Illumina MiSeq sequencing investigation on the contrasting soil bacterial community structures in different iron mining areas

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Abstract Mine activities leaked heavy metals into surrounding soil and may affect indigenous microbial communities. In the present study, the diversity and composition of the bacterial community in soil collected from three regions which have different pollution degree, heavy pollution, moderate pollution, and non-pollution, within the catchment of Chao River in Beijing City, were compared using the Illumina MiSeq sequencing technique. Rarefaction results showed that the polluted area had significant higher bacterial alpha diversity than those from unpolluted area. Principal component analysis (PCA) showed that microbial communities in the polluted areas had significant differences compared with the unpolluted area. Moreover, PCA at phylum level and Matastats results demonstrated that communities in locations shared similar phyla diversity, indicating that the bacterial community changes under metal pollution were not reflected on phyla structure. At genus level, the relative abundance of dominant genera changed in sites with degrees of pollution. Genera Bradyrhizobium, Rhodanobacter, Reyranella, and Rhizomicrobium significantly decreased with increasing pollution degree, and their dominance decreased, whereas several genera (e.g., Steroidobacter, Massilia, Arthrobacter, Flavisolibacter, and Roseiflexus) increased and became new dominant genera in the heavily metal-polluted area. The potential resistant bacteria, found within the genera of Thiobacillus, Pseudomonas, Arthrobacter, Microcoleus, Leptolyngbya, and Rhodobacter, are less than 2.0 % in the indigenous bacterial communities, which play an important role in soil ecosystem. This effort to profile the background diversity may set the first stage for better understanding the mechanism underlying the community structure changes under in situ mild heavy metal pollution.

Keywords Iron mine · MiSeq sequencing · Alpha diversity · Bacterial phyla · Potential resistance genera · Heavy metal

Introduction

Mining activities adversely affect the quality of surrounding soils and aquifers and threaten human health and ecosystems (Anjos et al. 2012; Rodriguez et al. 2009). Studies on the relationship between mining and microbial communities in surrounding soils mainly focused on nonferrous metals (e.g., Cu, Pb, and Sb), which are highly toxic to soil biota (Zhang et al. 2007). However, few studies have focused on iron mining despite its specific environmental impact. In iron mining area, great amounts of tailing dumps and iron deposit rocks...
piled up directly on the ground and bring different kinds of harmful heavy metals (i.e., Cu, Zn, Cr, Cd, Pb) to surrounding soil through wind and hydraulic erosion. Besides, several surveys also indicated that iron mining areas, even heavily polluted mining area or tailing dam, always contain lower heavy metals contents than other nonferrous metal mining areas (Marmolejo-Rodríguez et al. 2007; Wang et al. 2012b; Huang et al. 2013). Pollution caused by iron mine activities was wider, gentler, and more recondite, and the microorganism which widely distributed in the heavy metal-polluted area may present more unique and subtle community changes. Study on soil microbe community structure could not only guide us in exploring the potential risk of affected soil but also provide reference for soil remediation with several indigenous resistance bacteria.

Recent developments in molecular fingerprinting techniques, such as denaturing gradient gel electrophoresis (DGGE), terminal restriction fragment length polymorphism, and real-time PCR quantification, have allowed researchers to investigate the microbial diversity variations in soils contaminated with heavy metals (Berg et al. 2012; Cáliz et al. 2013; Giloteaux et al. 2011; Petrić et al. 2011; Zhao et al. 2014). Molecular fingerprint techniques have high resolution; however, the fingerprint complexity caused by the high richness of indigenous microbial communities in soil may mask the changes induced by heavy metals (Gomes et al. 2010). Rare ecologically significant species that may have been affected may avoid detection by DGGE because of the detection limit (Bruce et al. 2000). The recent application of the next-generation sequencing methods, such as pyrosequencing 454 and Illumina, may provide a more direct way to detect the microbial taxa, especially those with low-abundant species changes (Oberauner et al. 2013; Uroz et al. 2013). In addition, Illumina sequencing is cost-effective and could obtain tenfold or more sequences per sample than pyrosequencing 454, thereby allowing the analysis of a high number of detailed taxonomic profiles from samples (Kozich et al. 2013).

Recent studies have focused on either the intense in situ or exogenous heavy metal impact on taxonomic community diversity. Berg observed that the relative abundance of bacterial groups changed within a Cu gradient (20 to 3537 mg/kg), which originated from CuSO4 contamination over 85 years ago (Berg et al. 2012). Margarita discussed the diversity of taxonomic groups after a 60-day incubation period with Cd concentrations reaching 12.5 mg/kg (Ros et al. 2009). However, these previous studies ignored the ubiquitous mild heavy metal pollution effect on soil microbes, which was apparent in the iron mine in our studies. The present study was conducted in the Anzigou iron mine, located north of Miyun County approximately 10 km upstream from the Miyun Reservoir in Beijing City. Frequent mining activities have caused heavy metal pollution in the Miyun Reservoir and have directly affected water quality and safety. Several researchers have investigated the heavy metal distribution in the catchment area of Miyun Reservoir (Huang et al. 2013; Qin et al. 2014). However, few studies have characterized potential ecological risks of such distribution.

In this study, to obtain the comprehensive effect of bacterial communities structure, we compared all soil types located in three regions. The three regions were as follows: (1) ore treatment factory, ore mining region, and tailing dam from mining area; and (2) croplands, planted forest, and vegetable fields from two villages (X and Y). The soils from each location from the same region have different physical and chemical properties, but they have the same pollution level. The present study, therefore, aims to provide detailed variations of bacterial communities that came from different soil types under mild heavy metal pollution and explore the natural bioremediation consortia.

Materials and methods

Soil sampling and heavy metal determination

The study area was the Anzigou iron mine (40° 36’ N, 117° 08’ E), which is located in the north of Miyun County, Beijing City, China, upstream of the Miyun Reservoir. The total sampled surface soil (0 to 20 cm in depth) was collected from three regions, namely H, M, and C, respectively, at the Anzigou iron mine in October 2012. Three H samples were collected from the ore treatment factory, ore mining region, and tailing dam. Three M samples were collected from three different soil types (croplands, planted forest, and vegetable fields) in village X, which are 1 km downstream of the tailing dam near the Chao River. Three C samples were collected from three different soil types (croplands, planted forest, and vegetable fields) in village Y, which is 5 km downstream of the tailing dam (Fig. 1). Sample at each site was collected in triplicates. The soil samples were packed in sterile Ziploc bags, and each bag was sealed and transported to the laboratory for further study. A fraction of the mixed sample was stored at −20 °C for molecular analysis. Another fraction was dried at room temperature for a week and sieved through a 4-mm grid to remove stones and visible plant fragments. An aliquot was threaded through a 0.125-mm sieve for the metal content analyses. All samples were stored at 4 °C. Water content in samples was determined immediately after collection by drying at 105 °C until constant weight. The pH was determined with a pH meter (Starter-3C, OHAUS, USA) in a 1:2.5 suspension of ultrapure water. Organic matter was determined with a 2400II CHNS/O elemental analyzer (PerkinElmer, USA) after removing the inorganic C by digesting in 1 M HCl for 48 h, washing with ultrapure water until becoming neutral, and then drying at low temperature. Soil samples were extracted by
sodium bicarbonate to determine the available phosphorus using the molybdenum antimony colorimetric method. Ammonium nitrogen contents (NH$_4^+$) were determined using the indophenol blue colorimetric method (Keeney and Nelson 1982). Nitrate contents (NO$_3^-$) were measured by dual wavelength spectrometry (Norman et al. 1985). The total concentration of the metals was measured after digesting the soil sample in a mixture of H$_2$O$_2$, HNO$_3$, and HCl (1:4:2, v/v) in a sealed digestion tank at 190 °C for 30 h. The metal concentrations were determined using an inductively coupled plasma optical emission spectrometer (Varian, 720-ES, USA).

DNA extraction and PCR amplification

Microbial DNA was extracted from 0.5-g soil samples using the Power Soil DNA kit (MoBio Laboratories, Solana Beach, CA) according to the manufacturer’s protocols. The V3–V4 regions of the bacterial 16S rRNA gene were amplified using PCR (95 °C for 2 min, followed by 25 cycles at 95 °C for 30 s, 55 °C for 30 s, 72 °C for 45 s, and a final extension at 72 °C for 10 min) using the primers (Dennis et al. 2013) 5′-barcode-ACTCCTACGGGAGGCAGCA)-3′ and 806R 5′-GGACTACHVGGGTWTCTAAT-3′. The barcode is an eight-base sequence unique to each sample. PCR reactions were performed in triplicate in a 20-μL mixture containing 4 μL of 5× Fast Pfu buffer, 2 μL of 2.5 mM dNTPs, 0.4 μL of each primer (5 μM), 0.4 μL of Fast Pfu polymerase, and 10 ng of template DNA.

Illumina MiSeq sequencing

Amplicons were extracted from 2 % agarose gels and purified using the AxyPrep DNA Gel Extraction Kit (Axygen Biosciences, Union City, CA, USA) according to the manufacturer’s instructions and quantified using QuantiFluor™-ST (Promega, USA). Purified amplicons were pooled in equimolar and paired-end sequenced (2 × 300) on an Illumina MiSeq platform (Majorbio, Shanghai) according to the standard protocols.

Processing and analyzing of sequencing data

Raw FASTQ files were de-multiplexed and quality-filtered using QIIME (version 1.17) with the following criteria: (i) The 300-bp reads were truncated at any site that obtained an average quality score of <20 over a 10-bp sliding window, and the truncated reads shorter than 50 bp were discarded; (ii) exact barcode matching, two nucleotide mismatch in primer matching, and reads containing ambiguous characters were removed; and (iii) only overlapping sequences longer than 10 bp were assembled according to their overlapped sequence. Reads that could not be assembled were discarded. Operational taxonomic units (OTUs) with 97 % similarity cutoff were clustered using UPARSE (version 7.1), and chimeric sequences were identified and removed using UCHIME.

The rarefaction analysis based on Mothur v.1.21.1 (Schloss et al. 2009) was conducted to reveal the diversity indices, including the Chao, ACE, and Shannon diversity indices. The beta diversity analysis was performed using UniFrac (Lozupone et al. 2011) to compare the results of the principal component analysis (PCA) using the community ecology package, R-forge (Vegan 2.0 package was used to generate a PCA figure). Venn diagrams were implemented by Venn Diagram, while Mantel test, Redundancy analysis (RDA), and Heatmap figures were performed in Vegan packages in R.

Results and discussion

Pollution site background

Table 1 presents the soil characteristics and heavy metal contents determined in the present study. Except for CV, pH value from other site changes slightly from 7.23 to 7.82. The soil acidification in site CV may be caused by excessive application of nitrogen fertilizer (Guo et al. 2010). Soil water contents ranged from 7.69 to 22.42 %, reaching the lowest value in region H. The organic matter contents had no obvious change in different sampling sites. In addition, the various values of available phosphorus, NH$_4^+$ and NO$_3^-$, were likely due to the fertilizing habit or land use pattern. The average Cr and Cd concentrations were 298.33 and 0.708 mg/kg, respectively, and maximum values were observed in group H. The Cd, Cu, and Cr concentrations in all samples exceeded the Chinese national background value (0.20, 35, and 90 mg/kg, the primary standard set by Chinese National Environmental Quality), with the average Cr and Cd
| Sampling sites | pH          | Water contents (%) | Organic matter (g/kg) | Available phosphorus (mg/kg) | NH$_4^+$ (mg/kg) | NO$_3^-$ (mg/kg) | Cd (mg/kg) | Cr (mg/kg) | Cu (mg/kg) | Zn (mg/kg) | Fe (g/kg) | Nemerow index |
|---------------|-------------|--------------------|-----------------------|-----------------------------|-----------------|-----------------|-------------|-------------|-------------|-------------|------------|----------------|
| HF            | 7.23 (±0.12) | 7.69 (±0.35)      | 0.77 (±0.001)         | 4.12 (±0.11)                | 2.09 (±0.05)    | 0.14 (±0.01)    | 0.69 (±0.01) | 280.60 (±10.09) | 47.32 (±1.09) | 101.23 (±3.48) | 89.27 (±3.25) | 3.01 |
| HM            | 7.69 (±0.20) | 9.10 (±0.19)      | 0.91 (±0.001)         | 5.54 (±0.14)                | 7.11 (±0.29)    | 0.57 (±0.01)    | 0.75 (±0.03) | 324.44 (±9.20)  | 62.11 (±2.31) | 79.12 (±3.29)  | 104.01 (±4.08) | 3.17 |
| HT            | 7.81 (±0.35) | 8.34 (±0.31)      | 0.55 (±0.002)         | 3.79 (±0.32)                | 1.25 (±0.04)    | 6.03 (±0.04)    | 0.68 (±0.02) | 294.25 (±7.28)  | 61.29 (±3.37) | 102.30 (±3.24) | 125.70 (±2.75) | 3.02 |
| Mean value    | 7.58        | 8.38               | 0.72                  | 4.48                        | 3.48            | 2.24            | 0.71        | 299.76       | 59.91        | 94.22         | 106.33       | 3.06 |
| MC            | 7.26 (±0.11) | 19.72 (±0.98)     | 1.01 (±0.004)         | 60.27 (±2.92)               | 1.09 (±0.03)    | 0.22 (±0.01)    | 0.49 (±0.03) | 156.29 (±5.09)  | 35.76 (±0.97) | 95.52 (±4.12)  | 29.36 (±1.02) | 2.05 |
| MF            | 7.19 (±0.42) | 18.39 (±1.02)     | 1.14 (±0.004)         | 27.54 (±1.12)               | 0.78 (±0.01)    | 19.13 (±1.88)   | 0.50 (±0.02) | 134.76 (±4.29)  | 50.67 (±4.05) | 117.45 (±4.38) | 31.83 (±0.96) | 2.12 |
| MV            | 7.56 (±0.18) | 21.95 (±0.68)     | 1.39 (±0.002)         | 58.54 (±2.16)               | 0.21 (±0.01)    | 24.02 (±1.27)   | 0.49 (±0.01) | 199.30 (±8.68)  | 47.44 (±3.55) | 106.83 (±3.27) | 30.02 (±1.88) | 2.14 |
| Mean value    | 7.34        | 20.02              | 1.18                  | 48.78                       | 0.69            | 14.45           | 0.49        | 163.45        | 44.62        | 106.60        | 30.40         | 2.10 |
| CC            | 7.27 (±0.20) | 19.31 (±0.87)     | 0.87 (±0.001)         | 5.24 (±0.41)                | 8.57 (±0.02)    | 5.47 (±0.02)    | 0.19 (±0.01) | 110.49 (±3.94)  | 49.01 (±2.04) | 87.10 (±4.03)  | 24.14 (±0.52) | 0.98 |
| CF            | 7.71 (±0.36) | 17.91 (±1.05)     | 0.72 (±0.002)         | 26.12 (±1.52)               | 0.98 (±0.01)    | 12.42 (±0.54)   | 0.17 (±0.02) | 98.75 (±2.05)   | 32.49 (±1.56) | 76.38 (±2.40)  | 27.67 (±0.81) | 0.88 |
| CV            | 5.07 (±0.45) | 22.42 (±1.51)     | 1.07 (±0.005)         | 13.88 (±0.79)               | 1.56 (±0.01)    | 51.33 (±1.88)   | 0.09 (±0.02) | 132.37 (±5.91)  | 36.70 (±1.95) | 92.39 (±4.57)  | 28.22 (±0.74) | 0.76 |
| Mean value    | 6.68        | 19.88              | 0.89                  | 15.08                       | 3.70            | 23.07           | 0.15        | 113.84        | 39.40        | 85.29         | 26.68         | 0.87 |

Values represent average ± standard deviation

*HF* heavily polluted sites from ore treatment factory, *HM* heavily polluted sites from ore mining region, *HT* heavily polluted sites from tailing dam, *MC* moderately polluted sites from croplands, *MF* moderately polluted sites from planted forest, *MV* moderately polluted sites from vegetable fields from village X, *CC* clear sites from croplands, *CF* clear sites from planted forest, *CV* clear sites from vegetable fields from village Y.
contents reaching values three to four times higher than the primary standard. The Zn concentration in all sites was higher than the background value in Beijing (57.5 mg/kg) but was not higher than the background value in China (100 mg/kg). The contents of the other heavy metals (Pb, Ni, and Co) were below the Beijing background (24.6, 26.8, and 7.5 mg/kg, data not shown). Although the Fe concentration was extremely high in all samples, it is not regarded as an indicator of metal pollution. The Nemerow composite index, a widely used pollution indicator and has been utilized by Wang (Wang et al. 2012b), showed that the soil samples from region H had the highest index values (Table 1), thereby indicating that the site was heavily polluted. Soil from region C was clear and used as a point of comparison. The surroundings of the mining area (MC, MF, and MV) were moderately polluted by a low diffusion of contaminants into the surrounding environment. However, considering the special characteristics of the Miyun Reservoir as a centralized urban drinking water source, stricter criteria (the primary standard) were selected to evaluate the heavy metal pollution level, and the results were much more serious than the other surveys which based on the secondary standard (Wang et al. 2012b).

Sequencing results and diversity indices

A total of 110,409 reads and 10,444 OTUs were obtained from nine samples through 468 MiSeq sequencing analysis. Each library contained 7959 to 22,327 reads, with different phylogenetic OTUs ranging from 724 to 1483. All rarefaction curves tended to approach the saturation plateau, indicating that the data volume of sequenced reads was reasonable, and the discovery of a large variation in the total number of OTUs in different samples (Fig. 2). Compared with the contaminated soil (e.g., MC, MF, and MV), all samples from unpolluted soils have lower OTU density. The OTU densities of samples CF and CV were even lower than the average of most seriously contaminated soil samples (group H), and these samples’ reads were not the least among the samples. In addition, the average OTUs of group C attained the lowest value (978), but its average reads (15,048) were significantly higher than those of group H (9851, Table 2).

Moreover, the calculation of the alpha diversity species richness (Chao), evenness (ACE), and Shannon index all confirmed the increasing diversity in the contaminated soil, especially in group H compared with group C (Table 2). The results of ANOVA analysis of the Shannon diversity indices also show significant differences with P value of 0.037, and the following post hoc pairwise test showed that groups H and M were significantly different from group C. The average Shannon diversity indices of the three groups were 6.15 (group H), 6.11 (group M), and 5.57 (group C). These results indicated that the Shannon diversity of polluted areas H and M was significantly higher than that of the unpolluted area C.

Taxonomy composition of the iron mining area

Sequences that could not be classified into any known group were assigned as unclassified. The bacterial OTUs were assigned into 35 different phyla, 251 families, or 381 genera. Six different phyla (Proteobacteria, Acidobacteria, Actinobacteria, Bacteroidetes, Chloroflexi, and Gemmatimonadetes) out of the 35 total phylotypes were common to the nine libraries, which comprised more than 85 % of the total reads in every library. Proteobacteria was the most abundant group (Fig. 3a), comprising approximately 29.54 % (3086) of the OTUs and 39.95 %
(50,590) of the reads across all samples. *Acidobacteria*, the second most abundant phylum, (14.78 %, 1544 OTUs) consisted 18.70 % (23,682 reads) in all libraries. However, the proportion of *Proteobacteria* in the different samples showed high variability, in the range of 6.99 to 56.41 %. The read proportion reached lowest value in CV, which was different from that in group C (Fig. 3a). Meanwhile, *Acidobacteria* comprised the majority of the total reads in CV because of the strong acidic nature of the soil. The members from *Bacteroidetes* (13.61 %, 1422 OTUs), *Chloroflexi* (12.11 %, 1195 OTUs), *Actinobacteria* (10.45 %, 1091 OTUs), and *Gemmatimonadetes* (5.10 %, 533 OTUs) comprised 13.60 % (17,227 reads), 6.99 % (8848 reads), 8.11 % (10,266 reads), and 2.70 % (3424 reads) in all libraries, respectively. A much smaller fraction of the members of *Candidate division_TM7* (3.11 %, 325 OTUs; 1.84 %, 2333 reads), *Verrucomicrobia* (2.85 %, 298 OTUs; 1.99 %, 2530 reads), *Cyanobacteria* (1.49 %, 156 OTUs; 1.65 %, 2085 reads), and *Nitrospirae* (1.37 %, 143 OTUs; 1.70 %, 2333 reads) were found. The average reads of the unclassified group account for 2.27 %, but this value fluctuated in different sites.

The members of *Alphaproteobacteria* dominated the *Proteobacteria* phylum and occupied 9.79 % (1023 OTUs) of all libraries but accounted for 22.44 % of the total reads (Fig. 3b). The subdivision of *Betaproteobacteria*, *Deltaproteobacteria*, and *Gammaproteobacteria* comprised 9.65, 4.34, and 9.34 % of the total reads, respectively. The *Epsilonproteobacteria* only contained two OTUs (0.019 %) and comprised a significantly small fraction (0.06 %) in the total reads.

**Correlation between community structure and environmental factors**

Mantel test was performed to analyze the environmental factors responsible in shaping soil bacterial community structure along a pollution gradient. Mantel test results showed that soil bacteria was significantly correlated with heavy metal contents (mental \(r=0.647, P<0.05\); Table S1), whereas soil properties had no obvious effect on bacteria (mental \(r=0.364;\) Table S1) indicating that heavy metal pollution played a more important role in shaping local bacteria community.

To further identify the major environmental variables controlling the soil microbial community structure, RDA was performed (Fig. 4). Parameters that could explain the variance of bacteria community structure were ranked by RDA results (Table 3). Results indicated that both heavy metal contents (Cd, Cr, Cu, Fe) (\(P<0.05\)) and soil pH (\(P<0.01\)) were identified to be the most influential environment factors to drive changes occurring in the community composition.

**Comparison of bacterial community structure among groups**

Principal component analysis (PCA) was used to identify the community structure differences under different pollution levels (Fig. 5). The resolutions of PCA between phylum, genus, and OTU levels were different. Except for CV, separating other locations at phylum level was different, indicating that communities in most locations shared similar phyla diversity. A much better discrimination was exhibited at the genus level, and group H grouped together, whereas the other groups had no clear boundary. The best resolution was obtained at the OTU level with all reads counted (Fig. 5c). At OTU level, PCA demonstrated that different soil locations from groups H and M were clustered, and the locations from group M (moderately polluted area) gathered together and were closer to group H (heavily polluted area communities) than to group C (unpolluted area). In addition, sites from group H gathered closer (similar) than the other groups, thereby indicating the higher number of similar community structures in heavily polluted areas. These results demonstrated that the microbial communities in the polluted areas had significant differences compared with the unpolluted area. This finding was consistent with the abovementioned alpha analysis results. However,
the effect of soil properties on the community difference cannot be ignored. PCA at all levels demonstrated that the large gap between CV and the other regions in the unpolluted area was in accordance with the fact that the predominance of Acidobacteria in CV was closely related with the strong acidic nature of the site. Many research studies have confirmed that this phenomenon had close relationship with the strong acidic nature of soil (Jones et al. 2009).

Moreover, different from the PCA at genus and OTU level, PCA at phylum indicated that three groups have no clear boundary. To test the effect of heavy metal pollution on phyla structure, Matastats (White et al. 2009) were used to conduct differentially abundant features between three groups (Table 3). Matastats results demonstrated that three groups have significant difference on Bacteroidetes, groups H and C have significant difference on Actinobacteria, and groups have no obvious difference on the other phyla. Those findings raise the question of whether heavy metal pollution might have less affected on phyla structure.

Differences between the bacterial genera among different pollution levels

At genus level, the differences between the communities from various areas with different pollution levels were detected via a Venn diagram. A total of 341 genera were discovered, and 79.76 % of them belong to the shared genera (see Fig. S1). Group M alone contained more bacterial varieties (14 genera) than groups M (eight genera) and C (six genera), as shown in Fig. 6. An overlap between the detected genera of the three areas was also observed. The largest overlap was found between groups M/C (21 genera) followed by groups H/M (17) and groups H/C (3). Several new genera appeared in group H, indicating that these genera may thrive in heavily polluted areas. These genera included the following: Betaproteobacteria (36 reads, Sulfuricella, and 16 reads, Candidatus Nitrotoga), Bacteroidetes (4 reads, Ectricia and 3 reads, Larkinella), Actinobacteria (3 reads, Modestobacter), and Cyanobacteria (2 reads, Arthonema).

However, their abundance was low, and their relative abundance was less than 0.05 %. Compared with group C, the five most abundant characteristic genera present in the polluted region (groups H and M), according to the read abundance, were as follows: Alphaproteobacteria (258 reads and 0.23 %, Paracoccus), Cyanobacteria (180 reads and 0.16 %, Microcoleus), Cyanobacteria (112 reads and 0.11 %, Leptolyngbya), Betaproteobacteria (98 reads and 0.09 %, Thiobacillus), and Deltaproteobacteria (68 reads and 0.06 %, Peredibacter).

Based on the relative abundance of the genera from Fig. 7, the genera with an average abundance of >1 % in at least one group were defined as dominant. Combined with the Venn diagram, most dominant genera belonged to the genera shared by the three

Table 3  Significance of environmental variables in explaining the bacteria community structure obtained from the RDA results

|          | r² | P value |
|----------|----|---------|
| Soil     |    |         |
| Water contents | 0.309 | 0.315 |
| pH       | 0.952 | 0.001** |
| Organic matter | 0.066 | 0.826 |
| Available phosphorus | 0.002 | 0.994 |
| NH₄⁺  | 0.038 | 0.899 |
| NO₃⁻  | 0.042 | 0.832 |
| Heavy metal |    |         |
| Cd       | 0.862 | 0.001** |
| Cr       | 0.674 | 0.011*  |
| Cu       | 0.622 | 0.033*  |
| Zn       | 0.375 | 0.232 |
| Fe       | 0.576 | 0.046*  |

r² indicates the decision coefficient of environmental variables on the community structure

*P<0.05; **P<0.01
groups. Their relative abundance changed in sites with degrees of pollution, as shown in Fig. 8, except for the unknown genera. In group C, the dominant genera were Sphingomonas, Bradyrhizobium, Blastocatella, Bryobacter, Rhodanobacter, Reyranella, Rhizomicrobium, and Nitrospira. Sphingomonas was the most abundant in the three groups, as seen in Fig. 6. The genera Bradyrhizobium, Rhodanobacter, Reyranella, and Rhizomicrobium significantly decreased with increasing pollution degree (from groups C to H), and their dominance decreased. Meanwhile, the relative abundances of Blastocatella, Bryobacter, and Nitrospira slightly decreased, but their dominance was maintained. The relative abundances of genera Cellvibrio and Flexibacter increased and reached more than 3% in group M. However, their abundances significantly decreased in group H. The abundance of the following genera was still significantly higher in group H than in the other groups: Steroidobacter, Massilia, Arthrobacter, Flavisolibacter, and Roseiflexus, which belong to Gammaproteobacteria, Betaproteobacteria, Actinobacteria, Bacteroidetes, and Chloroflexi, respectively.

Bacteria with potential resistance to heavy metals

To verify the potential resistance of abundant genera in heavy metal pollution area, Pearson correlation between the genera relative abundance and heavy metal contents was performed. Arthrobacter, which is atypical Cr⁶⁺ reducing bacteria (Asatiani et al. 2004), turned out to be closely related to Cr content in mining area (R²=0.726, P<0.05) and became the dominant genera in group H. Thiobacillus was known as an important genus closely related to catalyzing ferrous ions and oxidizing the reducing inorganic sulfur (Dopson et al. 2003).In our samples, Thiobacillus had relative high abundance in the genera which only found in polluted area and its reads abundance in group H.
is tripled in group M. Moreover, *Thiobacillus* relative abundance had close relationship with Fe content ($R^2=0.691$, $P<0.05$). In addition, most *Cyanobacteria* can produce extracellular polymeric substances, mainly polysaccharide, which could adsorb heavy metals dispersed in the environment (Pereira et al. 2009, 2011). In this study, *Cyanobacteria* was a rare group, but its total read abundances in groups H (795 reads) and M (1202 reads) were significantly higher than in group C (92 reads). *Microcoleus*, accounted for 8.75 % of the total *Cyanobacteria* species, displayed a strong correlation with Zn content ($R^2=0.789$, $P<0.05$). The genera *Rhodobacter* was also reported to be resistant against several metals and could show passive or active uptake of metals (Italiano et al. 2009; Mishra and Malik 2013). Based on the Venn diagram, *Rhodobacter* only appeared in polluted area. Besides, Pearson analysis showed that Zn content had a substantial correlation with *Rhodobacter* ($R^2=0.801$, $P<0.01$). The population of resistant bacteria comprised less than 2 % of the indigenous microbial communities. However, the number of resistant bacteria can increase in contaminated soils, reaching 10 % or even 100 %, as the contaminants greatly change in terms of quantity and composition (Xu et al. 2012). Moreover, these bacteria serve a significant function in maintaining the ecological balance of the contaminated soil and could be used for the bioremediation of metal-polluted soil (Hayat et al. 2010; Kandeler et al. 2000). Numerous studies have reported a few naturally occurring microorganisms with high metal resistance in the rhizosphere that significantly increases the metal uptake in plants and reduces metal toxicity (Ma et al. 2011; Sessitsch et al. 2013).

Furthermore, several other genera were found similar to *Arthrobacter*, as the dominant genera in group H, and significantly related to heavy metals contents. *Steroidobacter* has
been reported to utilize testosterone under both oxic and anoxic conditions. The genus *Massilia* was mainly isolated from the blood of an immunocompromised patient (Lindquist et al. 2003) and has been recently found in environmental samples, such as soil samples from China and in the drinking water distribution system in Seville (Spain) (Gallego et al. 2006; Wang et al. 2012a). In this study area, *Steroidobacter* was found to closely relate to Cd content ($R^2=0.729$, $P<0.05$), while *Massilia* had significant correlation with Cd and Cr contents ($R^2=0.718$, $P<0.05$; $R^2=0.671$, $P<0.05$). However, none of them were discovered to be resistant to metals in recent research studies. Their wide spread in heavily polluted area and close relation with heavy metals may be affected by several other factors, such as the emission of beneficiation agents or wastewater, ore tailing accumulation, and availability of nutrients and oxygen, and water content in soil. These factors need further investigation.

**Conclusions**

In conclusion, the present study, using the high throughput Illumina sequencing method, provided a detailed picture of bacterial community variations on phylum and genus level under heavy metal pollution. Sequencing results and alpha diversity indices indicated that diversity of polluted areas, including ore treatment factory, ore mining region, tailing dam in H areas and three different fields (croplands, planted forest, and vegetable) in M areas which are only 1 km downstream of the tailing dam, was significantly higher than those of unpolluted areas. The total bacterial OTUs could be assigned into 35 different phyla, 251 families, or 381 genera. Among them, six different phyla (*Proteobacteria*, *Acidobacteria*, *Actinobacteria*, *Bacteroidetes*, *Chloroflexi*, and *Gemmatimonadetes*) were common to the nine libraries, which comprised more than 85 % of the total reads in every library. PCA and Matsats stats results implied that heavy metal pollution has less affected on these phyla structure (Table 4). At genus level, some new genera appeared in polluted area, and the five most abundant genera were as follows: *Paracoccus*, *Microcoleus*, *Leptolyngbya*, *Thiobacillus*, and *Pereidibacter*. Moreover, the relative abundance of dominant genera changed in sites with degrees of pollution, and several genera (e.g., *Steroidobacter*, *Massilia*, *Arthrobacter*, *Flavisolibacter*, and *Roseiflexus*) increased and became new dominant genera in the heavily metal-polluted area. The study found that some genera have close relation with heavy metals and may provide a new way to explore natural bioremediation genera. However, several genera which had not been reported for their resistance to metal

![Fig. 8](https://example.com/fig8.png)

Relative abundance of dominant genera in different groups. The genera with an average abundance of >1 % in at least one group were defined as dominant.

| Phylum                     | $P$ value between groups HF, HM, and HT and groups MC, MF, and MV | $P$ value between groups HF, HM, and HT and groups CC, CF, and CV | $P$ value between groups MC, MF, and MV and groups CC, CF, and CV |
|----------------------------|------------------------------------------------------------------|-----------------------------------------------------------------|-----------------------------------------------------------------|
| Proteobacteria             | 0.317                                                            | 0.159                                                           | 0.909                                                           |
| Acidobacteria              | 0.347                                                            | 0.609                                                           | 0.329                                                           |
| Actinobacteria             | 0.060                                                            | 0.028*                                                          | 0.432                                                           |
| Bacteroidetes              | 0.011*                                                           | 0.035*                                                          | 0.006**                                                          |
| Chloroflexi                | 0.926                                                            | 0.274                                                           | 0.472                                                           |
| Gemmatimonadetes           | 0.655                                                            | 0.946                                                           | 0.853                                                           |
| Candidate_division_TM7     | 0.699                                                            | 0.882                                                           | 0.794                                                           |
| Verrucomicrobia            | 0.324                                                            | 0.292                                                           | 0.377                                                           |
| Cyanobacteria              | 0.803                                                            | 0.349                                                           | 0.180                                                           |
| Nitrospirae                | 0.419                                                            | 0.324                                                           | 0.782                                                           |

$P$ value means an individual measure of the false positive rate

* $p<0.05$; ** $p<0.01$
turned out to be dominant in polluted area and needed further study to explore their influencing factors.

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