Functional metagenomic characterization of antibiotic resistance genes in agricultural soils from China

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1. Introduction

Environmental contamination with antibiotics and antibiotic resistance genes (ARGs) is becoming a global health problem (Bush et al., 2011). The successful treatment for infections using antibiotics is being threatened by antibiotic resistance, which has developed from resistance to one class of antibiotics to increasingly prevalent multidrug resistance and extreme drug resistance. Antibiotic resistance has been one of the most important sustained driving forces for antibiotic discovery. Most of the research on the risk of ARGs to human health focused on the clinical setting, however, ARGs from pathogens comprise only a tiny fraction of total ARGs. We identified 45 clones conferring resistance to minocycline, tetracycline, streptomycin, gentamicin, kanamycin, amikacin, chloramphenicol and rifampicin. The similarity of identified ARGs with the closest protein in GenBank ranged from 26% to 92%, with more than 60% of identified ARGs had low similarity less than 60% at amino acid level. The identified ARGs include aminoglycoside acetyltransferase, aminoglycoside 6-adenyltransferase, ADP-ribosyl transferase, ribosome protection protein, transporters and other antibiotic resistant determinants. The identified ARGs from the soil with manure application account for approximately 70% of the total ARGs in this study, implying that manure amendment may increase the diversity of antibiotic resistance genes in soil bacteria. These results suggest that antibiotic resistance in soil remains unexplored and functional metagenomic approach is powerful in discovering novel ARGs and resistant mechanisms.

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A B S T R A C T
Soil has been regarded as a rich source of antibiotic resistance genes (ARGs) due to the complex microbial community and diverse antibiotic-producing microbes in soil, however, little is known about the ARGs in unculturable bacteria. To investigate the diversity and distribution of ARGs in soil and assess the impact of agricultural practice on the ARGs, we screened soil metagenomic library constructed using DNA from four different agricultural soil for ARGs. We identified 45 clones conferring resistance to minocycline, tetracycline, streptomycin, gentamicin, kanamycin, amikacin, chloramphenicol and rifampicin. The similarity of identified ARGs with the closest protein in GenBank ranged from 26% to 92%, with more than 60% of identified ARGs had low similarity less than 60% at amino acid level. The identified ARGs include aminoglycoside acetyltransferase, aminoglycoside 6-adenyltransferase, ADP-ribosyl transferase, ribosome protection protein, transporters and other antibiotic resistant determinants. The identified ARGs from the soil with manure application account for approximately 70% of the total ARGs in this study, implying that manure amendment may increase the diversity of antibiotic resistance genes in soil bacteria. These results suggest that antibiotic resistance in soil remains unexplored and functional metagenomic approach is powerful in discovering novel ARGs and resistant mechanisms.

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Numerous reports describe that agricultural practices, such as direct application of antibiotics or animal manure, provide positive selective pressure for antibiotic resistant bacteria, resulting in increase number and resistant level of these bacteria and ARGs in agricultural soils, and thereby expand the level of native resistance in soil (Knapp et al., 2010; Popowska et al., 2012). In addition, a considerable amount of resistant bacteria added into soil through manure amendment would lead to the spread of resistance to soil bacteria (Ghosh and LaPara, 2007; Heuer et al., 2011). However, there are only a few studies focusing on identification and characterization of ARGs in agricultural soil using functional metagenomics. Using this method, a total of 8 new ARGs were identified from a Spanish agricultural field, conferring resistance to ampicillin, gentamicin, chloramphenicol and trimethoprim (Torres-Cortes et al., 2011). From an apple orchard soil, 13 ARGs were identified including two putative novel bi-functional protein encoding genes (Donato et al., 2010).

Recent studies have been carried out to quantify the ARGs from soil, water and sediment samples in China. In a study conducted in Haihe River, Northern China, Luo et al. revealed that sulphonamide resistance genes, sul1 and sul2, were prevalent in this river, and that the gene abundance in sediments is 120–2000 times higher than in water, suggesting that sediments are an important reservoir of ARG in Haihe River (Luo et al., 2010). In a recent survey of tetracycline resistance genes in soils adjacent to representative swine feedlots in China, 15 tetracycline resistance (tet) genes were commonly detected in soil samples and the absolute number of tet gene copies was strongly correlated with the concentrations of tetracycline residues in these soil samples (Wu et al., 2010).

Despite the widely recognized potential health risks of over- or misuse of antibiotics in human and animal industries in China, information on abundance and diversity of ARGs is still scarce. Therefore the aim of this study is to investigate ARGs conferring resistance to one of thirteen antibiotics and discover novel ARGs in agricultural soils using functional metagenomic screening. By using three different type of soil, we also aim to assess the impact of agricultural practices, such as manure amendment and application of antibiotics, on the prevalence and diversity of ARGs in agricultural soils.

2. Material and methods

2.1. Soil samples

Four soil samples were collected from different agricultural soils, including 1) one from a field soil (DC soil) grown with Allium fistulosum L with a history of manure amendment located at Tongzhou, Beijing, Northern China. 2) Two from paddy fields in Tianjin (QG and WG soils), northern China, without application of manure and organic compost. 3) One from shrimp pond sediment (YZC soil, previously paddy soil) at Zhangzhou, Southern China. For each sample, three 500 g subsamples of soil were taken from 0–10 cm depth and were mixed in situ to form a combined sample. Soil samples were stored in icebox for transport to laboratory and approximately 100 g subsamples were stored at −80 °C for DNA extraction.

2.2. Metagenomic library construction

High molecular weight community DNA was extracted by the freeze-grinding, SDS-based methods (Zhou et al., 1996) and was purified using a low melting agarose gel followed by phenol extraction. DNA concentration and quality was determined with NanoDrop ND-1000. 5 μg purified DNA was partially digested with 10 U Sau3AI (Takara) in a 100 μL reaction at 37 °C for 2 h. Digested DNA with a size range of 1 kb–3 kb was selected by agarose gel electrophoresis and extracted from agarose gel using QiAquick Gel Extraction Kit (Qiagen 28704). Recovered DNA was ligated with BamHI digested pUC19 vector using the following protocol: 1 μL 10× Ligation buffer, 80 ng partially digested DNA, 20 ng BamHI cut pUC19 vector and 0.5 μL NEB T4 ligase (M0202) in a 10 μL ligation reaction, followed by incubation at 16 °C for 16 h and heat inactivation at 65 °C for 20 min. 2 μL fresh ligation product was transformed by electroporation into 50 μL prepared electro-competent Escherichia coli DH5α cells (Sambrook and Russell, 2000). After transformation using 2500 V for a 2 mm electroporation cuvette, cells were recovered with 1 mL SOC medium in a shaker at 37 °C, 200 rpm for 1 h. For each sample, recovered cells were combined as a library after all the ligation products were transformed.

Libraries were titered by plating 1 μL and 0.1 μL recovered cells onto LB agar plates containing 100 μg mL−1 ampicillin followed by incubation at 37 °C for 16 h. The LB agar plates were spread with 40 μL 2% X-gal and 7 μL 20% IPTG before titering. After incubation, white colonies were selected for amplifying the insert using M13 primers flanking the BamHI site of the pUC19 vector. For each library, average insert size was calculated by PCR amplifying 15 random chosen inserts using M13 primers and agarose gel electrophoresis. Total library size was determined by multiplying average insert size with the number of colony forming unit (cfu). The rest of recovered cells were grown in 30 mL LB broth containing 100 μg mL−1 ampicillin at 37 °C 200 rpm for 6 h, cells were collected by centrifugation at 5000 rpm for 15 min and re-suspended in 10 mL LB containing 100 μg mL−1 ampicillin, which were frozen in liquid nitrogen with 15% glycerol and stored in −80 °C.

2.3. Screening of antibiotic resistant clones

For each library, 2 mL frozen stock was grown in 20 mL LB with 100 μg mL−1 ampicillin at 37 °C for 6 h. 100 μL amplified library was spread on LB agar plates containing 100 μg mL−1 ampicillin and one of the antibiotics listed in Table S1 and incubated at 37 °C (16 h), 30 °C (2 d) and 25 °C (4 d), with 3 replicates for each temperature. Resistant clones were transferred to fresh LB agar plates containing 100 μg mL−1 ampicillin and relevant antibiotics of which resistance had been selected. Insert of each clone was amplified with M13 primers. Insert size was determined by agarose gel electrophoresis. Amplified inserts with identical size were digested with MspI. Inserts with different RFLP pattern and different insert size were picked and sequenced using Sanger sequencing at BGI Shenzhen (BGI, Shenzhen, China). Picked clones were grown in LB medium containing 100 μg mL−1 ampicillin and relevant antibiotic to verify resistance before sequencing.

2.4. Identification of antibiotic resistance genes

Open reading frames (ORFs) of full length inserts were identified using ORFfinder (http://www.ncbi.nlm.nih.gov/projects/gorf/). Identified ORFs were compared to a non-redundant protein database using blastp and non-redundant nucleotide database using tblastx. The closest hit for each ORF was collected with blastp and phylogenetic analysis for all genes was performed categorized by the antibiotic class resistant phenotype. The amino acid sequences were downloaded and aligned using ClustalW methods, phylogenetic trees were constructed by the neighbor-joining method and bootstrap analysis (1000 replicates) was performed with MEGA 5.05 software package (Tamura et al., 2011).

3. Results and discussion

3.1. Soil metagenomic library construction and antibiotic resistant clone isolation

Four metagenomic libraries were constructed using DNA extracted from different agricultural soils: one from a shrimp pond sediment (YZC library), one from a field soil grown with A. fistulosum L (DC library) and two from paddy fields in Tianjin (QG and WG library). Each library contained about 2 × 108 recombinant clones with average
insert size of 2 kb. Thus each library contained about 400 Mb DNA extracted from the soil, representing approximately 100 bacterial genome (assuming an average bacterial genome size of 4 Mb).

The constructed metagenomic libraries were subjected to antibiotic resistant clone isolation using each of 13 antibiotics listed in Table S1. A total of 45 unique clones were identified conferring resistance to the following antibiotics: tetracycline (TC) minocycline and tetracycline; aminoglycoside streptomycin, gentamicin, kanamycin and amikacin; chloramphenicol and rifampicin. The resistant phenotypes of selected clones were validated by checking their ability to grow on relevant antibiotics containing LB medium, followed by sequence analysis for the characterization of antibiotic resistant genes. The identified ARGs include aminoglycoside acetyltransferase, aminoglycoside 6-adenyltransferase, ADP-ribosyl transferase, ribosome protection protein, transporters and other antibiotic resistant determinants. It is likely that the constructed metagenomic libraries could contain much more ARGs than that identified in this work due to the limitation of heterologous expression in the E. coli host. Given the considerable biodiversity of soil microbial communities, expanding the metagenomic library size could yield tens or hundreds more ARGs.

For all the putative resistant genes, the similarity with the closest protein in GenBank ranged from 26% to 92%, with only a minority of 2% of identified resistant genes had high similarity (>90% at amino acid level) to previous known genes, and 67% of identified resistant genes had low similarity less than 60% at amino acid level, which may indicate the novelty of these genes (Fig. 1). Most of the ARGs failed to identify genes with significant matches when using nucleotide blast search of NCBI nr database, which suggest that functional screening could discover genes whose function might not be obvious from their sequence.

Thirty one clones were derived from DC library, representing about 69% of the total resistant clones (Fig. 2). The DC library was constructed using DNA from field soil grown with Allium fistulosum L. which was fertilized with pig manure. Antibiotics are commonly used in swine farm for disease treatment and growth promotion, which could lead to high concentration of antibiotics in pig manure (Qiao et al., 2012) and subsequently increase the antibiotic concentration in soil amended with pig manure. Our recent study revealed strikingly diverse and abundant ARGs in Chinese swine farms (Zhu et al., 2013), the results in this study provided another piece of evidence that antibiotic resistance in the environment could be affected directly by the release of antibiotic-resistant bacteria and antibiotic residues for pig manure.

3.2. Rifampicin resistance

Four clones were identified to be resistant to rifampicin, three from WG library and one from YZC library, the size of inserts was between 1 and 4 kb. Putative ORFs responsible for rifampicin resistant were identified, and the similarity of amino acid sequences of rifampicin resistance genes ranged from 35% to 75% to their most similar proteins (Table S2, Fig. S1). ORF WGRif3028-2 encoded a rifampin ADP-ribosyl transferase, suggesting that the conferred resistance is due to ribosylation of rifampicin. ORF WGRif3063-1 encoded a transporter-like protein. Although ORF WGRif3065 encoded a protein similar to a hypothetical protein, tblastx search database showed that it was related to a putative efflux RND transporter protein. Three ORFs were identified in clone YZCRif3706, including 1 closely related to DEAD/DEAH box helicase and 2 transporter protein homologues, sub-cloning of each ORF to an E. coli host or transposon mutagenesis should be carried out to test the ability of individual gene to confer resistance to rifampicin.

3.3. Chloramphenicol resistance

A total of 11 ORFs predicted to confer chloramphenicol resistance were identified from 11 recombinant clones (insert size 0.8–2 kb). Most of the chloramphenicol resistant clones were derived from the DC library, except one from the WG library. These ORFs shared an amino acid identity between 32% and 89% with their closest genes in GenBank (Table S3, Fig. S2). Annotation of chloramphenicol resistant genes revealed a number of mechanisms which enable them to mitigate the inhibitory effect of chloramphenicol. ORFs DCChl2502 and DCChl2509 encoded proteins belong to the multidrug ABC transporter or the RND family efflux transporter, sharing a similarity of 70% to 48% with their closest protein, respectively. Phylogenetic analysis of the amino acid sequence of DCChl2502 and DCChl2509 revealed very low similarity with any of the known chloramphenicol/ florfenicol exporter genes (Fig. 3). DCCh2503, DCCh2510, DCCh2512 and DCCh2516-2 encoded proteins are involved in ribosome protection, recycling and biogenesis, which could weaken the bacteriostatic effect of chloramphenicol, resulting in the resistance to chloramphenicol (Dittrich and Schrempf, 1992; Gramman et al., 2006; Iwanaga et al., 2004).

A serine/threonine-protein kinase like protein was encoded by DCCh3005 and shared a very low similarity (32%) with the closest protein at amino acid level. Over-expression of this gene may lead to increased generation of persistent cells which could survive antibiotic treatment probably by entering into a dormant state and exhibit multidrug resistance (Correia et al., 2006). WGChl2502-1 encoded a PhoH family protein, which is important to starvation- and stationary-phase-induced resistance to membrane-permeabilizing antimicrobial agents (McLeod and Spector, 1996). The function of DCCh2506 encoded ATP phosphoribosyltransferase like protein, DCCh2508 encoded allantoinase like protein and DCCh2515 encoded alpha-N-
arabinofuranosidase like protein conferring chloramphenicol resistance remains unknown. The most common mechanism of chloramphenicol resistance is chloramphenicol acetyltransferases (CATs) mediated by acetylation, resulting in the inactivation of chloramphenicol (Schwarz et al., 2004), however, resistance gene encoding CATs like protein were not detected in this study.

3.4. Aminoglycoside resistance

We identified 23 ARGs from 23 clones conferring resistance to aminoglycoside: 2 conferring resistance to amikacin, 2 to gentamycin, 1 to streptomycin and 18 to kanamycin, with amino acid similarity between 26% and 92% to their closest proteins in GenBank (Table S4). Most of the aminoglycoside resistance genes were derived from DC library, except 2 from QG library and WG library respectively. Enzymes responsible for inactivation of aminoglycosides were classified into three major classes according to the type modification: AACs (acetyltransferases), ANTs (nucleotidyltransferases or adenyltransferases), and APHs (phosphotransferases) (Ramirez and Tolmasky, 2010; Wright, 1999). Five AAC like genes (DCGen3701, DCKan30A20-1, DCKan30A30-2, WGKan37B65 and WGKan37B73-2) and one ANT like gene (QGStr37-1) were identified in this study (Table S4). Phylogenetic analysis of the amino acid sequences of identified genes with reported AACs and ANTs showed that DCGen3701, sharing a low similarity of 38% with the closest match in GenBank database, was related to an AAC(3) cluster proteins, while it was separated from the other twelve AAC(3) proteins, indicating that it could be a novel AAC(3) cluster gene. WGKan37B65, WGKan37B73-2 and DCKan30A30-2 fell within the cluster of AAC(6′) enzymes which are by far the most common (Ramirez and Tolmasky, 2010), while DCKan30A20-1 was distant from other AAC(6′) forming a separate branch in the AAC(6′) cluster, suggesting that it may be a new AAC(6′) gene. DCKan30A20-3 was found to be related to an AAC(2′) gene and form a separate cluster (Fig. 4). QGStr37-1 shared a high similarity of 70% at amino acid level with an aminoglycoside 6′-adenyltransferase from Sphingobacterium spiritivorum (Table S4), they group together and form a new cluster distinctive from ANTI(6)-la proteins (Fig. 5).

Other putative inactivation enzymes were also identified in this study, including 2 fumarylacetoacetate (FAD) hydrolase like proteins (DCAmi3001-1, DCKan30A25-2) and 1 HAD family hydrolase like protein (DCKan30A12). Interestingly, we found DCKan30A10, which encoded an erthyromycin esterase like protein, exhibited resistance to kanamycin, while further experiments showed that clone DCKan30A10 could not grow in erthyromycin containing LB medium.

Ribosome alteration including methylation of 16S rRNA, mutation of 16S rRNA or ribosomal proteins would lead to aminoglycoside resistance (Doi and Arakawa, 2007; Galimand et al., 2005). Methylase like protein encoding genes were also detected conferring resistance to kanamycin, including ORFs DCKan30A22-2, DCKan30A43-2 and QGKan37-1.

Reduced uptake, decreased permeability or export by active efflux pump is another major mechanisms involved in aminoglycoside resistance (Ramirez and Tolmasky, 2010). DCGen3002-1 encoded a protein similar to phosphomannomutase and DCKan30A29 encoded a protein similar to nucleotide sugar dehydrogenase, both of which are responsible for alginate polymerization and the resultant alginate layer would cause a mucoid phenotype and provide a protective barrier against host immune defenses and antibiotics (May et al., 1991; Naught et al., 2002). DCKan30A03 encoded 2-oxoglutarate dehydrogenase is involved in glutamine utilization and could reduce cell wall hydrophobicity, which has previously been reported to be important for multiple antibiotic resistance (Wolff et al., 2009). Efflux pump mediated resistance was found in ABC transporter like protein encoded DCKan30A31-2.

Other mechanisms for aminoglycoside resistance were also found in this study, such as attenuation of oxidative stress caused by aminoglycoside (DCAmi3003-2, DCKan30A27), regulation of marRAB multiple
antibiotic resistance operon by DCKan30A19-2 encoded transketolase-like protein (Domain et al., 2007) and DCKan30B13-2 encoded tRNA synthetase mediated resistance (Struble and Gill, 2009). Although previous reports showed that UvrA homolog confers resistance to nogalamycin and daunorubicin (Goosen and Moolenaar, 2008), the function of DCKan30B21 encoded UV damage endonuclease in the kanamycin resistance remains unknown. Three ORFs were identified in clone DCKan30A08 and were closely related to aminotransferase class-III, Isoprenylcysteine carboxyl methyltransferase and glyoxalase/bleomycin resistance protein/dioxygenase, respectively. The ORF in clone DCKan30A08 responsible for kanamycin resistance remained to be determined.

3.5. Tetracycline resistance

A total of seven clones were identified: five conferring resistance to tetracycline and two to minocycline. Two of the TC resistant clones were derived from DC library, one from QG library, and four from YZC library, respectively (Table S5). Efflux of TCs out of cells and ribosome protection are the most widespread mechanisms in TC resistance, and most of their genes are acquired via transferable plasmids and/or transposons (Thaker et al., 2010). TC efflux pumps like genes were identified in this study, they were DCTet3003-1, YZCMin2541 and YZCTet3735. Phylogenetic analysis indicated that DCTet3003-1 and YZCTet3735 were related to tetV, while YZCMin2541 was related to otrC (Fig. S3).

Tetracycline inactivation enzymes were relatively few with only three genes, tet(X), tet(34) and tet(37), were reported conferring tetracycline resistance due to alteration of tetracycline (Roberts, 2005). While, in this study, QGMin2501 was identified encoding an amidohydrolase like protein which could hydrolyse of methylamino of minocycline, resulting in inactivation of minocycline. The serine/threonine protein kinase mediated resistance, as mentioned before, was also observed in DCTet3002 (nucleotide sequence identical to DCChl3005), YZCTet3012 and YZCTet3064-1.

ARGs conferring resistance to beta-lactams (Cefotaxime), fluoroquinolone (Ciprofloxacin), macrolides (Erythromycin), trimethoprim and nitrofurantoin were not detected in this study, which may be due to the cloning bias and selective heterogeneous expression of environmental DNA in E. coli (Forns et al., 1997). Although functional screening of metagenomic libraries is a powerful approach in the study of ARGs, it has the disadvantage that the gene expression is necessary for the selection, cryptic genes or gene unable to be expressed in the recipient strain could be missed during the functional screening. Given that the quantity and diversity of microbes in soil, the constructed metagenomic library...
represents only a small fraction of microbial community in soil, and thus, the true size of the soil ARGs reservoir could be much larger (Monier et al., 2011). Some of the ARGs were closely linked with a number of widespread mobile genetic elements (MGEs), including plasmids, transposons and integrons, which were involved in horizontal transfer of ARGs among environmental bacteria (Fluit and Schmitz, 1999; Fricke et al., 2009; Li et al., 2012; Zhang et al., 2011). These ARGs could be missed when screening a microbial chromosomal DNA derived metagenomic library.

Our results showed that most of the identified ARGs had low similarity at amino acid level with the closest protein in GenBank, suggesting that soil ARGs might remain largely uncovered. Antibiotics in the natural environment exist at a concentration much lower than the therapeutic dose, and play multifaceted roles in nature, including inter-species competition, signal communication, host–parasite interaction, virulence modulation and quorum sensing (Sengupta et al., 2013). Therefore, unlike clinical environment, in which the main function of ARGs is to confer protection against the lethal concentrations of antibiotics, ARGs harbored in the natural environment also serve for response to antibiotic signaling (Aminov, 2009) and confer resistance not only to antibiotics but also to a number of structurally unrelated compounds (Sengupta et al., 2013). This could be one reason for the disparity between soil ARGs and clinical resistance. Future work should be carried out ascertain the effect and mechanisms of novel ARGs.

4. Conclusion

This study is the first effort to characterization antibiotic resistosome for multiple antibiotics from agricultural soil in China using functional metagenomic method. We highlight the advantage of functional metagenomic technology in discovering novel ARGs and resistant mechanisms rather than identifying genes on the basis of sequences. We identified a variety of ARGs conferring resistance to rifampicin, chloramphenicol, aminoglycoside and tetracycline, through almost all known mechanisms underlying relative antibiotic resistant phenotype. The novelty and diversity of identified ARGs suggest that antibiotic resistance in soil remains unexplored and could potentially threat human health if they transfer to pathogens. The identified novel ARGs and resistant mechanisms will extend our knowledge and understanding of the ecology of microbial antibiotic resistance in different environments, and will shed new insights on the emergence of ARGs in natural environment.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.envint.2013.12.010.

References


Fig. 5. Neighbor-joining tree of amino acid sequences of the ANTs (nucleotidyltransferases or adenyltransferases) like aminoglycoside resistant genes found in this study. The relative enzymes were presented with enzyme names, followed by genera, species and GenBank accession numbers. Some of the branches were compressed to improve the visibility of the tree, followed by the number of the grouped sequences in parentheses.


