrounding the roots (Rhizosphere) and the Bulk Soil at 50 cm from the trunk and at 40 cm depth. These results confirm the overall decontamination found at 14 months, although some heavy metals (i.e. V, Sn and Pb) were still found to exceed their legislation threshold values [8]. At 55 months, the roots grown inside the planted plots increased the volume of the rhizosphere habitat and positively influenced the entire belowground microbial community. In fact, at 55 months the poplar trees reached 6 meters in height and the belowground root biomass had been able to spread fourfold compared to the previous 14-month sampling, reaching a volume of c.a. 16.000 cm$^3$. This means that the planting layout of the poplars (0.5 × 2m) was effective in ensuring that their root development positively stimulated the overall microbial community in each planted plot. The positive “rhizosphere effect” has been reported by other
authors [26,27] and was presumably due to the root surface releasing several organic compounds into the adjacent soil. Roots release low-molecular-weight exudates (e.g. sugars, polysaccharides, amino acids, aromatic acids, aliphatic acids, fatty acids, sterols, phenolic compounds, enzymes, proteins, plant growth regulators and secondary metabolites) which attract soil microorganisms into the rhizosphere where they multiply by several orders of magnitude [28]. In turn, microorganisms strongly influence plant development and health by mineralizing organic matter, degrading organic pollutants, including persistent ones such as PCBs, improving heavy metal bioavailability by varying the degree of reaction of the soil (pH), releasing chelating substances (organic acids, siderophores) and promoting oxidation/reduction reactions [8,20]. In line with the above, the DNA sequencing data showed higher percentages of bacteria genera (i.e. *Stenotrophomonas, Pseudomonas, Mesorhizobium, Rhizobium, Skermanella* and *Gp6*), involved in biogeochemical cycles (in line with an increase in soil C and N) and xenobiotic degradation (Figure 3C), in the Rhizosphere and Bulk Soil than in the Control [29,30,31,32,33].

On the other hand, the Control microbial community was quite different from that of the planted plots and some genera such as *Nitrososphaera* (*Archaea*), *Nocardioides* (*Actinobacteria*) and *Steroidobacter* (*Gammaproteobacteria*) were significantly more abundant than in the other conditions.

The fact that the plant presence promoted a *Proteobacteria* dominance in all plots was demonstrated by both DNA (average value 58%) and even more by cDNA (average value 77%) results. The *Proteobacteria* increase was in line with the rise in microbial abundance and viability, showing their active role in removing chemicals and increasing soil quality. The comparison of DNA and cDNA showed that in fact different bacterial genera or percentages of each taxonomic group were active in the soil studied, highlighting not only the role of *Proteobacteria*, but also some differences in the percentages of some classes between the bulk soil and rhizosphere. The *Proteobacteria* group includes most bacterial species involved in the main biogeochemical cycles and is typically the most abundant phylum found in a good quality state soil [34] and several *Alpha-, Beta-,* and *Gamma-proteobacteria* species have been reported to be able to degrade PCBs and increase with a plant presence [13,35]. Interestingly, in cDNA a higher percentage of *Firmicutes* was also found and in this group there are several bacteria able to resist and remove HMs [36,37,38].

The DNA predictive functional analysis confirmed overall significant differences between the planted plots and Control (Figure S4A, Additional File 1). That it is to say that the poplars were able to change both the structure and functioning of the belowground microbial community. However, the predictive functional analysis showed that some genes involved in biphenyl degradation were present inside the control microbial community, demonstrating that the latter was potentially able to degrade PCBs and that the plant stimulated the specific microbial populations able to recover the soil. Interestingly, the cDNA predictive functional analysis showed that the overall bacterial populations inside each planted plot (Rhizosphere and Bulk Soil) were metabolically similar, showing how the plant positively influenced the functioning of the overall microbial community. In fact, the number of bacteria associated with some specific genes (e.g. *bphA, bphB* and *bphC*) significantly increased in the planted plots (Rhizosphere and Bulk Soil), demonstrating the effectiveness of the poplar in promoting microbial transformations. *bphA* is a multi-component enzyme consisting of terminal dioxygenase and electron transfer components, which
starts the degradation process of the biphenyl ring; \textit{bphB} catalyzes the conversion of dihydrodiol to a dihydroxy compound and \textit{bphC} is involved in ring meta-cleavage [39]. Chlorobenzoate compounds are one of the final products of aerobic PCB degradation and can be degraded through the meta-cleavage pathway of aromatic compounds to pyruvate and acetyl-CoA [40]. In line with these results, numerous species of \textit{Pseudomonas}, the dominant genus found in the planted plots, have these genes [41,42,43,44,45]. The fact that both \textit{bphB} and the meta-cleavage pathway of aromatic compounds had higher values in the Bulk Soil than in the Rhizosphere, shows that some degradation steps were presumably more favoured in proximity of the rhizosphere.

The presence of heavy metals is known to cause bacterial physiological stress with the production of reactive oxygen species (ROS), [46]. In this work, we found an increase in the planted plots in bacteria able to resist HM stress by producing these protective enzymes. In fact, many bacteria are able to reduce oxidative stress by producing catalase (CAT), superoxide reductase (SOD), peroxidase, glutathione reductase (GR) thioredoxin (Trx) and thioredoxin reductase (TrxRs) [47]. CAT and SOD are able to convert ROS into oxygen and water and make the maintaining of cellular integrity possible [48,49,50]. SOD, a metalloenzyme which converts highly toxic superoxide into oxygen and less toxic hydrogen peroxide, is similar [47] (Banerjee et al., 2015).

Similarly, we found an increase in the planted plots in bacteria able to respond functionally to toxic molecules using ABC transporters. Prokaryotic ABC transporters have a key role in the import of nutrients and export of unwanted molecules such as toxic ones [37]. The overall functional profiling of the microbial community showed how the plant presence favoured an increase in bacterial genera able at the same time to transform and degrade all contaminants and to respond functionally in order to maintain their homeostasis [37].

Conclusions

This work has investigated the shifts in the natural microbial community in a soil where PCBs and HMs were removed by PABR. The overall results showed how the synergic relationships established between plants and belowground microorganisms were able to increase soil quality, by enriching the soil with active bacteria able to degrade and contain pollutants, resist stresses and increase soil nutrient content.

At the same time the poplar trees were able to grow healthily and transform a degraded area into one that could be used advantageously. Plants and their microbiota can be considered “metaorganisms”, i.e. able to overcome biotic and abiotic stress. The PABR strategy is recommended as an effective and good example of a nature-based solution, as mentioned and recommended by the European Commission.

Material And Methods

Study area and sampling
The area (≈4.5 Ha) near the city of Taranto in Puglia (Italy) had been contaminated for several decades by wastes and pollutants (mainly heavy metals and polychlorinated biphenyls) because it was used as an illegal city dump. In particular polychlorinated biphenyls (PCBs) accumulated over time due to spilling and improper disposal of dielectric fluids from a power station with transformers nearby. A preliminary characterization of the experimental site showed PCB hot spots and widespread HM contamination with values higher than the legal limits, as described in detail in Ancona et al. [8]. In April 2013 about 650 poplar cuttings were planted in a site sub-area of 785 m², in 8 rows 2 m apart. Inside each row the cuttings were placed at a distance of 0.5 m from each other. A first sampling campaign performed at 14 months from their planting showed a general decrease in PCBs and HMs thanks to the poplar-assisted bioremediation strategy applied [8]. In the present study we report the results of a further sampling performed at month 55 after planting. As in the previous sampling, three contaminated plots were sampled. They were selected for the highest initial values of PCB contamination (i.e. average values of about 240 ng/g), which exceeded the national legislative limit (Italian Decree152/06) of 60 ng/g for green areas [8].

The area of each plot was 1 m², with the plant at its centre. Each plot polluted by PCBs and HMs was sampled considering two different distances from the tree: four aliquots of soils below the trunk (at 0 cm and at 5-30 cm depth) and around the tree roots (Rhizosphere samples) and four aliquots at 50 cm distance from the trunk and at 20-40 cm depth (Bulk Soil samples) were collected. The composite soil samples were obtained from the latter, considering four cardinal points with each tree at its center, as described in detail in [8]. Moreover, composite soil samples (Controls) were collected outside the planted area (ca 20 m distant).

From each plot, two composite sub-samples (2 g each) were immediately stored in a Life-Guard Soil Preservation Solution (Qiagen, Manchester, UK) in order to halt any possible degradation of the microbial RNA and consequently preserve the gene expression profiles and microbial community structure composition of any active cells. In a similar way, other two soil sub-samples (1 g each) were immediately fixed in a phosphate buffer saline solution with formaldehyde 2% for DAPI counts. Finally, two fresh soil sub-samples (1g each) were used for evaluating the microbial activity.

Two composite samples (1000 g each) were also collected for chemical analysis and stored at room condition. Each datum from each different condition (Rhizosphere, Bulk Soil and Control) is reported as an average value of the 3 replicate plots.

**PCB Chemical analysis**

PCBs were extracted from soil samples (Rhizosphere, Bulk and Control) with an Accelerated Solvent Extractor (ASE 300 DIONEX, USA). About 500 mg of powdered soil were mixed with previously washed diatomaceous earth to form a free-flowing powder. The mixture was extracted in an ASE extraction cell and purified using Silica Gel activated with concentrated sulphuric acid (3:2) in accordance with 210 DIONEX Technical note. The extraction solvent used was n-hexane operating at 100 °C and 1500 psi. The
organic extracts were then evaporated to incipient dryness under a gentle nitrogen stream (using a Caliper Life Sciences TurboVap II Concentration Workstation) and re-solubilized into 0.5 mL Nonane. GC-MS analysis of 31 PCB congeners (Table S4, Additional File 5) was performed using a ThermoElectron TRACE GC Ultra coupled with a PolarisQ Ion Trap (Thermo Electron, Austin, TX) mass spectrometer equipped with a PTV injector and a TriPLUS RSH autosampler. Details for GC-MS analyses are described in Additional File 5. PCB quantification analysis showed that only 8 congeners - 6 PCB markers and two dioxin-like ones (105, 118) - were detected in the soil samples collected at different times of investigation (t=0 and t=55 months).

**Heavy Metals analysis**

Soil samples were acid digested in closed PTFE vessel devices using temperature control microwave heating (Ethos Touch Control, Milestone, Microwave Laboratory Systems). Soil mineralization was performed using Aqua Regia extraction by treating 500 mg of powdered samples with 9 mL of concentrated HCl and 3 mL of HNO$_3$ [51]. Heating was achieved in a two-step procedure: 10 min to reach 200 °C followed by 15 min at 200 °C. Resulting solutions were diluted with ultrapure water in order to obtain a maximum content of 5% of acids and of 0.2% dissolved solids.

The quantification of 14 different mineral elements (Be, V, Cr, Co, Ni, Cu, Zn, As, Se, Cd, Sn, Sb, Tl, Pb) was performed by mass spectrometry with an inductively coupled plasma source (ICP-MS) equipped with a 7700x Agilent (Agilent Technologies, Japan).

**Organic Carbon and total Nitrogen analysis**

Total and organic carbon and total nitrogen were determined with an elemental analyser (Carlo Erba NA 1500 series 2 C/H/N/O/S) equipped with an autosampler. For the organic carbon analyses, the samples were acidified with 20 μl 5 M ultrapure HCl and kept at 50–60 °C for 30 min in order to remove inorganic carbon. The method used is based on the complete and instantaneous oxidation of the solid sample by combustion in an O$_2$ enriched environment. Each sample, contained in tin or silver capsules for the determination of total and organic carbon, respectively, was introduced into the combustion reactor and maintained at about 1000°C for 20 minutes. Helium was used as a carrier gas to transport the combustion gases (CO$_2$, N$_2$, H$_2$O) through a catalyst (Al$_2$O$_3$-WO$_3$) in order to inhibit the formation of nitrogen.

The combustion mixture, together with the excess of oxygen, passed through a reduction column, consisting of granules of pure reduced copper, maintained at about 650 °C, where the excess oxygen was adsorbed and any nitrogen oxides present were transformed into elemental nitrogen. The resulting gases (CO$_2$, N$_2$, H$_2$O) then passed through an absorbent filter (magnesium perchlorate), which retains the water, to a gas chromatographic column that separates N$_2$ from CO$_2$. The column connected to a thermal conductivity detector (TCD) provided an output signal proportional to the quantity of the individual
components in the mixture. From the measurement of the chromatographic peak area, corrected for white, the amount of carbon and nitrogen was obtained by comparison with a calibration curve.

**Total Microbial abundance and Cell viability**

Microbial abundance and cell viability were analysed using epifluorescence microscope-based methods which do not need DNA extraction from soil. The microbial abundance (No. cells/g dry soil) was assessed performing total direct counts. This method makes it possible to detect all the microbial cells present in a soil sample regardless of their physiological state and metabolic activity, thanks to the DAPI dye (4',6-diamidino-2-phenylindole), which is a DNA fluorescent intercalant. Fixed soil subsamples (1 g each) were processed as reported in detail in Barra Caracciolo et al. [52].

The cell viability (% live cells/live + dead) was evaluated in fresh soil subsamples (1 g each). The two dyes used to measure the ratio of live to dead cells were propidium iodide and SYBR Green II (Sigma-Aldrich, Germany) respectively, as described in detail in previous works [53,54].

**DNA and RNA extraction and sequencing**

Extraction of environmental RNA and DNA (eDNA) from each soil sample (Rhizo, Bulk and Control) was carried out using the RNA Power Soil Total RNA Isolation Kit and the DNA Elution Accessory kit (Qiagen), following the manufacturer's recommendations. In order to evaluate the putative active bacteria in Rhizosphere and Bulk Soil, the same soil samples were used for RNA extraction; the latter was extracted as reported above and ds-cDNA synthesized in accordance with [24]. Both eDNA and ds-cDNA were used as templates for MiSeq Illumina sequencing of the hypervariable V4-V5 of the 16S rRNA gene, using the U519F and U926R primers [55].

The amplification with MiSeq Illumina was performed by Bio-Fab Research s.r.l. (Rome, Italy).

**Data Processing**

A total of 10 Mb reads were generated and were analysed using SEED2, a user interface-based sequence editor and NGS data analysis pipeline [56], and Mothur [57] software packages. The forward and reverse reads were joined using SEED2 software with a 15% of maximal difference and minimal overlap of 40bps. The primers, sequences with an ambiguous base and average quality score lower than 30 were removed. Before performing the chimera cleaning, with USEARCH (http://www.drive5.com/usearch) from the SEED2 software package, we de-replicated the sequences. After the chimera cleaning the sequences were re-replicated. To avoid bias, the number of sequences in each sample was rarefied to the lowest one.

The resulting filtered and trimmed sequences were clustered into Operational Taxonomy Units (OTUs) using UPARSE [58] from the SEED2 package and applying the average neighbour algorithm with a similarity cut off of 97%. The OTUs with less than 0.005% of high-quality reads were then filtered to avoid an overestimation of the diversity [59,60].
With the Mothur software, the centroid sequence of each cluster was selected as the most representative of each OTU and was taxonomically classified at the genus level according to Ribosomal Database Project (RDP, http://rdp.cme.msu.edu) trainset 16 with a confidence threshold of 80%.

**Predictive functional analysis**

The PICRUSt2 software tool (https://github.com/picrust/picrust2) was used for predicting functional abundances based on the 16S rRNA gene amplicon data sets [61]; the OTUs table generated by UPARSE was used as an input. The prediction of KO relative abundances was performed with hidden-state prediction [62] and was used to infer pathways abundances [63]. The statistical analysis was carried out with STAMP (version 2.1.3, https://pypi.org/project/STAMP) in order to evaluate the significant differences between metagenome metabolic profiles under the different conditions.

**Statistical analysis**

An Alpha diversity table was created using SEED2 software with several diversity indices (Species richness, Shannon and Evenness indices). Principal Coordinate Analysis (PCoA) was performed with the online tool Microbiome Analyst (http://www.microbiomeanalyst.ca) using a Bray-Curtis index dissimilarity method and PERMANOVA as the statistical method in order to analyse the Beta diversity. The t-tests were performed to determine if there was a significant difference between the means of two groups (between Rhizosphere and Bulk Soil and DNA and cDNA). All the histograms and the PCoA graphs were made using MS Excel.

**List Of Abbreviations**

CAT: Catalase; GSH TS: glutathione transport system; GSHRs glutathione reductase; HM: heavy metal; NGS: Next-Generation Sequencing; POD peroxidase; SOD: superoxide dismutase; Trx: thioredoxin; TrxRs: thioredoxin reductase.

**Declarations**

**Ethics approval and consent to participate**

Not applicable

**Consent for publication**

Not applicable

**Availability of data and material**

The datasets generated during the current study are available in the ENA repository: https://www.ebi.ac.uk/ena/browser/view/PRJEB37712
Competing interests

The authors declare that they have no competing interests

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Authors' contributions

ABC and VA designed this study. VA, GA, CC collected the soil samples from the study area. GLG, PG, LR performed microbiological analyses. VA, GA, CC performed chemical analysis. MFL, AFG, PV, GLG performed the bioinformatics and data analysis of raw data. ABC, VA, PG, GLG wrote the manuscript. All authors read and approved the final manuscript.

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Additional Files

Additional File 1.docx
PCB and heavy metals concentrations tables and figures; Alpha diversity statistical analysis; Bray-Curtis PCoA graph of DNA and cDNA samples and PCA of functional genes profiling of DNA and cDNA samples.

Additional File 2.xls

Number of genera found in all samples considering DNA and cDNA.

Additional File 3.xls

List of KEGG Orthologs (KO) and their abundances in all samples considering DNA and cDNA.

Additional File 4.xls

List of pathways and their abundances in all samples considering DNA and cDNA.

Additional File 5.docx

PCB extraction methods and instrumental conditions for GC-MS analyses.

**Figures**
Figure 1

A

N. cells/g soil

OC: 1.5%

OC: 1.9%

OC: 2.0%

Control

Rhizo

Bulk

B

% live cells/(lives+deads)

Control

Rhizo

Bulk
Microbial abundance (No. cells/g soil); B. Microbial viability (% live cells/live + dead) OC: organic carbon
Figure 2

Prokaryotic communities in Rhizosphere (Rhizo), Bulk Soil (Bulk) and unplanted soil (Control). A. Phylum level, B. Class level and C. Genus level in DNA analysis.
A

Rhizo DNA
Rhizo cDNA
Bulk DNA
Bulk cDNA

B

Rhizo DNA
Rhizo cDNA
Bulk DNA
Bulk cDNA

C

Rhizo DNA
Rhizo cDNA
Bulk DNA
Figure 3

Prokaryotic communities in both DNA and cDNA in Rhizosphere (Rhizo) and Bulk Soil A. Phylum level, B. Class level and C. Genus level.

Figure 4

Functional gene analyses on DNA data. Number of gene families encoding enzymes of the biphenyl degradation: biphenyl 2,3-dioxigenase (bphAa, bphAb); cis-2,3-dihydro-2,3-dihydroxybiphenyl dehydrogenase (bphB); 2,3-dihydroxybiphenyl 1,2-dioxigenase (bphC). B. Number of gene families encoding enzymes related to oxidative stress: superoxide dismutase (SOD); thioredoxin (Trx); thioredoxin reductase (TrxRs); glutathione transport system (GSH TS); Catalase (CAT); peroxidase (POD); glutathione reductase (GSHRs). C. Number of gene families related to germination, sporulation and ABC transporters. D. Abundance of meta cleavage pathway of aromatic compounds.
Figure 5

Comparison of functional gene analyses on DNA and cDNA samples. A. Number of gene families encode enzymes of the biphenyl degradation: biphenyl 2,3-dioxigenase (bphAa, bphAb); cis-2,3-dihydro-2,3-dihydroxybiphenyl dehydrogenase (bphB); 2,3-dihydroxybiphenyl 1,2-dioxigenase (bphC). B. Number of gene families encoding enzyme and proteins related to oxidative stress: superoxide dismutase (SOD); thioredoxin (Trx); thioredoxin reductase (TrxRs); glutathione transport system (GSH TS); Catalase (CAT); peroxidase (POD); glutathione reductase (GSHRs). C. Number of Gene families related to germination, sporulation and ABC transporters. D. Abundance of meta cleavage pathway of aromatic compounds.

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