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Assessing the genetic diversity of Cu resistance in mine tailings through high-throughput recovery of full-length *copA* genes

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Characterizing the genetic diversity of microbial copper (Cu) resistance at the community level remains challenging, mainly due to the polymorphism of the core functional gene *copA*. In this study, a local BLASTN method using a *copA* database built in this study was developed to recover full-length putative *copA* sequences from an assembled tailings metagenome; these sequences were then screened for potentially functioning CopA using conserved metal-binding motifs, inferred by evolutionary trace analysis of CopA sequences from known Cu resistant microorganisms. In total, 99 putative *copA* sequences were recovered from the tailings metagenome, out of which 70 were found with high potential to be functioning in Cu resistance. Phylogenetic analysis of selected *copA* sequences detected in the tailings metagenome showed that topology of the *copA* phylogeny is largely congruent with that of the 16S-based phylogeny of the tailings microbial community obtained in our previous study, indicating that the development of *copA* diversity in the tailings might be mainly through vertical descent with few lateral gene transfer events. The method established here can be used to explore *copA* (and potentially other metal resistance genes) diversity in any metagenome and has the potential to exhaust the full-length gene sequences for downstream analyses.

Novel metal resistance genes in the environment, particularly mining-impacted soils, are valuable resources for industrial biomineralization¹ and soil remediation². *copA* is one of the core determinants for microbial resistance to Cu and its diversity has been examined in soils in a limited number of recent studies^{3–7}. These studies are all amplicon-based and rely on the availability of degenerate primers to target conservative regions of *copA*. Unfortunately, available *copA* sequences in the literature are thought to be highly polymorphic and therefore available primers only cover a subset of *copA* sharing high similarity. This hinders the assessment of *copA* diversity and the discovery of novel *copA* in the environment. To overcome this difficulty, we report a method to recover full-length *copA* from a tailings metagenome by combining the methods of metagenome assembly, local BLASTN and evolutionary trace (ET) analysis. The metagenomic strategy has been successfully applied to annotate Cu resistance genes in an activated sludge metagenome⁸, and ET analysis can be used to check the reliability of candidate *copA* genes detected in a metagenome by screening for key conserved domains of their CopA proteins.

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copA as a part of *cop* system and its homolog gene *pcoA* as a part of *pco* system were first identified in copper resistant strains of *Pseudomonas syringae* (PscopA) and *Escherichia coli* (EpcpA), respectively. PscopA and EpcpA were both plasmid-borne but many *copA* homologs were identified soon after, both chromosomal- (e.g. *Enterococcus hirae*) and plasmid-borne (e.g. *Xanthomonas* sp.). Interestingly, many chromosomal *copA* were found to be typical P-type ATPase⁹, while *pcoA* was then identified as multicopper oxidase¹⁰. Therefore, *copA* is thought to be highly polymorphic^{3,5} and the nature of *copA* has always been described as species-dependent in the literature^{11–14}. This caused confusion when studying *copA* diversity in the environment. It is known that multicopper oxidase and ATPase are different protein families using different energy sources^{15,16}. We thus hypothesize that all *copA* named based on sequence similarity in the literature are not homologs and can be divided into two groups encoding for multicopper oxidase and P-type ATPase, which are both highly conserved.

ET analysis is a method to extract functionally important residues from sequence conservation patterns in homologs, with the assumption that active site residues of a protein family are more conserved during its evolutionary history¹⁷. While complete or partial crystal structure of some CopA proteins has been available in the literature^{18–21}, it is feasible now to identify the common active sites which may be highly conserved among CopA proteins. Therefore, ET analysis is possible to reveal the underlying root of the polymorphism of *copA*/CopA.

A metagenome obtained by high-throughput sequencing of environmental DNA conceptually provides all the gene resources in a given environment²². Metagenomic sequencing successfully overcomes the difficulties facing traditional isolation-based and conventional amplicon-based molecular methods in recovering genetic information from the environment. By means of bioinformatic tools (e.g. MIRA)²³ which assemble the short reads in metagenomes into longer contigs or even nearly complete genomes, metagenomics allow us to explore a functional gene of interest in a high-throughput fashion²⁴. BLAST is the most commonly used tool to find homologous sequences based on sequence similarity for functional attributes²⁵. By building a local database of antibiotic resistance genes (ARGs), a local BLASTN procedure has been established to screen for the presence of ARGs in bacterial genomes²⁶; yet this local-BLASTN method has not been applied to metal resistance genes or to explore metagenomes.

Therefore, this study aimed to develop a method for the recovery of full-length *copA* sequences in a tailings metagenome. A metagenomic library was obtained by assembling 7 individual tailings metagenomes using MIRA. The library was searched using the embedded BLASTN method in BioEdit against a local *copA* database built in this study, followed by recovering full-length *copA* by annotating the contigs containing putative *copA*. All identified putative *copA* were then examined for the presence of highly conserved metal-binding motifs found by ET analysis.

Results and Discussion

In generating the metagenomes, the MiSeq sequencing yielded >3.9 billion bp and >14.8 million reads after quality control for the 7 tailings samples. The 7 individual tailings metagenomes were pooled together for high-quality assembly and subsequent *copA* gene recalling. In total, 8,566,357 reads (greater than 300 bp) were used for metagenomic assembly which generated 82,334 contigs with an N_{50} of 1,700 bp and a longest contig of 123,516 bp. The data amount used for assembly in this study is much higher than earlier studies^{27–29} and comparable to recent studies^{30–32} where the characterized soil microbial communities have much higher complexity. This allows a deeper coverage of the functional gene diversity in the tailings microbial communities. Moreover, though metagenomes of microbial communities in acid mine drainage have been well studied^{33,34}, this is, to our knowledge, the first report on microbial metagenomes from neutral mine tailings which more closely resemble natural soils than acidic tailings. While natural soil always harbors a microbial community with a complexity whose fine decoding is still beyond the capacity of current bioinformatics tools³⁵, the tailings microbial metagenomes of lower diversity and community structure complexity characterized here may be a useful proxy for studying soil ecosystem functioning.

The Mt Isa Cu-Pb-Zn tailings are neutral and saline ($EC > 2$ mS/cm; in the dry season it is much higher as found in our previous study³⁶) substrates and contain an average Cu concentration more than 16-fold greater than the background soil values³⁷ (Table 1). In addition to the high levels of salinity and Cu, the tailings are also high in total Pb and Zn concentrations. The saline and metal stresses may have exerted a strong selective pressure on the microorganisms within the system, as indicated by the extremely low microbial biomass (Table 1) and the microbial diversity, which is dominated by either thermophiles or halophiles as found in our previous studies^{38,39}. The dominance of extremophiles may explain the high GC contents (all > 60%) of the metagenome libraries⁴⁰.

The strong selective effects of metal stresses were also reflected by the significantly high abundance of resistance genes for heavy metals in the metagenomes, based on the annotations by MG-RAST pipeline. In comparison with a local soil close to the tailings site sampled, the tailings metagenomes contain average *cop*, *czc* (coding for multiple metal resistance) and *ars* (coding for arsenic resistance) gene abundances of 2.8, 2.5 and 1.7 folds, respectively, of those of the soil (unpublished results; the metagenome data will be released soon in MG-RAST). The heavy metals in the soil are close to background crustal values (Fig. 1). Increased levels of toxins can lead to the enrichment of relevant resistance genes, as reported for both antibiotic and heavy metal resistance genes^{41–45}. Enrichment of *copA* has also been reported in various metal-contaminated environments such as paddy soil³, arable soil¹⁴ and sediment⁴⁶.

Sample	pH	EC ($\mu\text{S}/\text{cm}$)	CEC (cmol/kg)	TOC (g/kg)	MBC (mg/kg)	Total Cu (mg/kg)	Total Pb (mg/kg)	Total Zn (mg/kg)	Raw metagenome data size (Mb)	Mean GC content after quality control (%)	Total gene copy annotated	Number of contigs after assembly	Maximum contig size (bp)	N50 of the contigs
Tailings_1	6.70	5,490	30.47	5.55	132.35	1316	5080	3914	706	63 \pm 9	2,153,536	82,334	123,516	1,734
Tailings_2	6.80	5,420	48.30	5.85	92.65	1294	5300	3234	707	62 \pm 10	1,046,050			
Tailings_3	6.90	3,860	39.04	5.54	73.64	1285	4700	2620	551	60 \pm 11	1,136,742			
Tailings_4	6.90	2,080	42.06	4.95	51.38	1247	5090	3164	551	64 \pm 7	1,044,817			
Tailings_5	6.70	3,120	15.63	4.43	38.49	1205	6350	2991	544	62 \pm 9	1,115,745			
Tailings_6	6.80	3,040	12.19	3.77	28.04	1056	5860	3261	542	62 \pm 9	1,265,601			
Tailings_7	6.80	3,390	20.51	3.54	27.78	1143	6140	3485	611	62 \pm 9	1,130,746			

Table 1. General descriptors of the tailings samples and the metagenomes analysed in this study. Notes: EC, electrical conductivity; CEC, cation exchange capacity; TOC, total organic carbon; MBC, microbial biomass carbon.

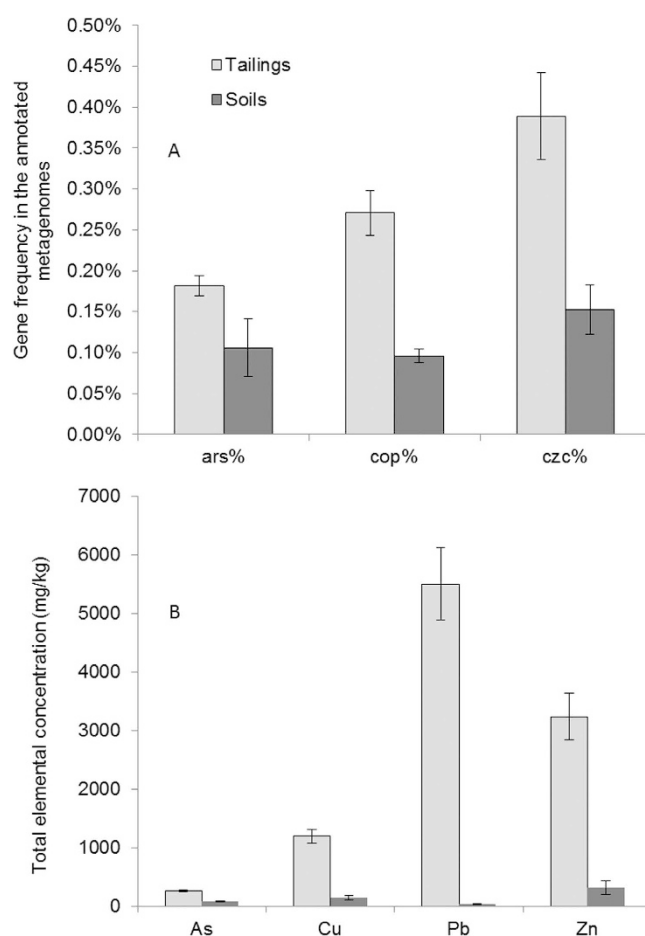


Figure 1. Total elemental concentrations of As, Cu, Pb and Zn in the tailings samples ($n=7$) and a reference soil ($n=3$) used in this study (**B**) and the corresponding abundances of resistance genes for these metals in the metagenomes from the tailings/soil samples (**A**). *ars*, resistance genes for As; *cop*, resistance genes for Cu; *czc*, resistance genes for multi-metals (e.g., Co, Pb, Zn).

However, to our knowledge, statistical results from shotgun metagenomic sequencing for *cop* genes have not been reported until recently⁸. Considering the polymorphism of *cop* systems, the results in this study may cover a deeper diversity of Cu resistance genes than those which are amplicon-based and rely on the specificity of primers.

A *copA* dataset containing 122 sequences annotated as *copA* obtained from full genomes available in Genbank has been built in this study. The abundance of *copA* selected from each phylum generally reflects the abundance of the available *copA* for that phylum in Genbank. Specification tests of the local BLASTN were done using *copA* sequences from the local database as well as full genomes containing putative *copA*. For the *copA* sequences from the database, the method was able to unambiguously identify all these genes with sensitivity and specificity at 100% and full coverage; at an elevated e-value, fragmented sequences of identity >80% can be found. At an e-value of 10^{-4} , we were also able to locate *copA*-like genes in the three genomes of *Acidithiobacillus ferrooxidans*, *Rubrobacter radiotolerans* and *Thioalkalivibrio sulfidophilus*, who are phylogenetically close to those of abundant species in the tailings, with more than 10 hits with an identity >80% and an alignment length >40 bp (the cutoff used for subsequent contig annotation in this study). Additionally, 6 hits were returned with an identity >86% and alignment length >40 bp when the BLASTN was applied to the genome of *Thermodesulfobacterium geofontis*. These results demonstrate a good coverage of *copA* diversity in the database and the tool established here is theoretically able to cover the novel *copA* diversity in our tailings metagenome.

In total, 99 full-length putative *copA* genes were recovered from the tailings metagenome. All of these genes had high similarity (>70%) to *copA* sequences from the genomes in Genbank. As the local BLASTN method is similarity-based, the number of *copA* genes identified can vary largely upon how novel the subjected microbial gene pool is and the diversity of the *copA* database. The threshold of alignment length in local BLASTN results was arbitrarily set as 40 bp to limit the number of contigs subject for subsequent *copA* gene annotation. However, the number of *copA* sequences recovered can be increased if the threshold is reduced, since Gupta *et al.*²⁶ found a minimum alignment length of 17 bp for putative new genes using a similar method for ARGs. Considering that the tailings microbial communities in this study are fairly low in species diversity, the local BLASTN method we have established has the potential to exhaust the *copA* diversity therein and can also be a novel tool to explore *copA* diversity in more complex soil environments.

Annotation of new genes is mostly sequence-similarity-based⁴⁷, including the early studies detecting *copA* homologs. While both multicopper oxidase-like and P-type ATPase-like *copA* have been referred to in the early literature^{9,48}, more recent studies state these as two “types” of *copA*¹⁴. However, we propose that the so-called two types of *copA* are not homologs and are incorrectly associated due to the limited number of available sequences used for similarity comparison in early studies. A wide range of protein sequences was used for phylogenetic analysis in this study. These were sequences that were verified experimentally or annotated as CopA and CopA-like Cu translocating proteins obtained from reported Cu resistant microorganisms. The sequences clearly separated into two distinct groups, one containing typical multicopper oxidases CueO and the other containing the typical P-type ATPase CtpA and ZosA (Fig. 2; Table 2). The former group includes the protein products of *copA* genes detected in early studies from the plasmids of *P. syringae* and *E. coli.*, and the latter group includes P-type ATPase Cu translocating proteins from *A. fulgidus*, *E. hirae* and *L. pneumophila* whose crystal structure has been resolved. These two groups of CopA may have distinct roles in Cu resistance, as implied by the experimental assays *in vitro* of typical CopA. Members of multicopper oxidase group CopA have been reported to be able to oxidize substrates of laccase, like phenol^{10,49}, while members of ATPase CopA have been found to bind and transport Cu(I)^{19,50,51}. Meanwhile, in the *E. coli* genome, the model microorganism for *cop* system studies, both groups of *copA* are present and their roles are found to be different. Plasmid-borne *pcoA* functions synergistically with chromosomal *copA*, and the protein PcoA can oxidise Cu(I) carried by PcoC and substitute the role of chromosomal CueO^{10,52,53}. Our analysis here indicated that many model species for *copA* studies contain both groups of *copA*, such as *E. hirae* and *P. syringae* (Fig. 2). The different roles of the two groups of *copA* may also be implied by their different sequence and tertiary structure. Multicopper oxidase CopA has no crystal structure resolved so far but prediction by SWISS-MODEL⁵⁴ showed that EcPcoA and PsCopA have similar crystal structure with laccase, which is distinct from the transmembrane figuration of the ATPase CopA detected in *A. fulgidus* and *L. pneumophila*^{16,19,50,55}. ET analysis indicated that the two groups of CopA have different metal binding motifs (Supplementary Figure 1). The ATPase group typically uses Cys as a metal binding residue in the motifs of CXXC (with HXXH as a variant) and CXC (the typical transmembrane metal binding motif), while the multicopper group uses His for metal binding in the form of HXH which is highly conservative within the group. Taken together, we suggest that CopA homologous to PcoA should be named as PcoA which is a multicopper oxidase, to differentiate it from ATPase CopA. Consequently, the metal binding motifs detected here can be used to screen for *copA* candidates with a high potential of functioning in Cu resistance.

The 99 full-length putative *copA* sequences were translated and aligned with the known CopA used for ET analysis. The presence of metal binding motifs abovementioned was screened in the putative CopA; For ATPase like CopA, the ATP binding motifs (GDGIN) were also used for screening. In total, 70 putative CopA were thought to have a high chance of functioning in Cu resistance.

Phylogenetic analysis was performed on the CopA sequences affiliated with the dominant species detected in the tailings (Fig. 3). The topology of the tree is largely consistent with that of the 16S rRNA gene based phylogeny. This indicates that the *copA* gene diversity was mainly controlled by vertical descent in the history of tailings community evolution, at least among the dominant species. Similar conclusions have been drawn for *merA* genes responsible for mercury resistance². P-type ATPase metal

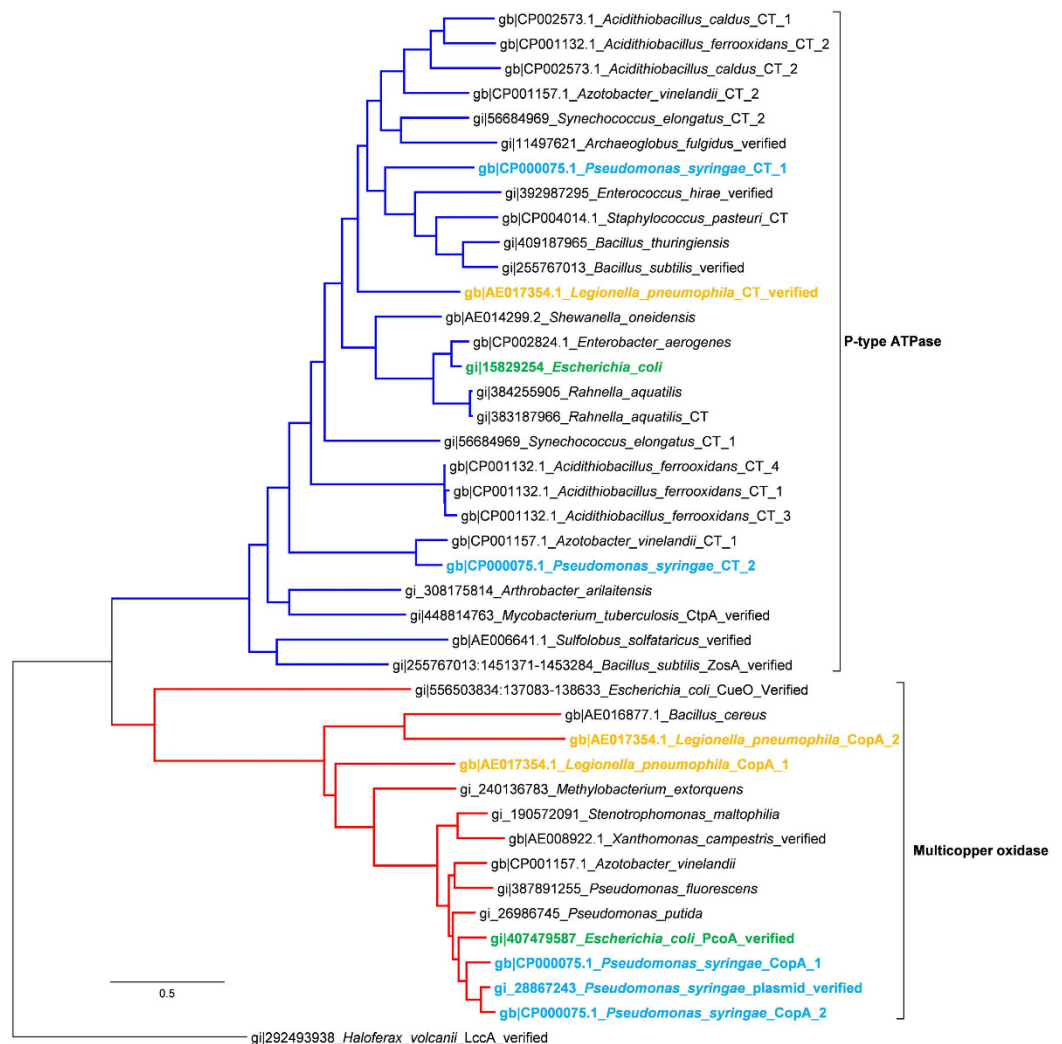


Figure 2. A phylogenetic tree based on the alignment of the CopA and reference protein sequences used for ET analysis in this study. Microbial species containing CopA sequences from both the P-type ATPase and Multicopper oxidase groups are highlighted in colour. When the Cu resistance function has been verified this is mentioned for those taxa.

homeostasis genes are probably ancient genes and have been essential for microbial survival, and thus lateral gene transfer plays a minor role in their evolution⁵⁶.

It is worth noting that the 29 putative *copA* sequences with no metal binding or phosphatase binding domains found may also function for Cu-resistance. For one thing, it is possible the combined presence of all the metal binding motifs found here are not essential for Cu resistance. Studies on EcCopA have found that the metal binding domain C⁴⁷⁹PC⁴⁸¹ motif is essential for Cu resistance but the other two C¹⁴XXC¹⁷ and C¹¹⁰XXC¹¹³ are not^{57,58}. Meanwhile, some Cu resistance ability may have evolved with sequences containing alternative metal binding domains. For example, in the alignment of ATPase group CopA, about half of the members lack the C⁸⁴XXC⁸⁷ motif; and CopA of *Sulfolobus solfataricus*, in which the Cu resistance function has been experimentally verified, uses YPC instead of CPC as a transmembrane metal binding domain (Supplementary Figure 1). Furthermore, it must be highlighted that the CopA proteins are relatively divergent in terms of both sequence composition and length. Inaccurate alignments can be made and lead to failure for detecting motifs when the sequences are too divergent⁵⁹. Therefore, the accuracy of alignment methods may largely determine the ability to detect the metal binding motifs of the subject sequences. For example, the alignment method used in this study failed to align CueO of *E. coli* with the others, and manual adjustments were required to locate the three experimentally-verified metal binding domains⁶⁰.

Species	MIC (mM) ^a
<i>Acidithiobacillus caldus</i> DSM 9239	24 ¹
<i>Acidithiobacillus ferrooxidans</i> isolate N39-30-03	800 ¹
<i>Agrobacterium tumefaciens</i> CCNWR533-2	>5 ⁷⁷
<i>Archaeoglobus fulgidus</i>	Protein functioning verified ^{b50}
<i>Arthrobacter arilaitensis</i> 11J	3.1 ⁷⁸
<i>Azotobacter vinelandii</i> GZC24	4.7 ⁷⁹
<i>Bacillus cereus</i> BC21	3.9 ⁷⁹
<i>Bacillus subtilis</i>	Protein functioning verified ⁸⁰
<i>Bacillus thuringiensis</i> N2	>5 ⁷⁷
<i>Cyanobacterium Synechococcus</i>	Unknown ^{c81,82}
<i>Enterobacter aerogenes</i> NTG-01	2 ⁸³
<i>Enterococcus hirae</i>	Protein functioning verified ¹⁷
<i>Escherichia coli</i> RJ92	400 ⁸⁴
	20 ⁸⁵
	Protein functioning verified ¹⁸
<i>Methylobacterium extorquens</i>	Unknown ⁸⁶
<i>Pseudomonas fluorescens</i> 09906	1.6 ⁸⁷
<i>Pseudomonas putida</i> CZ1	>5 ⁷⁷
<i>Pseudomonas putida</i> S4	>1 ⁸⁸
	1.0–3.2 ⁸⁹
	1.2–2.0 ⁹⁰
<i>Pseudomonas syringae</i>	Protein functioning verified
<i>Rahnella aquatilis</i> MT7	3.9 ⁷⁹
<i>Staphylococcus pasteurii</i> N2	>5 ⁷⁷
<i>Stenotrophomonas maltophilia</i> AAP56	>0.4 ⁹¹
<i>Synechococcus elongatus</i>	Unknown ⁹²
<i>Xanthomonas campestris</i> BrC2	0.3 ⁹³
	Protein functioning verified ⁴⁹
<i>Haloferax volcanii</i> LccA	Protein functioning verified ⁷²
<i>Escherichia coli</i> CueO	Protein functioning verified ⁷³
<i>Mycobacterium smegmatis</i> CtpA	Protein functioning verified ⁷⁴
<i>Bacillus subtilis</i> ZosA	Protein functioning verified ⁷⁵

Table 2. Selected microbial species used for evolutionary trace analysis in this study. The MIC of Cu, and whether the protein function has been determined is included. Genomes of these species all harbour *copA* or *copA*-like copper translocating genes. ^aminimum inhibitory concentration as of the strains of the species studied and under the specific test conditions in the corresponding references; ^bfunctions of *copA* genes have been experimentally verified or the crystal structure of CopA proteins has been resolved; ^creferred as Cu resistant but MIC is unknown or not provided.

Conclusions

A local BLASTN method was established in this study to recover full-length *copA* genes in an assembled tailings metagenome. The detected *copA* genes were further screened for potentially functioning *copA* by screening active sites in their encoding proteins, which were inferred by ET analysis using known CopA. The method established here can be used to recover full-length *copA* (and potentially other resistance genes) in any assembled metagenomes.

Materials and Methods

Sampling and DNA extraction. Tailings samples were sampled in June in 2013 from a field trial site located at Mount Isa (Mt Isa) tailings impoundment. Mt Isa (20.73°S, 139.5°E) is located in northwest Queensland, Australia. Mt Isa has a semi-arid climate with an annual pan evaporation of 2800 mm and an average rainfall of 400 mm, with the vast majority of the rain falling during the wet season (November to February). The tailings used for field trial were highly weathered and collected from a tailings storage facility (TD5) that contained mixed streams of Cu and Pb-Zn tailings and was decommissioned about

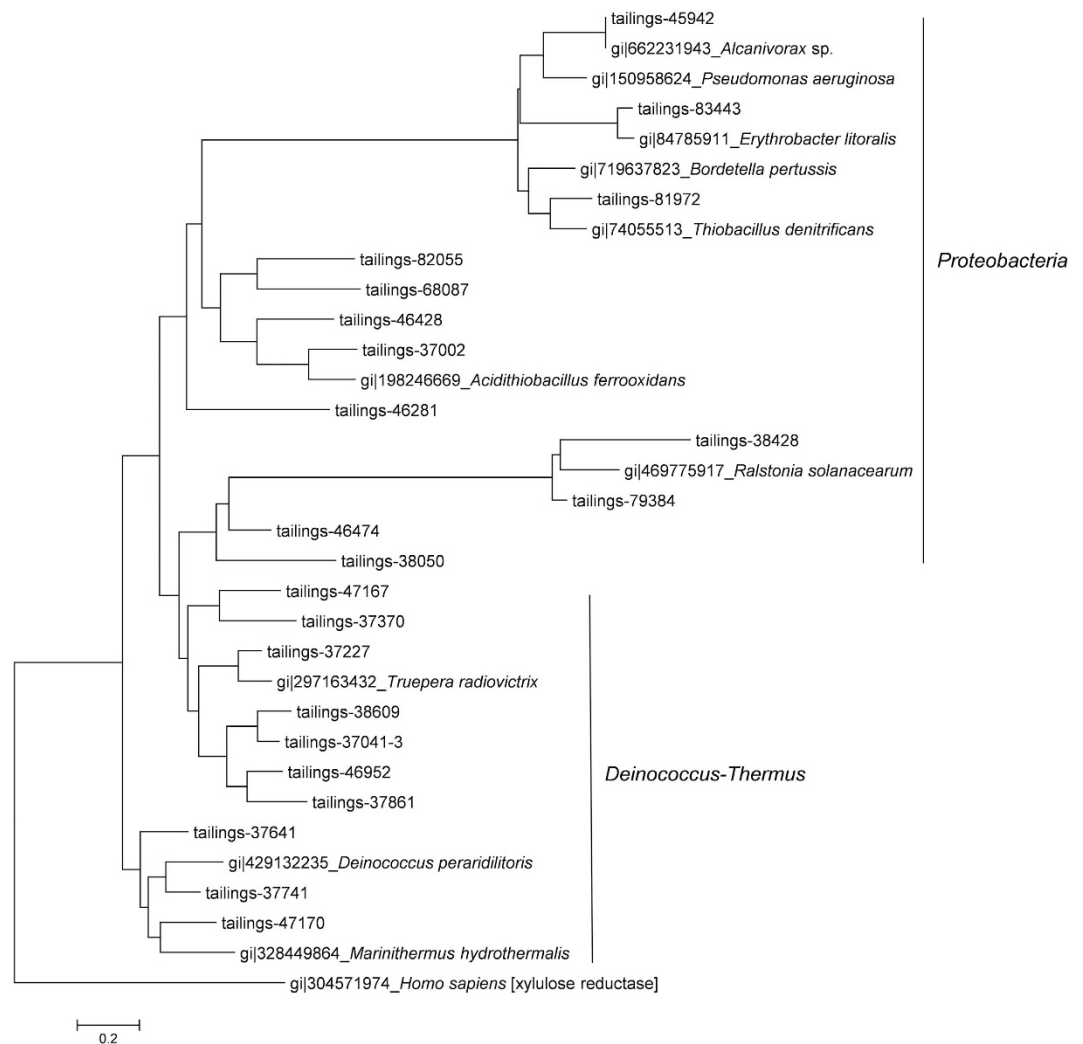


Figure 3. A phylogenetic tree showing the representative *copA* sequences affiliated with *Proteobacteria* and *Deinococcus-Thermus* recovered from the tailings metagenome after assembly.

40 years ago, which had received streams of Cu and Pb-Zn tailings for decades. The details of tailings properties and treatment setup can be found in our previous studies^{38,39}. Briefly, the tailings mainly consist of quartz, dolomite, pyrite, gypsum and kaolinite, and contain $0.13 \pm 0.03\%$ of total Cu, $0.18 \pm 0.02\%$ of total Pb, and $0.29 \pm 0.01\%$ of total Zn. A field revegetation trial was established in 2010, and within 3 years the woodchips amendment and/or revegetation treatments did not substantially change the tailings physiochemical properties and the dominant microbial species. Dominant microbial species were affiliated with *Rubrobacter* spp. of *Actinobacteria*, *Truepera* spp. of *Deinococcus-Thermus* and *Thioalkalivibrio* spp. and *Thiobacillus* spp. of *Proteobacteria*^{39,61}.

Tailings sample were stored in an iced container and shipped to the laboratory within 24 hours for DNA extraction. For physiochemical analyses part of the tailings were oven-dried at 40 °C, sieved through a 2 mm screen and mixed thoroughly before use. Methods for physiochemical analyses, such as electrical conductivity (EC), cation exchange capacity (CEC), total organic carbon (TOC), microbial biomass carbon (MBC) and total elemental concentrations (Table 1), can be found in our previous studies^{39,61}.

DNA was extracted from 24 tailings samples (8 plots with 3 replicates each plot). DNA extraction was done using commercial kits after cell enrichment by sucrose density centrifugation; detailed methods can be found in our previous studies^{39,61}. For MiSeq shotgun sequencing, replicates were combined to obtain enough DNA, and one sample of pure tailings failed the quality test for sequencing. Therefore, 7 independent tailings DNA samples representing the gene pool in the tailings landscape were sequenced using Illumina MiSeq platform in this study. The DNA concentrations and quality were measured using the Qubit® 2.0 Fluorometer (Thermo Fisher Scientific). DNA concentrations of the 7 samples ranged from 4.5 to 25.2 ng/μl, giving between 0.23 to 1.51 μg of DNA to use for the sequencing.

MiSeq sequencing. DNA sequencing libraries were prepared using the Illumina TruSeq DNA LT Sample Prep Kit (Illumina), following the standard manufacturer's protocol (Part #15026486 Rev. C July 2012) with modifications as described below. Genomic DNA was fragmented by sonication (Covaris S2) using the "Whole-genome Resequencing" settings in the protocol, after which the DNA was purified using AMPure XP beads. The purified, fragmented DNA was end repaired and again purified using AMPure XP beads. The purified, end repaired DNA was size-selected using a double-SPRI method to obtain insert sizes of approximately 600 bp (actual average insert size ranges from 456–667 bp). The size-selected DNA was A-tailed and then had adapters ligated, followed by an AMPure XP purification. This ligated product was amplified by PCR to produce the final library. The final individual libraries were visualized and quantified on the Agilent BioAnalyzer 2100 using the High Sensitivity DNA Kit. The libraries were then pooled in an equimolar ratio, and the pool was quantified by qPCR using the KAPA Illumina Library Quantification Kit (KAPA Biosystems). The library pool was sequenced on the Illumina MiSeq (MiSeq control software v2.0.5/Real Time Analysis 1.18), using the MiSeq Reagent Kit v3 (600-cycle) with paired-end 300 bp reads.

Shotgun metagenomic analyses. For functional annotations, the paired-end raw data was uploaded to MG-RAST server (Rapid Annotation using Subsystems Technologies for Metagenomes)⁶². Sequences were annotated to functional categories against M5NR database using BLASTX at an e-value cutoff of 1×10^{-5} . Results of general descriptors and annotations were downloaded from MR-RAST server for download analyses. Gene abundance was counted manually for *cop*, *czc* and *ars* genes, which are responsible for Cu, Pb/Zn and As resistance, respectively. For metagenomic assembly, the raw reads were trimmed using Skewer⁶³ to have a $Q > 25$ and a minimum length of 200 bp. Quality trimmed reads were used for *de-novo* assembly using the open source software package Mira (4.0rc5)²³ on a 24 core, 192 GB server with multithreading enabled. The seven tailings libraries were combined to improve the quality of assembly⁶⁴. The contig dataset was then subjected to local BLASTN for searching of *copA*-like genes.

Local BLASTN. A local BLASTN method, similar to ARGannot by Gupta *et al.*²⁶, was established to recall *copA*-like genes in the assembled metagenomic database. Briefly, nucleotide sequences annotated as *copA* in Genbank were retrieved manually and formed a local *copA* database (Supplementary Material 1). Then the assembled metagenomic dataset was aligned against the *copA* database using the embedded BLASTN method in BioEdit⁶⁵. Then the results were sorted based on aligned length and a threshold of 40 bp was used for screening *copA*-like genes in the metagenomic dataset. All the contigs containing candidate *copA* were picked out manually and then subjected to gene finding using Glimmer⁶⁶. For *copA*-like gene finding, the selected contigs were aligned against the Genbank database using BLASTN one-by-one and the sequence region annotated as *copA* or heavy metal resistance genes were compared with the open reader framework (ORF) found by Glimmer. The candidate ORF was then manually curated again against the Genbank database. The closest *copA* sequences were also retrieved for phylogenetic analysis. Maximum-likelihood phylogenetic trees were constructed for all the *copA* sequences in the database and selected found *copA* sequences from the tailings metagenome, using the method described below.

Local BLASTN against the *copA* database was done on a Window 7 computer equipped with dual-core 2.8 GHz CPU and 8 G RAM. The assembled dataset was cut into 6 sub-datasets and BLASTN for each dataset used up to 6 computing hours.

Specificity tests were done for the local BLASTN method using 1) 4 *copA* sequences included in the database (gi_294009986_ *Sphingobium japonicum*; gi_258541105_ *Acetobacter pasteurianus*; gi_347756788_ *Micavibrio aeruginosavorus*; gi_162145846_ *Gluconacetobacter diazotrophicus*), 2) three complete genomes (gi_198282148_ *A. ferrooxidans*, gi_627776062_ *R. radiotolerans*, and gi_220933193_ *T. sulfidophilus* who are phylogenetically close to those of abundant species in the tailings) containing *copA* novel but with homologs in the *copA* database and 3) a complete genome (gb_CP002829.1_ *T. geofontis*) belonging to phylum *Thermodesulfobacteria* which is not included in the database and probably containing novel *copA*.

Evolutionary trace analysis. Evolutionary trace was done following the method by Wilkins *et al.*⁶⁷. Since microorganisms harbouring *copA* homologs can also be Cu-sensitive⁶⁸, gene sequences for ET analysis in this study were selected based on three criteria (Table 2): 1) the microorganisms must be reported as Cu-resistant; 2) the full genome and/or complete sequence of plasmid(s) must be available in the Genbank; and 3) the gene must be annotated as *copA* or *copA*-like translocating genes in the full genome. In addition, five CopA protein sequences whose gene functions have been verified experimentally through mutagenesis or whose crystal structure has been resolved were also obtained from UniProt⁶⁹. Information on active sites of these verified CopA was gathered from the literature^{10,16–19,50}. Information on active sites of plasmid-encoded PsCopA was obtained from the predictions of UniPro since no crystal structure has been resolved so far for this group of CopA, and crystal structure of EcCueO and other multicopper oxidases were used as references. Furthermore, two multicopper oxidases, LccA (laccase) of *Haloferax volcanii* and CueO of *E. coli* for Cu homeostasis, and two P-type ATPases, CtpA of *Mycobacterium smegmatis* for cadmium transport and ZosA of *Bacillus subtilis* for Zn transport, 38 CopA sequences of known or highly possible to have Cu-resistance functions were aligned using the method of MUSCLE⁷⁰ embedded within MEGA 6⁷¹. The crystal structures of reference

proteins, LccA, CueO, CtpA and ZosA, have also been resolved^{72–75}. For active sites comparison, the Cu-binding motifs were searched manually based on the available information of verified proteins said above. For phylogenetic analysis, the alignment in FASTA format was used to determine the conserved regions using Gblocks 0.91 b online⁷⁶. Two highly conserved regions, one at the C-terminal and one at the N-terminal, were found, and the variable beginning and tail parts were trimmed in MEGA 6. The aligned regions of the alignment was then used for the construction of a Maximum-Likelihood tree using default values within MEGA.

Screening of CopA containing active sites found in ET analysis. All the CopA sequences detected from the tailings metagenome were aligned with the reference CopA (those with verified functions) using the alignment method of MUSCLE, as described above. The active site regions were checked manually based on the alignment for screening of CopA, gaps were manually adjusted to refine the alignment, and those containing the conserved motifs were determined as highly possible to be functioning as P-type ATPase Cu translocating proteins.

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Author Contributions

All authors commented on the manuscript and helped with interpretation of the data. X.L. and Y.Z. initiated the concepts. X.L. and L.H. designed the experiments. X.L. performed the experiment, did the downstream bioinformatics analyses and wrote the draft of the paper. L.H. did all of the sampling. T.B. and X.L. did the MiSeq metagenomics sequencing. B.S. processed the raw Illumina sequencing data.

Additional Information

Supplementary information accompanies this paper at <http://www.nature.com/srep>

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