Fungal and metabolome diversity of the rhizosphere and endosphere of *Phragmites australis* in an AMD-polluted environment

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**ABSTRACT**

Symbiotic associations with rhizospheric microbial communities coupled with the production of metabolites are key adaptive mechanisms by metallophytes to overcome metal stress. However, little is known about pseudo-metallophyte *Phragmites australis* interactions with fungal community despite commonly being applied in wetland phytoremediation of acid mine drainage (AMD). In this study, fungal community diversity and metabolomes production by rhizosphere and root endosphere of *P. australis* growing under three different AMD pollution gradient were analyzed. Our results highlight the following: 1) Ascomycota and Basidiomycota were dominant phyla, but the diversity and richness of taxa were lower within AMD sites with *Penicillium*, *Candida*, *Saccharomycetales*, *Vishniacozyma*, *Trichoderma*, *Didymellaceae*, and *Cladosporium* being enriched in the root endosphere and rhizosphere in AMD sites than non-AMD site; 2) non-metric multidimensional scaling (NMDS) of 73 metabolomes revealed spatially defined metabolite exudation by distinct root parts (rhizosphere vs endosphere) rather than AMD sites, with significant variability occurring within the rhizosphere correlating to pH, TDS, Fe, Cr, Cu and Zn content changes; 3) canonical correspondence analysis (CCA) confirmed specific rhizospheric fungal taxonomic changes are driven by pH, TDS, heavy metals, and stress-related metabolomes produced. This is the first report that gives a snapshot on the complex endophytic and rhizospheric fungal community structure and metabolites perturbations that may be key in the adaptability and metal phytoremediation by *P. australis* under AMD environment.

1. Introduction

In the last century, tremendous global development has been associated with anthropogenic activities that have led to the degradation of soil and water environments. Acid mine drainage (AMD), an outflow of highly acidic and metal-rich water from mineral mining sites, is one of the worldwide problems leading to ecological destruction of watersheds and the contamination of human water sources by inorganic acids, coupled with heavy metals toxicity. Due to stringent federal and state environmental regulations, AMD wastes must be treated before discharge. Therefore, several AMD remediation technologies, mostly focusing on the treatment of already produced AMD before their discharge into water bodies, are currently being applied across the globe [1]. Phytoremediation is one of the emerging technologies for the reclamation of AMD-polluted water and soil environments due to higher costs associated with conventional AMD remediation approach [2].

Under a metalliferous environment, metal tolerant plant species (either obligate/facultative metallophytes or hyperaccumulators) have developed adaptive mechanisms that enable them to survive the elevated metal levels. Stress tolerant and pseudometallophyte *Phragmites australis*, a common perennial rhizomatous and stoloniferous wetland reed frequently used in passive treatment systems of AMD-polluted and other contaminated environments [3, 4, 5, 6], possesses the ability to accumulate more phosphorous and potassium than other macrophytes, in addition to enhancing the removal of excessive nutrient ions including heavy metals in the environment. It has also been reported as a pesticide degrader including organochlorine compounds via peroxidase pathway [7], and removing heavy metals from polluted soils by accumulating them in the roots, stems, and leaves [8, 9, 10]. Under such environment, the symbiotic plant-microbe-metal interactions involving root exudates and microorganisms in rhizosphere ecology play significant role in
promoting host adaptation to metal toxicity, phytoremediation and other ecological stress [11, 12].

Generally, plant exudates a cocktail of small and high molecular organic and inorganic molecules in their rhizosphere, creating a nutrient-rich microenvironment that may select for specialized metal-tolerant microorganisms (bacteria and fungi) [13, 14]. In turn, the microbes acting as plant growth promoting microorganisms (PGPMs), may play critical role in adaptation to metal phytotoxicity and stimulate plant growth either indirectly through inducing plant defense mechanisms against phytopathogens, and/or directly through the mineral nutrients solubilization (nitrogen, phosphate, potassium, iron, etc.), production of plant growth promoting substances (e.g., phytohormones), and secretion of specific enzymes (e.g., 1-amincyclopropane-1-carboxylate deaminase) [11, 12, 15]. For example, P. australis has been reported to maintain the carbon-rich rhizosphere through the production of metabolites such as phenols, short-chain organic acids, sugars, and amino acids, that act as an abundant source of energy and nutrients for rhizospheric microbes [10]. Additionally, the rhizosphere maintains constant oxygenation of the rhizosphere that enhances the growth of rhizospheric microorganisms that promotes the oxidation of ammonia to nitrate for plant absorption [16], thereby supporting the atmospheric nitrogen cycle. These beneficial interactions help the reed to flourish by colonizing a wide soil volume, quickening photosynthesis, protection from plant pathogens, and enhances their survival in stressed environments. However, such interactions are complex and multifaceted, and thus rhizobiome assembly and the involvement of the P. australis in these processes are currently enigmatic. It can be hypothesized that any change in the environment may result in the corresponding change in the utilization of plant root exudates and the production of stress-induced metabolites that may influence the diversity of metal-tolerant plant growth-promoting rhizospheric and metallophyte-associated endophytic microbial communities. Thus, there has been increased research interest in the last two decades to understand the mechanisms of plant-microbe-metal interactions under stressful environmental conditions that will facilitate wider application of metallophytes in phytoremediation.

There is ample evidence that Phragmites harbors rich endophytic and epiphytic fungal communities with the potential to play a key role in their adaptability under different environments [17, 18, 19]. According to Dolinar and Gaberscik [20], rhizosphere fungi possess the ability to form living root-soil links, a zone of the soil generally referred to as rhizosphere, capable of adjusting the soil properties, microbial networks, or potential root exudates which include metabolites to promote the growth of associated host plant. The role of fungal endophytes in synthesizing defensive compounds and metabolites that enhance the host’s survival in stress conditions has also been reported [11, 21]. However, there is a paucity of knowledge on fungal endophytes and ectophytes as well as the community composition of fungi which colonize the rhizosphere of P. australis, especially under AMD-polluted environment, despite commonly being applied in wetland remediation of AMD sites. To the best of our knowledge, the impact of the differential production of metabolites in the rhizosphere-root continuum of P. australis in AMD environments and its perturbations on rhizospheric fungal community and functional diversity and dynamics has also not been reported elsewhere.

In the present study, we adopted a microcosm-based culture-independent metagenomic and metabolomic approach to gain more knowledge into the nature of in situ microbial community structure and dynamics, and the rhizospheric metabolites production within P. australis rhizosphere under AMD-polluted environments. Specifically, the study examined: (1) the community structure and dynamics of rhizospheric soil and endophytic fungi associated with the P. australis roots under different AMD-pollution gradient environment; (2) the impact of the AMD pollution gradient on primary metabolites production; (3) the link between fungal community structure, primary metabolites production and the physicochemical characteristics of the AMD environment.

## 2. Materials and methods

### 2.1. Study area and sampling

Two sites in Randfontein, Gauteng Province, South Africa, Lan 3 (S 2607.820, E02746.680 and elevation of 1693 m), a tailing dam of Minto, and Florida Lake (S2610.625, E02754.220 and 1673 m elevation), a tailing dam of Sibanye Gold Mine were selected for the study. Whereas Florida Lake is characterized by dense and robust growth of P. australis on the banks, the two AMD sites had lesser densities of the reed plant, especially at Lan 3 site. All the sites have been in operation for about 20 years. Sampling was conducted under permission from the Johannesburg City Parks and the two mining companies in June 2018.

At each site, we excavated 3 whole plants including the surrounding soils from each of the three quadrats (20 cm × 20 cm) 50 m apart on the shore of the AMD ponds. A total of 9 samples (plants and surrounding rhizospheric soil) were collected in plastic bags and transported on ice (≤4 °C) to the laboratory, and root systems immediately processed (by separating the rhizosphere soil and root compartments that forms the root endosphere) within 6 h for metabolites analysis. Fractionation of the rhizosphere soil and the root endosphere was done based on the procedure described by Yamamoto et al. [22]. For metagenomic experiments, samples were stored at -80°C until analysis.

### 2.2. Physicochemical analysis

Water was scooped with a bucket from each sampling point and physicochemical parameters, namely, pH, salinity, total dissolved solutes (TDS) and electrical conductivity (EC) were measured on-site using Hanna HI9828 multi-parameter ion-specific meter (Hanna Instruments Pty) Ltd, Bedfordview, South Africa). Rhizospheric soil were initially freeze dried, followed by homogenization by grinding in a pestle and mortar before passing through a 200-mesh sieve. Metal content in mg/L (Fe, Ca, Mn, Cd, Cr, Co, Na, and Pb), was analyzed in an Inductively Coupled Optical Emission Spectrometer (Agilent Technologies 700 series ICP-OES), following acid microwave digestion of 0.5 g of the soil samples as described previously [23]. One-way analysis of variance (ANOVA) and Tukey’s Honest post hoc test (p < 0.05) was done using SAS version 9.4 (SAS Institute, Cary, NC, USA) to compare the differences in metal contents among the sampling sites.

### 2.3. DNA extraction, PCR amplification, and MiSeq high-throughput sequencing

Root endosphere samples were ground into fine particles suitable for DNA extraction. Fecal/Soil Total DNA™ extraction kit (Zymo Research Corporation, CA, USA) was used to extract total DNA from 0.25 g of rhizosphere soil or ground root samples according to manufacturer’s protocol. The quantity and the quality of the resultant environmental DNA was checked on a 1.5% agarose gel and Biodrop Lite spectrophotometer (Biochrom Ltd, Cambridge, UK). Extracted DNA used for downstream PCR and sequencing analysis had A260:A280 ratios between 1.8–2.0 and DNA concentrations of 20–150 ng/μL. Equimolar concentration (5 ng) of the environmental DNA of a replicate sample type (endospheric or rhizospheric) for each site were pooled together (9 replicates) to get a representative sample for downstream library preparation.

Libraries of nuclear ribosomal ITS region were amplified from each resultant pooled DNA products using ITS1 (5’-CTTGGTCCTAGAGGACGTAAGTCACGCGGGGGGCGGCGG-3’) and ITS4 (5’-TCCCTCCTGCTTATGATATGCAGCAGCCG-3’) primers fused with Illumina MiSeq adapters and barcode sequence (underlined) unique to...
each sample according to MiSeq standard protocol (https://support.illumina.com/documents/documentation/chemistry_documentation/16s/16s-metagenomic-library-prep-guide-15044223-b.pdf). The resultant amplified DNA were cleaned and sequenced by paired-end (300 bp reads each) sequencing on the Illumina MiSeq Platform at Inqaba Biotechnology (Pretoria, South Africa).

2.4. Data processing and ITS gene-based community analysis

The obtained Fastq datasets were initially scrutinized for PCR artefacts and low-quality reads (reads with >50% bases having a quality score < 2) using ngsShoRT (next generation sequencing short reads) trimmer as described by Chen et al. [24]. The merging of the forward and reverse sequences was done following the protocols of Mago et al. [25]. The resultant sequences were further strictly filtered to remove low quality sequences as described by Bokulich et al. [26], before being subjected to QIIME pipeline (V2.0, https://qiime2.org/) [27]. According to the method described by Edgar [28], chimeric sequences were removed, and the sequence clustered into operational taxonomic units (OTU) at 97% sequence similarity [29]. Finally, the effective sequence was assigned to representative microbial taxa using the RDP (Ribosomal Database Project) classifier [30].

Alpha diversity indices of Chao1, Shannon, Simpson, evenness, Fisher alpha dominance and Good's coverage were calculated at the genetic distance of 0.03 using the plot richness function of phyloseq package in R [31] to reflect the diversity and richness of the fungal community in different samples. The dominant OTUs at different taxonomic levels were used to generate heatmap using ampvis2 package in R [32], respectively, to visualize the variations and distributions of present fungal communities. β-diversity-based Bray-Curtis dissimilarity distance and canonical correspondence analysis (CCA) to visualize the community relationships between and within each AMD environments with explanatory environmental variables was also performed using Paleontological statistics software package Version 4 (PAST4) [33]. OTU counts were used as “species” data, while AMD site physicochemical data were included in the analysis as “environmental” variables. The most significant variables were initially picked by forward selection at $p < 0.01$ baseline before being included in the final model whose significance was tested with 999 permutations.

2.5. Metabolome analysis

For metabolite extraction, 10 g freshly collected rhizosphere soil and root samples were ground before mixing with 1 ml 75%:25% v/v methanol: water solution. The mixture was vortexed briefly, then sonicated for 10 min and centrifuged at 13,000 rpm for 5 min. The supernatant was then filtered with a 0.22 µm Whatman filter membrane and transferred into vials. Then, the samples were directly analyzed to determine their metabolomic profiles including organic acids, vitamins, and signaling molecules using LCMS-8040 triple quadrupole mass spectrometer (Shimadzu Corporation, Kyoto, Japan), equipped with Shim-pack Velox® PFPP (pentafluorophenylpropyl) column (Shimadzu Corporation, Kyoto, Japan). The instrument settings for the LC/MS/MS were as follows; total flow = 0.4 mL/min, injection volume = 1 µL, oven temperature = 40 °C/max 85 °C, nebulizing gas flow = 3 L/min, drying gas flow = 15 L/min and the mobile phase was 50%/50% acetonitrile/water. Peak detection, data mining, alignment and normalization, removal of isotope masses, and library searching was performed using LabSolutions Insight® multi-analyte quantitation software (Shimadzu Corporation, Kyoto, Japan).

To compare the peak abundance of molecular features, peak area values were exported from LabSolutions Insight® to Microsoft Excel. The major metabolites peak areas were used to generate stacked bar charts and heatmap using ggplot2 [34] and heatmap.2 packages [35] in R, respectively, to visualize the variations and distributions of present metabolites in the rhizospheric soil and roots samples. Using metMDS and anosim function of the vegan package in R [36], non-metric multidimensional scaling (NMDS) and ANOSIM, respectively, was calculated to explore the metabolite data structure based on dissimilarity measurement (Bray-Curtis dissimilarity) between three AMD environments and localization (rhizosphere versus endosphere). The NMDS plots were generated using ggplot2 package in R. When the ANOSIM analysis suggested significant differences in metabolome structure between groups, we then determined which metabolites were driving those differences by

![Figure 1. Map showing the sampling sites (Mogale Gold Mine, Sibanye Gold Mine and Florida Lake).](image)
identifying indicator species (metabolites) using the R package \textit{indicspecies} \cite{37}. This analysis generates an indicator value (\textit{IndVal.g}) that measures the association between OTUs with each group or combination of groups and then identifies the group corresponding to the highest association value. In this study, indicator metabolites based on an \textit{IndVal} of >0.30 and a \( p < 0.05 \) assessed after 999 permutation tests.

2.6. Data availability

All the data analysis results obtained during this study are included in the manuscript (and its Supplementary 36 Information files). All the raw datasets from Illumina sequencing have been deposited at the NCBI database (https://www.ncbi.nlm.nih.gov/) sequence archive (SRA) as BioProject ID PRJNA640287.

3. Results

3.1. Physicochemical profiles of AMD samples

Physicochemical parameters of the three sampling sites (two acid mine sites (Lan 3 and Wuinze 17) and the control site (Florida Lake)) showed significant differences for each of the parameters measured (Table 1). The acid mine sites had significantly (\( p < 0.05 \)) higher EC, salinity, TDS, and low pH when compared to the non-AMD control site (Table 1). Further, the heavy metal analysis also showed significantly higher (\( p < 0.05 \)) values in the AMD sites rhizospheric soil samples, that were >5-fold higher, than the values for the non-AMD control site, especially with Fe, Na, Cr, and Zn (Table 1).

3.2. Root endosphere and rhizosphere fungal community diversity and distribution under different AMD pollution

3.2.1. Diversity analysis of fungal communities

Fungal diversity analysis based on the nuclear ribosomal ITS region analysis resulted in 382,391 quality reads and 707 OTUs across six samples (Table 2). Good’s coverage across the sample was >98%, implying that the sampling depthness was enough to estimate the diversity enclosing all major fungal groups inhabiting the studied root endosphere and rhizosphere of \textit{P. australis} under the different AMD environments (Table 2). This was further supported by the rarefaction curves (Figure 2a) and rank abundance plot (Supplementary Information Figure 1) that asymptotically approached a plateau, suggesting that the curves accurately reflected the fungal community.

According to observed OTU numbers, the sample FL.A (154) had the richest diversity, followed by FL.E (122), Wuinze 17.A (119), and Lan3 A that had comparable moderate diversities, whereas rhizosphere samples Wuinze17.A (114) and Lan3.E (90) had less richness. Similarly, the calculated alpha diversity indices (dominance, Chaol index, Shannon index, Simpson’s index, and evenness) showed that FL samples (both rhizosphere and root endosphere) had more representation and evenness of the taxa than other sites (Table 2). All the biodiversity indices separated the acid mine sites (Lan 3 and Wuinze 17) from the non-acid mine site (Florida Lake). The shared fraction of core fungal OTUs associated with the rhizosphere and root endosphere is summarized in Figure 2b. Based on the Venn diagram analysis, 70 and 63 OTUs constituted the core microbiota in the rhizosphere and root endosphere, respectively, accounting for ~99% of the total abundances. In the rhizosphere, 20 OTUs (78.7% abundance) were common in all the three AMD sites, with 3 OTUs shared between Lan 3 and Wuinze 17, 5 OTUs between Florida Lake and Lan 3, and 6 OTUs between Florida and Wuinze 17.

3.2.2. Taxonomic composition of the AMD environment fungal communities

Based on relative abundances, fungal groups in the different \textit{P. australis} root endosphere and rhizosphere grouped into 8 phyla, 28 classes, 84 orders, and 259 genera. Figure 3 summarizes the abundance of the fungal community at phylum, class, and genera level. Overall, members of phylum \textit{Ascomycota} and \textit{Basidiomycota} were the most abundant across all the sampling sites, accounting for >75% of the observed taxa (Figure 3a). Approximately 1–20% of the sequences were unidentified even to a phylum level with the BLAST-based classifications employed, with most unclassified fungi observed in the non-AMD site rhizosphere (FL.E, 20%) and AMD site root endosphere (Lan E, 18.5%). However, subtle variations in classified fungal taxa within and between sampling sites were discernible. Florida Lake site was characterized by a higher abundance of phylum \textit{Basidiomycota}, accounting for 40% and 90% of taxa in the rhizosphere and root endosphere, respectively. In contrast, \textit{Ascomycota} constituted the major phylum in AMD sites, accounting for ~90% and 70% abundance of taxa in the rhizosphere and root endosphere, respectively. The other minor phyla (<1% relative abundances) identified included \textit{Entorrhizohyphomycota}, \textit{Gomerochomycota}, \textit{Monoblepharomyctota}, \textit{Macromycotota}, and \textit{Rozellomyctota}. At the class level, \textit{Eurotiomycetes}, \textit{Saccharomycetales}, \textit{Tremellomycetes}, \textit{Agraricomycetes}, \textit{Microbotryomycetes}, \textit{Ustlaginomycetes}, \textit{Dothideomycetes}, \textit{Sordariomycetes}, \textit{Leotiomycetes}, and \textit{Cystobasidimycetes} were the abundant classes with variation in their abundance across the sampling sites (Figure 3b).

The relative abundance of the fungal communities at the genera level (Figure 3c) also provided more insight into the taxonomic differences between the rhizosphere and root endosphere of \textit{P. australis} in AMD and the control site. In the rhizosphere samples, the two AMD sites clustered together separating them from the genera found in the non-AMD site. The most abundant genera occurring in the two sites included \textit{Aspergillus}, \textit{Penicillium}, \textit{Candida} and unclassified \textit{Saccharomycetales}. In the root

Table 1. Summary of the mean (±SD) physicochemical variables water and heavy metal content of the rhizospheric soil of \textit{P. australis} growing in two acid mine sites (Lan 3 and Wuinze 17) and non-acid mine site (Florida Lake) (n = 9).

| Variables      | Lan 3          | Wuinze 17       | Florida Lake  |
|----------------|----------------|-----------------|---------------|
| EC (μS/cm)     | 5537 ± 2.7a   | 2180 ± 1.2a     | 113 ± 0.1a    |
| TDS (mg/L)     | 4305 ± 1.6a   | 1296 ± 0.8a     | 102 ± 0.4a    |
| Salinity (g/kg) | 3.20 ± 0.3a   | 1.26 ± 0.1b     | 0.10 ± 0.002c |
| pH             | 5.01 ± 0.27c  | 6.50 ± 0.8b     | 7.81 ± 2.2a   |
| Fe (mg/L)      | 193.6 ± 7.89a | 54.21 ± 6.69a   | 25.24 ± 1.64a |
| Cd (mg/L)      | 0.02 ± 0.001a | 0.02 ± 0.003c   | 0.02 ± 0.001a |
| Cr (mg/L)      | 0.17 ± 0.02a  | 0.09 ± 0.01a    | 0.03 ± 0.003c |
| Co (mg/L)      | 0.39 ± 0.09a  | 0.05 ± 0.004a   | 0.01 ± 0.006c |
| Cu (mg/L)      | 0.27 ± 0.05a  | 0.13 ± 0.002c   | 0.03 ± 0.002c |
| Na (mg/L)      | 1.29 ± 0.10a  | 0.40 ± 0.03c    | 0.26 ± 0.02c  |
| Pb (mg/L)      | 0.10 ± 0.02a  | 0.04 ± 0.005c   | 0.06 ± 0.005c |
| Zn (mg/L)      | 0.50 ± 0.05a  | 0.21 ± 0.02c    | 0.11 ± 0.02c  |

Means with similar small letters (a-c) within a row are not significantly different (\( p < 0.05 \)). EC-electrical conductivity, TDS-total dissolved solute.
endosphere, despite no clear clustering between the AMD and non-AMD sites, the most abundant taxa were genus Vishniacozyma, Aspergillus, Penicillium, unclassified Capnodiales, and Saccharomycetales. Core microbiome analysis demonstrated that unique OTUs differed within the sites and between rhizosphere and endosphere (Supplementary Information Table 1). In root endosphere, non-AMD site (FL) had highest number of unique taxa (18 OTUs) while AMD sites had 9 and 1 OTU for Lan3 and Wuinze17_E, respectively. In contrast, 17, 5 and 7 unique OTUs was observed in the rhizospheric samples of FL, Wuinze17 and Lan3, respectively.

3.3. Metabolome profiling

A total of 73 metabolites were annotated based on matching retention time and mass spectra to the in-built LabSolutions Insight® platform database. The metabolites included amino acids (22), nucleotide/nucleosides (17), carbohydrates (9), organic acids (7), vitamins (2), and other known plant metabolites (16) (Supplementary Information Table 2). Figure 4 presents the heatmap showing both the hierarchical clustering analysis (HCA) and relative spectral abundance of the 73 metabolites detected in both rhizosphere and endosphere compartments of *Phragmites australis* roots impacted by different AMD environment. The presented dendrogram indicates sub-clusters containing different primary metabolites with varying degrees of similarities.

Overall, metabolites of the rhizosphere from the two AMD sites (Lan 3 and Wuinze 17) clustered together indicating the closeness of the metabolite profiles in the two sites, but with subtle variation in their spectral abundances (Figure 4). Interestingly, choline was relatively high in all three rhizosphere soil samples. In the root samples, however, clustering was not observed between the two AMD sites but only between an acid mine site (Lan 3) and a control site (Florida Lake).

Beta diversity statistical analyses were also done to determine the statistical significance of spatial heterogeneity in the relative spectral abundance of metabolites. The NMDS plots revealed that samples clustered more by type (whether rhizosphere or root endosphere) (ANOSIM, *R* = 0.8567, *p* < 0.001) rather than by AMD sites (ANOSIM, *R* = -0.1381, *p* = 0.9890) (Figure 5). The stress of the NMDS ordination was 0.103. Similarly, goodness of fit analysis also revealed that localization (rhizosphere vs root endosphere) (*r*² = 0.712, *p* = 0.006) had a significant correlation with metabolites composition compared to habitat (AMD sites) that had a poor and non-significant correlation (*r*² = 0.149, *p* = 0.934). However, closer inspection of the data showed that variability in primary metabolites was more significant in the rhizosphere than the root endosphere, with rhizospheric samples of AMD sites (Wuinze17 and Lan3) clustering closer together than non-AMD site (Florida Lake) (Figure 5). The study also investigated the effects of physiochemical parameters on metabolites composition and abundance in the rhizospheric samples. To achieve this, the direction and length of the vectors influencing the metabolite variability. Vector-fitting of the significant environmental variables to the NMDS ordination space as defined by the metabolite composition is illustrated in Figure 5.

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**Table 2.** Summary of sequencing outputs and diversity indices for rhizosphere and root endosphere fungal communities associated with the roots of *Phragmites australis* growing in two acid mine sites (Lan 3 and Wuinze 17) and non-acid mine site (Florida Lake-FL).

| Indices          | Rhizosphere | Root endosphere |
|------------------|-------------|-----------------|
|                  | Lan3_E      | Wuinze17_E      | FL_E | Lan3_A | Wuinze17_A | FL_A |
| Observed OTU     | 90          | 110             | 122  | 114    | 119        | 154  |
| Quality reads    | 58 217      | 106 724         | 25 248| 96 400 | 46 416     | 49 386|
| Simpson_1-D      | 0.181       | 0.135           | 0.080| 0.112  | 0.135      | 0.099|
| Shannon_H        | 2.292       | 2.538           | 3.059| 2.641  | 2.753      | 2.756|
| Evenness_eH/S    | 0.110       | 0.115           | 0.175| 0.123  | 0.132      | 0.102|
| Fisher_alpha     | 10.43       | 12.11           | 16.66| 12.77  | 14.78      | 19.67|
| Chao-1           | 92.63       | 113.5           | 132.9| 116.1  | 123.1      | 156.2|
| Good's coverage (%) | 99.5   | 99.9            | 98.9 | 99.9   | 99.3       | 99.8 |

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**Figure 2.** (a) Rarefaction analysis showing the sequence coverage. (b) Venn diagram showing number of OTUs at each location and the shared fraction of core fungal microbiome associated with rhizosphere and root endosphere of *Phragmites australis* growing under AMD environments.
To identify differentially abundant metabolites between different groups (rhizosphere vs root endosphere), a metabolite-group association analysis using the multipatt function of the “indicspecies” R package [37] was performed. As shown in Table 3, nine metabolites (adenosine, acetylcholine, symmetric dimethylarginine, asymmetric dimethylarginine, dopa, guanosine monophosphate, kynurenine, isoleucine, and leucine) were significantly \((p < 0.05)\) associated with the root endosphere. In contrast, niacinamide, creatinine, ophthalmic acid, 2-ketogluutaric acid, serotonin, histamine, guanosine, methionine, dimethylglycine, thymidine monophosphate, thymine, choline, cystathionine, proline, and valine were significantly associated with the rhizosphere \((p < 0.05)\).

3.4. Relationship between fungal community, primary metabolites, and physicochemical parameters

To examine which of the physicochemical parameters and primary metabolomes in both root endosphere and rhizosphere were most likely associated with variation of the fungal community, the study explored their relationship in a multivariate canonical correspondence analysis (CCA) (Figure 6). The CCA triplot showed that the concentration of metabolites (dimethylglycine, ophthalmic acid, histamine, guanosine, methionine, dimethylglycine, thymidine monophosphate, thymine, choline, cystathionine, proline, and valine) were significantly associated with the rhizosphere \((p < 0.05)\). In Florida Lake’s rhizosphere sample, pH was the major physicochemical parameter that influenced the fungal community and it showed a strong positive correlation with fungi belonging to the genera Ascomycota_uc, Capnodiales, Cystofilobasidiales, and Sporidiobolales. In contrast, pH negatively correlated \((p < 0.01)\) to rhizosphere microbiome diversity in Lan_3. The result showed that the AMD sites induced the production of more primary metabolites by the reed but selected for fewer associated fungi.

Inferences from the CCA analysis also showed that a correlation exists between the fungal community in the root endosphere of \(P. australis\) and the primary metabolites (Figure 6b). The fungal community at the order level and primary metabolites showed close ordination indicating the impact of AMD pollution on the metabolites produced by the root tissues and the diversity of endophytic fungi. Fungal taxa such as Ascomycota_uc, Capnodiales, and Saccharomycetales showed a strong positive correlation with kynurenine, acetylcholine, and dopa in Lan_3, whereas Eurotiales, Hypocreales, Ustilaginales, and Dothideales positively correlated with adenosine, asymmetric dimethylarginine, symmetrical dimethylarginine, and leucine in Wuinze 17. In contrast, orders belonging to phylum Basidiomycota such as Filobasidiales, Sporidiobolales, Holtermanniales, and Tremellales in Florida Lake showed a strong positive correlation with guanosine monophosphate and isoleucine.

4. Discussion

4.1. Rhizosphere and endosphere fungal community of \(P. australis\) under AMD conditions

This study describes the diversity of rhizospheric fungal communities associated with AMD heavy metal pollution gradient and plant root exudate profile. Overall, the results showed that the diversity and richness of the fungal community in both the root endosphere and rhizosphere in \(P. australis\) growing under the non-AMD control site were higher than in the
The rhizoplane and the endosphere are colonized by a subset of the microbial community has been postulated to occur in a two-step fashion. Generally, the assembly of the root microbial communities in soils, leading to reduced microbial diversity, richness, and metabolic activity [38, 39, 40]. However, soil fungal communities are less affected by heavy metals compared to soil bacterial communities [39, 41].

Interestingly, fungal diversity was generally higher in the root endosphere than the rhizospheric samples (Table 2). Unlike our study, previous reports reported in the rhizosphere than the endophytic compartments of poplar trees [44], and a subtropical island shrub Mussaenda kwangtungensis [45]. In these studies, it is postulated that the rhizosphere, generally characterized by presence of root exudates, mucilage produced by the root caps, and the release of sloughed-off root cells, is an active transition zone that provides suitable ecological niches for the growth, development, and reproduction of microbial communities [46]. On the other hand, environmental filters related to requirement for expression of genes associated with the production of cell-wall-degrading enzymes and resistance to a range of plant innate immune responses for successful endophytic compartment colonization, may account for the observed lower microbial diversity [44, 45, 47]. Collectively, the reported higher fungal diversity and richness of the aforementioned taxa may indicate that the local P. australis rhizospheric microenvironment has a complex effect on the fungal community, selecting for more endophytic metal tolerant groups under AMD-stress conditions. However, whether these observations could also be attributed to the inherent experimental challenge of low sample size used, varying copy numbers for the tandemly repeated nuclear ribosomal operon [48], PCR biases [49], and efficiency differences of DNA extraction from endophytic and rhizospheric compartment samples [50], also warrants further investigation.

In this study, Ascomycota and Basidiomycota were the dominant phyla (>1% of high-quality sequences) across all AMD samples. Congruent to our findings, Baker et al. [51] also reported that members of the phylum Ascomycota and Basidiomycota primarily dominating the sub-surface low-pH biofilms in Richmond mine (Iron Mountain). At the class level, Saccharomycetales, Eurotiomycetes, Tremellomycetes, Agraricomycetes, Microbotryomycetes, Cystobasidiomycetes, Ascomycota, Basidiomycota, Eurotiomycetes, Tremellomycetes, Agraricomycetes, Microbotryomycetes, Ustilaginomycetes, Dothideomycetes, Sordariomycetes, Leotiomycetes, and Cystobasidiomycetes were the dominant groups. Interestingly, both rhizosphere and root endosphere in two AMD sites had significantly higher levels of phylum Ascomycota compared to non-AMD samples. As co-adaptation between plants and its rhizospheric soil microbiota is essential to cope with edaphic stresses under extreme environment, the observed higher abundance of members of phylum Ascomycota belonging to class Dothideomycetes and Eurotiomycetes imply they could be playing important roles in the ecology and stability of
P. australis under such extreme environment. Several studies have reported that filamentous fungal genera such as Aspergillus, Mucor, and Trichoderma [52, 53, 54], Rhodotorula [55], Rhizopus [56], and Penicillium [57] are adaptable to environmental heavy metal contamination, detoxify them through inherent mechanisms such as transformation, crystallization, extracellular precipitation, complexation, and cellular absorption [57, 58, 59]. In this study, members of genera Penicillium, Candida, Saccharomyces, Vishniacosyagma, Trichoderma, Cladosporium, and unclassified Didymellaceae were highly enriched in the root endosphere and rhizospheric samples from AMD sites compared to the non-AMD site (Figure 3c). Yamaji et al. [60] also reported the tolerance of Clethra barbinervis to heavy metal as impacted by root fungal endophytes. Like ectomycorrhizal fungi, metal resistant root fungal endophytes utilize the enhancement of the antioxidative system, mediating heavy-metal distribution in plant cells and detoxification of heavy metal to induce tolerance of plants to heavy-metal stress [61]. Further, endo- and ectomycorrhizal fungi produce siderophores and enzymes important in the reduction of plant ethylene levels and thereby enhancing plant growth [60]. Thus, root fungal endophytes and ectophytes (Ascomycota and Basidiomycota) reported collectively may be contributing to the many

| Metabolites | Type | Endosphere | Rhizosphere | p-value |
|------------|------|------------|-------------|---------|
| Adenosine  | Nucleotide | 0.862 | - | 0.0001 |
| Acetylcholine | Signal molecule | 0.665 | - | 0.0094 |
| Symmetric dimethylarginine | Amino acid | 0.636 | - | 0.0003 |
| Asymmetric dimethylarginine | Amino acid | 0.599 | - | 0.0015 |
| Dopa       | Other | 0.595 | - | 0.0061 |
| Guanosine monophosphate | Nucleotide | 0.522 | - | 0.0032 |
| Kynurenine | Others | 0.488 | - | 0.0451 |
| Isoleucine | Amino acid | 0.451 | - | 0.0006 |
| Leucine    | Amino acid | 0.381 | - | 0.0005 |
| Nicotinamide | Others | - | 0.823 | 0.0001 |
| Creatinine | Organic acid | - | - | 0.661 |
| Ophthalmic acid | Organic acid | - | - | 0.654 |
| 2-Ketoglutaric acid | Organic acid | - | - | 0.643 |
| Serotonin  | Others | - | - | 0.632 |
| Histamine  | Others | - | - | 0.611 |
| Guanosine  | Nucleotide | - | - | 0.608 |
| Methionine | Amino acid | - | - | 0.602 |
| Dimethylglycine | Amino acid | - | - | 0.573 |
| Thymidine monophosphate | Nucleotide | - | - | 0.570 |
| Thymine    | Nucleotide | - | - | 0.568 |
| Choline    | Others | - | - | 0.556 |
| Cystathionine | Amino acid | - | - | 0.533 |
| Proline    | Amino acid | - | - | 0.504 |
| Valine     | Amino acid | - | - | 0.435 |

* The generalized indicator value (IndVal.g) and corresponding p-values used to assess the predictive value of a metabolite for either rhizosphere or endosphere as implemented in indicpecies in R [37] is given.
critical ecological functions including metal tolerance and detoxification, carbon, and nitrogen cycling [11, 60], important for the fidelity and proliferation of the reed plant within the extreme AMD environment.

4.2. Spatially defined root metabolome in *P. australis* is dependent on AMD gradient

Under conditions of heavy metal stress, plants have evolved numerous defense mechanisms ranging from reduced metal uptake, sequestration of metal into vacuoles, binding to phytochelatins/metallothioneins, activation of various antioxidants, to build-up of large quantities and diverse primary and secondary metabolites [62, 63]. These studies further reported that Cd, Cu, Ni, and Zn preferentially in human physiological conditions. On the other hand, fungal communities have been reported to be less sensitive to soil pH variations [41], which is inconsistent with the current findings. However, low pH is generally associated with high bioavailability and toxicity of heavy metals [82, 83]. This may explain the lower fungal diversity observed under high AMD pollution conditions. Under mild AMD conditions (site Wuinze 17), the impact of primary metabolites on rhizospheric fungal diversity was more discernible, with dimethylglycine, ophthalmic acid, histamine, serotonin, 2-ketoglutaric acid, thymidine monophosphate, proline, and choline levels being significantly associated with members of the order *Agaricales* and *Saccharomycetales*. NMDS analysis also showed higher variability between such metabolites in the rhizosphere of the *P. australis* reed than the root tissues under AMD-polluted. This may imply that the production of metabolites and the fungal communities present in the acid mine drainage sites at the rhizospheric level are more environmentally dependent than in the root tissues. It is worth noting that rhizosphere microbiota may also play significant role in the differential root exudation profile, in addition to producing some of these metabolites on their own in order to increase their fidelity in the rhizospheric environment. For example, the role of microbiome-reprogrammed systemic root exudation in promoting soil conditioning has been reported in tomatoes [84]. In our study, we focused only on primary metabolome, which were all assumed to be of plant origin. Since, plant root exudation is a dynamic process, involving diverse primary and secondary metabolites likely dependent on a plethora or transporters that are mostly uncharacterized [11, 85], result of this study has only scratched the surface of *P. australis* metabolic diversity and rhizomicrobiome assembly. It is anticipated that further studies will continue to advance the mechanistic understanding of plant-microbe-heavy metal interactions in this important pseudometallophyte.

5. Conclusions

Overall, the results of this study indicate that spatially defined exudation as modulated by abiotic factors plays a key role in the rhizobionome assembly with the establishment of distinct fungal communities that were observed to be associated with specific root parts of *P. australis*. It is plausible that these interactions (between root exudates and fungal community in the rhizosphere) are a critical component of *P. australis* growth and adaptability to the heavy metal-laden and low pH AMD ecological niche. While the fungal community may be important in the metal mobilization, transformation, and detoxification leading to the plant heavy metal tolerance and remediation, they are also greatly influenced by the plant root exudates, that enrich the rhizosphere in nutrients, including acting as messengers to initiate interaction between plant and the beneficial microorganisms. However, it is worth noting that the synergistic action of plant and microbes and their mechanism for metal phytorhizoremediation under the AMD environment may involve a complex interaction involving numerous cryptic crosstalk and regulatory networks that are still largely unknown. The success of microbial colonization of the rhizosphere is also dependent on several factors, such as...
chemotaxis, substrate specificity, competitiveness, and cooperativeness, whose understanding is currently limited. Further, plant root exudation is a dynamic process, involving diverse primary and secondary metabolites likely dependent on a plethora of transporters that are mostly a dynamic process, involving diverse primary and secondary metabolites likely dependent on a plethora of transporters that are mostly.

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