RESEARCH ARTICLE

Antibiotic resistome mostly relates to bacterial taxonomy along a suburban transmission chain

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HIGHLIGHTS

 \bullet The $\alpha\mbox{-diversities}$ of resistome were lower in manure and compost than in soils.

• There were significant correlations between the resistome and bacterial taxonomy.

• Bacterial taxonomy was the highest in explaining resistome variances.

GRAPHIC ABSTRACT



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ABSTRACT

Antibiotic resistance genes comprising antibiotic resistome are of great concern due to their increase in the environment. Recent evidence of shared resistomes between soils and animal husbandry has imposed potential risks to human health. However, the correlation between a given community's resistome and bacterial taxonomic composition is controversial. Here, a transmission chain of resistomes from swine manure to compost and compost-amended soil were analyzed in five suburban areas of Beijing, China, with unamended agricultural soils as control soils. Antibiotic resistomes and bacterial taxonomic compost, and compost-amended soils, the β -diversity of the resistome and bacterial taxonomic composition was significantly correlated, while no correlation was detected in control soils. Bacterial taxonomic composition explained 36.0% of total variations of the resistome composition, much higher than environmental factors. Together, those results demonstrated that antibiotic resistome was closely related to bacterial taxonomic composition along the suburban transmission chain.

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1 Introduction

Bacterial resistance to antibiotics is a huge threat to human health, undermining medical treatments for pathogenbased infections worldwide. An increasing body of evidence illustrates that the widespread presence of antibiotic resistance genes (ARGs), termed the resistome when referring to all of detected ARGs, in human pathogens is associated with antibiotic usage (Marshall and Levy, 2011; Smillie et al., 2011; Forsberg et al., 2012). Therefore, resistome is regarded as an emerging biological hazard in various ecosystems (Smillie et al., 2011; Zhu et al., 2013; Su et al., 2017; Peng et al., 2020), whose dissemination is accelerated by human activities (Pruden et al., 2012; Chen et al., 2019).

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In modern societies, antibiotics are routinely used in livestock farming, which heightens the environmental burden of resistome (Zhu et al., 2013). China has become the largest producer and consumer of antibiotics in recent decades (Zhu et al., 2013). Until recently, unmonitored use of antibiotics to treat animal disease and improve growth has caused high concentrations of antibiotic residues in the environment (Qiao et al., 2012). Animal manure is a major route of antibiotic contamination because of poor absorption of veterinary antibiotics by the animals and common practice to use manure as organic fertilizer, leading to further diffusion (Alcock et al., 1999; Leclercq et al., 2016; Wu et al., 2020). Furthermore, the composting process of manure could be a hotbed causing enrichment of resistome and animal-associated pathogens. As a result, an increase in resistome is found in manure (Looft et al., 2012), compost (Liao et al., 2019), manure-amended soils (Ghosh and LaPara, 2007), riverine water, and sediments (Pruden et al., 2012).

Massive antibiotic use could escalate horizontal gene transfer (HGT) (Smillie et al., 2011; Forsberg et al., 2012; Gillings and Stokes, 2012; Ji et al., 2020). Accordingly, transferable, ARG-carrying plasmids were shown to be more abundant and persistent in manure-amended soils than in control soils (Jechalke et al., 2013; Jechalke et al., 2014). Because resistome is mobile, resistome may not be related to bacterial taxonomic composition (Smillie et al., 2011; Stokes and Gillings, 2011). However, the possibility was denied by observations that soil resistome composition was correlated with the microbial community composition across and within soil types, suggesting that bacterial taxonomy was influential for soil resistome (Forsberg et al., 2014). Similarly, Clostridium, Acinetobacter, and Pseudomonas were related to ARG enrichment in manureamended soils (Leclercq et al., 2016). A possible explanation is that resistome originating from manure amendments to soils promotes the growth of its bacterial hosts, resulting in significant correlations (Udikovic-Kolic et al., 2014).

As most studies have focused on a single ecosystem, it remains unclear how possible correlations between resistome and bacterial taxonomy vary with humaninduced dissemination of antibiotics. To this end, samples in manure, compost, compost-amended agricultural soil, and nearby unamended agricultural soil (control soil) were collected from five swine farms in Beijing's suburbs. Using the techniques of functional gene microarray named GeoChip and 16S rRNA gene amplicon sequencing, correlations between resistome and bacterial taxonomy were examined along the transmission chain. High concentrations of antibiotics impose environmental selection or pressure, potentially reducing the diversities of the resistome or associated microorganisms. Therefore, the resistome and microbial community profiles could vary with concentrations of antibiotics. Here, three hypotheses were aimed: 1) there are low diversities of resistome in

manure and compost samples because of high concentrations of antibiotics; 2) both resistome and bacterial taxonomy are distinct in different samples; and 3) resistome is correlated with the bacterial taxonomy, showing stronger correlations in manure and compost samples.

2 Materials and methods

2.1 Sample collection and environmental factor measurements

Four manure and four aerobic compost samples were collected in April 2015 from each of the five swine farms (i.e., CP in Changping district, DX in Daxing district, HZ in Haidian district, JG and ZZ in Shunyi district) within the suburbs of Beijing, China. Four samples of surface soil (0–15 cm) in agricultural fields receiving compost from each farm (i.e., compost-amended soil) for one month were collected, together with four control soil samples nearby (~1 km) unamended agricultural fields, resulting in 80 samples in total. Each soil sample is a composite by mixing three soil cores (5 cm diameter) for compost-amended soils and unamended agricultural soils.

Thermometers were used to measure temperature onsite, and hygrometers were used to measure water content, which was performed in triplicates and averaged. Commonly used antibiotics containing three categories of tetracyclines, sulfonamides, and quinolones in livestock practices, including chlortetracycline (CTC), oxytetracycline (OTC), tetracycline (TCN), norfloxacin (NOR), ofloxacin (OFL), sulfamerazine (SMR), sulfadimidine (SMN), sulfamethoxazole (SMX), and sulfadiazine (SDZ), were measured in the laboratory (Gao et al., 2020). In brief, those antibiotics were extracted by 10 mL of a phosphate buffer (pH = 3.0) with 0.1 g of Na₂EDTA and acetonitrile (C₂H₃N) (1:1 vol/vol). All samples were sonicated for 30 min before centrifugation at $7000 \times g$ for 10 min. The procedure was repeated three times to ensure efficient extraction. Three supernatants were mixed before dilution to 500 mL using deionized water, followed by filtering through 0.45 µm filters and pH adjustment to 3.0. Subsequently, samples were then extracted with 6 mL of Oasis HLB extraction cartridges and used for analyses with liquid chromatography-tandem mass spectrometry (LC-MS/MS) (Applied Biosystems, Foster City, CA, USA). The average recovery rate of antibiotics across samples was 90%.

2.2 DNA extraction from samples

For each sample, 1.5 g of sample was used to extract and purify DNA by the MoBio PowerSoil DNA isolation kit (MoBio Laboratories, Carlsbad, CA, USA). A NanoDrop ND-1000 spectrophotometer (NanoDrop Technologies Inc., Wilmington, DE, USA) was used to evaluate DNA quality based on the absorbance ratios of 260/280 nm and 260/230 nm. DNA concentrations were determined by PicoGreen with a FLUOstar Optima fluorescence plate reader (BMG Labtech, Jena, Germany). DNAs were then stored in freezers at -80° C for subsequent analysis.

2.3 Bacterial 16S rRNA gene amplicon sequencing and data processing

Sequencing and library construction were performed as previously described (Wu et al., 2017). The primer pair 515F (5'-GTGCCAGCMGCCGCGGTAA-3') and 806R (5'-GGACTACHVGGGTWTCTAAT-3') was used to amplify the V4 hypervariable region of bacterial 16S rRNA genes. A MiSeq benchtop sequencer (Illumina, San Diego, CA, USA) was used to perform sequencing in a $2 \times$ 250 pair-end manner. Raw sequences were uploaded to the online Galaxy sequence analysis pipeline. Then, raw sequences were assembled with FLASH (Magoč and Salzberg, 2011). The Joined sequences with vague bases or lengths less than 245 bp were discarded. Bacterial operational taxonomic units (OTUs) were clustered by UPARSE7 at a threshold of 97% identity. For the remaining sequences, singletons and chimeras were also discarded, and all sequences were then resampled to a depth of 31109 sequences for each sample. Representative sequences were used to annotate OTU from the Ribosomal Database Project (RDP) Classifier with a 50% confidence evaluation (Wang et al., 2007).

2.4 GeoChip hybridization experiments and data processing

GeoChip 5.0M, a functional gene array targeting genes associated with biogeochemical cyclings, was utilized to examine functional genes present in microbial communities (Zhang et al., 2017), with 11 ARG types including ABC transporter (ATP binding cassette transporter), MATE transporter (Multi-antimicrobial extrusion protein), Mex (Mex proteins in the RND family), MFS transporter (Major facilitator superfamily), SMR (small multidrug resistance), tet (tetracycline resistance), Fosfomycin Mod*ifying Enzymes*, *vgb* (virginiamycin B Lyase), β -lactamase, qnr (quinolone resistance), and van (vancomycin resistance). Briefly, the DNA of each sample was labeled by a fluorescent dye named Cy-3 dUTP, then hybridized to GeoChip at 67°C in a rotator/incubator with 20 r/min for 24 h. The arrays were washed after hybridization and then scanned by a NimbleGen MS200 Microarray Scanner (Roche, San Francisco, CA, USA). Data were extracted from the array images using Agilent Feature Extraction software version 11.5 and uploaded to the online Microarray Data Manager, as previously described (Ma et al., 2019). Detected signals with the signal-to-noise ratios < 2 were removed, and signals among arrays were normalized by the universal standard of spot intensities.

2.5 Statistical analyses

The α -diversity includes richness, Simpson index, and Shannon's index. The β -diversity was calculated based on Bray-Curtis dissimilarity. Tukey HSD test was used to determine the differences in α -diversity. PERMANOVA (permutational multivariate analysis of variance) (Anderson, 2001), MRPP (multi-response permutation procedure), and ANOSIM (analysis of similarities), all based on Bray-Curtis dissimilarity, were used to determine the differences of the resistome or bacterial taxonomic compositions between sample types or between sampling sites. Heatmaps were generated based on the Z scores of the relative abundances. For each ARG or bacterial phylum, a Z score was calculated as its average relative abundance in a type of samples (e.g., control soil) subtracted by the mean and then divided by the standard deviation of the four average relative abundances in four sample types. Pearson correlation was used to examine the relationships between resistome and bacterial taxonomic composition. Mantel tests were used to examine correlational relationships between the resistome and bacterial taxonomic community based on Bray-Curtis dissimilarity (Legendre and Legendre, 2012). The multiple regression on distance matrices (MRM) analysis was carried out to investigate relationships among the bacterial taxonomic composition of the community, the resistome present, and environmental factors (Lichstein, 2007). All analyses were performed in R.

3 Results

3.1 The resistome composition

There are 9277 ARG probes in the GeoChip microarray, which can be divided into 34 ARG subtypes belonging to 11 ARG types (Figs. S1(a) and S2). All of the 34 ARG subtypes were detected across samples. Those ARGs represent classical mechanisms of antibiotic resistance, including antibiotic efflux, antibiotic inactivation, and target protection/redundancy. MFS transporter, Van, and Mex were the most abundant, followed by ABC and MATE transporters (Fig. S1(b)). In contrast to the observation that antibiotics levels were much higher in manure and compost samples than in compost-amended soils and unamended soils (control soils) (Table S1), the α diversities of resistome were significantly lower in manure and compost than in soils (P < 0.050 by Tukey HSD test, Fig. S1(c)). Van, MFS, and MATE transporters were the most abundant in manure and compost samples, while MFS, Mex, and ABC transporters were the most abundant in compost-amended soil and unamended soil (Fig. S1(b)). In contrast, the α -diversities and abundance of resistome were not significantly different among CP, DX, HZ, JG, and ZZ sites, wherein each of the five dissemination chains

of animal farm-compost-compost amended soil was established, as well as control soils (P>0.050 by Tukey HSD test, Fig. S1(d)).

The overall composition of resistome was distinct among manure, compost, compost-amended, and control soils, as revealed by three non-parametric dissimilarity tests (MRPP, ANOSIM, and PERMANOVA, Table 1 & Fig. 1(a)). In compost-amended soil and control soil samples, B_lactamase_C, MFS_antibiotic, ABC_antibiotic transporter 2, B lactamase 5f0 D, ABC antibiotic transporter, b lactamase, Mex 5f0 RND antibiotic, fosA, MFS_antibiotic_5f0_other, B_lactamase_B_2, Van, MATE antibiotic 5f0 gammaproteo, and SMR antibiotics 2 showed relatively high abundances (Fig. 1(a)). In contrast, manure and compost samples had relatively high abundances of B lactamase A 5f0 2, MFS antibiotic 5f0 entero, fosB, Van 2, B lactamase A 5f0 firmicutes, MFS antibiotic 5f0 Firmicutes, Tet, vgb, MFS antibiotic 5f0 gammaproteo, MATE_antibiotic_5f0_other, and fosX (Fig. 1(a)).

3.2 The bacterial taxonomic composition

A total of 74676 bacterial OTUs were assigned to 54 phyla. Consistent with observations in resistome, the bacterial

taxonomic composition was distinct among manure, compost, compost-amended, and control soils (MRPP, ANOSIM, and PERMANOVA, Table 1 & Fig. 1(b)). Also, the bacterial taxonomic composition was distinct among the five sites. At the phylum level, Diapherotrites, Thaumarchaeota, Woesearchaeota, Acidobacteria, Armatimonadetes, BRC1, candidate division WPS-1, candidate division WPS-2, candidate division ZB3, Candidatus Saccharibacteria, Chlamydiae, Chlorobi, Chloroflexi, Cvanobacteria, Deferribacteres, Deinococcus-Thermus, Elusimicrobia, Fibrobacteres, Gemmatimonadetes, Hydrogenedentes, Ignavibacteriae, Latescibacteria, Marinimicrobia, Microgenomates, Nitrospirae, Omnitrophica, Parcubacteria, Planctomycetes, Synergistetes, Thermodesulfobacteria, and Thermotogae were more abundant in soils than in manure and compost (Fig. 1(b)). In contrast, SR1 and Tenericutes bacterial phyla were more abundant in manure and compost than in soils, consistent with a previous study showing Tenericutes as one of the dominant bacterial phyla in the manure composting process (Liu et al., 2020).

3.3 Linkages between resistome and bacterial taxonomy

Correlations between resistome and bacterial taxonomic



Fig. 1 Heatmaps of the (a) resistome and (b) bacterial phyla in manure, compost, compost-amended soil, and control soil, with 80 samples in total. The scale represents the relative abundance of each ARG or each bacterial phylum, which was normalized as a Z score detailed in the method.

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composition were tested in all samples. The richness of resistome and bacterial OTUs were significantly correlated (R = 0.486, P < 0.001 by Pearson correlation, Fig. 2(a)). Similar results were observed when the Shannon index (R= 0.596, P < 0.001, Fig. 2(b)) and the Simpson index (R =0.471, P < 0.001, Fig. 2(c)) were used as alternative indices of α -diversities. A strong, significant correlation was also observed for β -diversity between resistome and bacterial taxonomic composition (Bray-Curtis dissimilarity, R = 0.600, P < 0.001, Fig. 2(d)). Within each of five sampling sites, significant (P < 0.050) correlations between β-diversity of resistome and bacterial OTUs were observed (Table S2). However, when we divided samples into manure, compost, compost-amended, and control soils, there was no significant (P > 0.050) correlation for α -diversities (Table 2). In contrast, there were significant (P < 0.050) correlations for β -diversity in manure, compost, compost-amended soils, but not control soils (Table 2), which was verified by Mantel tests (Table S3). Since antibiotics concentrations were the lowest in control soils (Table S1), it was likely that its resistome was least selected by antibiotics, resulting in a largely stochastic resistome unlinked to bacterial taxonomy.

We examined correlations between individual bacterial genera and resistome to reveal bacteria with the strongest linkages to resistome, similar to earlier studies (Tao et al., 2016; Zhang et al., 2019). A total of 15 pairs of bacterial genera and ARGs showed positive correlations (P < 0.05) in manure samples and also in compost samples (Figs. 3(a) and 3(b), & Table S4). The ARGs are β -lactamase, fosfomycin modifying enzyme, MFS transporter, tet, Van, and vgb. Eight out of 13 bacterial genera were affiliated to Firmicutes, four were affiliated to Proteobacteria, and one was affiliated to Actinobacteria. Only two bacterial genera were associated with more than one ARGs (genus Acinetobacter ~ B lactamase A 5f0 firmicutes, genus Acinetobacter ~ tet; genus Alkalibaculum ~ fosX, genus Alkalibaculum $\sim vgb$), suggesting a minor to modest risk to form "super-bugs" (i.e., bacteria with multiple resistance to

Table 1 Non-parametric multivariate dissimilarity analyses of resistome and bacterial taxonomic composition

Group	Tractment	PERMANOVA ^{a)}		MF	RPP	ANOSIM	
Group	Treatment	R^2	Р	δ	Р	R	Р
Resistome ^{c)}	Sample types	0.425	0.001 ^{b)}	0.169	0.001	0.549	0.001
	Sample sites	0.074	0.001	0.216	0.067	0.032	0.082
Bacterial taxonomic composition	Sample types	0.321	0.001	0.715	0.001	0.735	0.001
	Sample sites	0.106	0.001	0.821	0.001	0.065	0.022

Notes: a) All three tests of the permutational multivariate analysis of variance (PERMANOVA), multiple response permutation procedures (MRPP), and analysis of similarity (ANOSIM) are based on Bray–Curtis distance. b) Significant *P*-values less than 0.050 are shown in bold. c) There are 80 samples in total.

Table 2 Pearson correlations between richness, Shannon index, Simpson index, and β -diversity (based on Bray-Curtis dissimilarity) of ARGs and bacterial OTUs in manure, compost, compost-amended soil, and control soil

Types	Indices	r	Р
Manure	Richness	0.145	0.543
	Shannon index	0.211	0.372
	Simpson index	0.215	0.363
	β-diversity	0.346	<0.001 ^{a)}
Compost	Richness	0.441	0.059
	Shannon index	0.284	0.238
	Simpson index	0.107	0.663
	β-diversity	0.430	< 0.001
Compost-amended Soil	Richness	-0.163	0.493
	Shannon index	-0.069	0.773
	Simpson index	-0.164	0.489
	β-diversity	0.318	< 0.001
Control Soil	Richness	-0.029	0.902
	Shannon index	-0.146	0.540
	Simpson index	-0.050	0.835
	β-diversity	-0.018	0.808

Notes: a) Significant P-values less than 0.050 are shown in bold.



Fig. 2 Pearson correlations between richness (a), Shannon index (b), Simpson index (c), and β -diversity (based on Bray-Curtis dissimilarity) (d) of ARGs and bacterial OTUs. The significance of the correlation was determined by P < 0.050. There are 80 samples in total.



Fig. 3 Significant positive Pearson correlation pairs between bacterial genera and resistome in the group of (a) manure, (b) compost, (c) compost-amended soil, and (d) control soil samples. The thickness of the edge represents correlation values. Green circles represent bacterial genus. Pink circles represent resistome. The size of the circle represents the degree of connectedness. The significance of the correlation was determined by P < 0.050.

different antibiotics). However, *fosB* genes were correlated with *Olsenella*, which is involved in endodontic infections in humans (Siqueira and Rôças, 2008). *Van* genes were correlated with *Tissierella*, which can cause chronic osteomyelitis (Schweizer et al., 2016) and prosthetic knee joint and bloodstream infections (Caméléna et al., 2016). Some *Acinetobacter* spp. are multidrug-resistant and can act as major infectious agents in debilitated patients (Towner, 2009). Similarly, *Stenotrophomonas* caused opportunistic infections in patients with cystic fibrosis, cancer, and HIV (de Oliveira-Garcia et al., 2002). Those results suggest that these bacterial species may act as hubs for HGT of resistome and increase potential public health risks.

A total of 17 pairs of bacterial genera and ARGs showed positive correlations (P < 0.05) in compost-amended soils and also in control soils (Figs. 3(c) and 3(d), & Table S5), which did not overlap with 15 pairs in manure and compost samples. The ARGs are *ABC transporter*, β -lactamase, MFS transporter, fosA, and Van. Twelve out of 16 bacterial genera were affiliated to the phyla Proteobacteria, Actinobacteria, and Firmicutes, while others were affiliated to Bacteroidetes, Acidobacteria, Armatimonadetes, and Chloroflexi. One bacterial genus was associated with both b lactamase and Van. In contrast with the situation of multiple pathogens detected in manure and compost samples, only van gene was related to Succiniclasticum, an opportunistic pathogen and endotoxin producer (Neubauer et al., 2019), and Aquabacterium, in compost-amended and control soils, indicating a relatively less risk to public health than manure and compost samples.

3.4 Factors influencing the resistome

The multiple regression on distance matrices (MRM) analysis was performed to identify major factors influencing environmental resistome. The bacterial taxonomic composition explained 36.0% (P = 0.001) of the resistome (Table 3), much higher than the explanatory power of soil

moisture (19.5%, P = 0.001) and the sample type (2.5%, P = 0.001). In contrast, the sample site and temperature were not important in shaping the resistome.

When samples were divided into manure, compost, compost-amended, and control soils, the bacterial taxonomic composition explained the resistome in manure (12.0%, P = 0.003), compost (18.5%, P = 0.001), and compost-amended soil (10.1%, P = 0.001). The sampling site explained 14.5% of the resistome (P = 0.001) in manure samples and 7.5% of the resistome (P = 0.005) in compost samples. These five different sample sites contained various antibiotics concentrations, which could be important in shaping the resistome. Soil moisture explained only 6.4% of the resistome (P = 0.040) in compost samples.

4 Discussion

In this study, significant correlations between resistome and bacterial taxonomy were detected in manure, compost, and compost-amended soil, but not controls soils, suggesting that interconnection between the resistome and bacterial taxonomy was environment-dependent. The results conflict with a previous study showing that bacterial phylogeny structures resistome in soils (Forsberg et al., 2014). The inconsistency may result from different soil types or technologies used across various studies. In addition, the results showed that control soils contained a large diversity of resistome and low antibiotic concentration settings (Fig. S1(c) & Table S1). As a result, high functional redundancy with a negligible environmental selection in control soils could obscure correlations with bacterial taxonomy.

Distinct profiles of ARGs regardless of similar geographical locations were observed in urban wastewater treatment plants (Fernandes et al., 2019). Similarly, variations of resistome and bacterial communities throughout the composting process were divided into two distinct stages, and total ARGs increased during composting (Cao

 Table 3
 Contribution of bacterial taxonomic composition and environmental factors in shaping the resistome by multiple regression on distance matrices (MRM) in all samples, manure, compost, compost-amended soil, and control soil

Influence – factors	All Samples		Manure		Compost		Compost-amended Soil		Control Soil	
	$R^{2a)}$	$\begin{array}{c} \text{coef} \\ (R^2 = 0.445) \end{array}$	R^2	$\begin{array}{c} \text{coef} \\ (R^2 = 0.230) \end{array}$	R^2	$\begin{array}{c} \text{coef} \\ (R^2 = 0.228) \end{array}$	R^2	$\begin{array}{c} \text{coef} \\ (R^2 = 0.118) \end{array}$	R^2	$\begin{array}{c} \text{coef} \\ (R^2 = 0.027) \end{array}$
Bacterial taxonomic composition	0.360*** ^{b)}	0.286***	0.120**	0.064*	0.185***	0.086*	0.101***	0.107***	0.0003	0.026
Sample Type	0.025***	-0.004	-	_	—	-	—	-	_	-
Sample Site	0.002	-0.002	0.145***	0.027**	0.075**	0.006	0.006	-0.003	0.008	-0.007
Temperature	0.003	-0.010**	0.022	0.002	0.057	0.005	0.001	-0.004	0.0004	0.003
Moisture	0.195***	0.018***	0.026	-0.016	0.064*	0.011	0.004	0.0001	0.014	-0.007

Notes: a) R^2 represents explained variance in the distance matrix regression model for each influence factor; and coefficient (coef) estimates the correlation of each factor in multivariate regression model of distance matrices. b) Significant *P*-values indicated as $*P \le 0.050$, $**P \le 0.010$, and $***P \le 0.001$.

et al., 2020). In this study, manure and compost samples were distinct from compost-amended and control soils (Fig. 1). Van genes encoding vancomycin resistance genes are often found in human pathogens, such as glycopeptideproducing Actinomycetes and nonglycopeptide-producing Actinomycetes (Hong et al., 2008). MATE (multidrug and toxic compound extrusion) transporters are also responsible for antibiotics uptake, conferring multidrug resistance to pathogens (Kusakizako et al., 2020). Both genes were among the most abundant in manure and compost samples (Fig. S1(b)), revealing potential health risks of human pathogens in manure and compost. In contrast, MFS (major facilitator superfamily) and ABC (ATP binding cassette) transporters are the most abundant in compost-amended and control soils (Fig. S1(b)). These two kinds of transporters can transport a wide spectrum of compounds in addition to antibiotics (Pao et al., 1998; Yan, 2013).

Firmicutes was the most abundant phylum in manure and compost samples (Fig. 1(b)). As well-known producers of antibiotics (Mannanov and Sattarova, 2001), Firmicutes was related to resistome (Huerta et al., 2013), leading to ARGs' dissemination as carriers. High abundances of Firmicutes in manure and compost coincided with the high antibiotic concentration (Table S1), imposing potential risks of resistome's dissemination. As the most abundant phylum in soil samples (Fig. 1(b)), Proteobacteria contain many multidrug-resistant strains (Forsberg et al., 2014) and are common carriers of class 1 integronsintegrase (Martinez et al., 2013). Feeding antibiotic-laced feed to pigs increased the abundance of Proteobacteria in the pig gut (Looft et al., 2012). The higher abundance of Proteobacteria increases antibiotic resistance in soils (Forsberg et al., 2012), representing a major conduit through which resistome is disseminated in the environment. It also increased the abundance of resistome for antibiotics classes not added to the feeding stock, promoting HGT of resistome within Proteobacteria or among different phyla (Klümper et al., 2015).

In manure and compost samples, tetracycline-resistance genes (tet) were correlated with several pathogenic bacterial genera such as Pseudocitrobacter, Subdoligranulum, Catonella associated with endodontic infections (Siqueira and Rôças, 2006), Clostridium IV, and Acinetobacter (Figs. 3(c) and 3(d)). Pseudocitrobacter is a novel genus isolated from hospitalized patients, while Subdoligranulum and Catonella are affiliated to the order Clostridiales, which includes several significant human pathogens. Clostridium perfringens, belonging to Clostridium IV, is a gastrointestinal disease pathogen that produces severe histotoxic hypoxia (Mendez et al., 2008). Tetracycline has been extensively used for treatment in various clinical conditions of humans and animals. In past decades, tetracycline resistance emerges due to the wide application of tetracycline for clinical, veterinary, and agricultural purposes, becoming a rising concern for clinical treatments. However, the correlations suggest

that these pathogenic taxa mentioned above could be potential hosts for tetracycline-resistance genes, differing from previous studies showing that the distribution of some tetracycline-resistance genes was limited to a few genera such as *Sphingobium* and *Enterobacteriaceae* (Ghosh et al., 2009; Zhang et al., 2009). Therefore, more alternative bacterial carriers of tetracycline-resistance genes remain to be discovered, including pathogens and bacteriophages.

Consistent with previous studies showing that bacterial community shaped the antibiotic resistome (Liao et al., 2019; Zhu et al., 2019; Zhang et al., 2020), there were significant correlations between resistome and bacteria colonizing in manure, compost, and compost-amended soils (Table 2). MRM also indicated that bacterial taxonomy was the primary factor influencing resistome in manure, compost, and compost-amended soil (Table 3). Those results suggest that the spread of resistome by potentially pathogenic bacteria could impose a challenge to medical treatments.

These findings have important implications because animal husbandry and manure composting are mature industries in modern society. As incremental anthropogenic activities could impose pathogenic threats, interconnected resistome and microbiome suggest that bacteria could concurrently cause the spread of resistome. Our finding is an important step toward understanding how resistome is disseminated in our environments and assessing the ecological impact of manure composting for potential pathogenicity, which is indispensable for knowledge-based policy management to control the environmental dissemination of resistome.

5 Conclusions

In a nutshell, the resistome and bacterial community were distinct among samples, with more potential pathogens related to ARGs identified in manure and compost. The resistome was significantly related to bacterial taxonomy in swine manure, compost, and compost-amended soils, but not in unamended agricultural soils. Therefore, bacterial taxonomy was the primary determinant of the resistome in the transmission chain of antibiotics.

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Data Availability GeoChip data are available in the NCBI GEO database under project no. GSE132839. DNA sequences of the 16S rRNA gene are available in the NCBI Sequence Read Archive under project no. PRJNA516026.

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