Occurrence and Distribution Characteristics of Antibiotic Resistance Genes in Sediments Between Urban and Rural of the Liaohe River Basin, China

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Abstract

Antibiotic Resistance Genes (ARGs) are considered to be emerging pollutants related to human activities. The rapid development of global urbanization has expanded human activities, thereby exacerbating the global human health risks caused by antibiotic resistance genes. The effects of urban and rural environments are multifarious, which makes the source and distribution of ARGs in the environment diversification. Understanding the distribution and spread of ARGs is essential for studying the environmental behavior of ARGs. In this study, the occurrence 296 genes were detected by the high-throughput qPCR technology, and FC value was used to analyze the diversity of ARGs and Mobile Genetic Elements (MGEs) in sediments between urban and rural areas of the Liaohe River Basin, China. The co-occurrence of MGEs and ARGs was analyzed using network to decipher core genes. A total of 187 ARGs and 10 MGEs were detected in all sediment samples. The average number of genes detected in urban sites is 89 higher than that in rural sites. The high abundance and various types of ARGs and MGEs detected in urban river sediments indicates that the occurrence of urban ARGs is more complex. MGEs were detected high levels and were significantly correlated with the abundance and diversity of ARGs in river sediments providing evidence that MGEs were related to the occurrence and distribution of ARGs and \textit{tnpA}(tnpA-07, tnpA-01 and tnpA-03) gene were at the key position of co-occurrence of various types of ARGs.

Introduction

Antibiotics have antibacterial properties, prevent diseases and promote growth (Danner et al 2019), so they are widely used in the prevention and treatment of human and livestock diseases, as well as in the breeding of livestock and aquaculture animals, which caused an increase in resistance to various antibiotics, posing a major threat to modern health work (Davies et al, 2010). Studies have shown that antibiotic resistance is a natural phenomenon, and human activities accelerate the spread of ARGs (Chen et al, 2013). At the same time, a variety of abiotic factors such as antibiotics, heavy metals and organic compounds are related to the occurrence of antibiotic resistance (Li et al, 2018). The appearance and distribution of ARGs in various environmental media have been found in sewage treatment plants (Rizzo et al, 2013), soil (Wang et al, 2014), sediments (Zhu et al, 2017), surface freshwater (Peng et al, 2019) and other locations. The river environment can provide an ideal environment for the physical transportation, acquisition and dissemination of ARGs, which is related to the comprehensive influence of the differences between urban and rural (Marti et al, 2014).

Antibiotics exhibit a pseudo-persistent state in the natural environment, which means antibiotics will continue to exert selective pressure on bacterial communities, increasing the possibility of spontaneous mutations and horizontal gene transfer in bacterial communities to produce emergence and spread of of ARGs (Grenni et al, 2018) among different microbial species through plasmids, phages and genome islands. In addition to the horizontal transfer of ARGs mediated by MGEs, ARGs can also promote ARGs to spread more frequently among different microbial species through plasmids, phages and genome islands. The comparative research (Nadine et al, 2015) on the functional river antibiotic resistance before
and after adding wastewater from sewage treatment plants showed that potential way to spread ARG in rivers through sewage treatment plants, and treatment significantly increases the copy number of aminoglycoside resistance genes in downstream rivers. At present, most researches focus on the characteristics of the occurrence of antibiotic resistance genes in river waters (Wang et al, 2016; Martie et al, 2018), while ARGs in sediments are 120 to 2,000 times higher than in water samples (Luo et al, 2010) and research on sediments of urban and rural is lacking.

The Liaohe River Basin has obvious characteristics of trans-regional rivers. In the urban clusters of the Liaoning privilege section of the Liaohe River Basin, the river water is overloaded to absorb the industrial wastewater and domestic sewage of the cities along the way. According to statistics, a total of 24,748 tons of antibiotics (Zhang et al, 2015) and $9.47 \times 10^{13}$ copies/person/day of ARGs in China have been released into rivers and related waterways (Su et al, 2017). Correspondingly, incomplete urban infrastructure construction has caused a large amount of urban sewage to flow into the river, which has brought huge pressure on the local environment during the rapid urbanization and industrialization process and has a huge impact on the ecology (Pan et al, 2012), causing contamination of antibiotics resistance genes (Wu et al, 2019). Relatively, rural pollution is mainly pollution from agriculture, fertilizers, and domestic waste (Tang et al, 2008, Wang et al, 2017). Comparison of urban and rural pollutant differences can better understand the speed and direction of the pollution process in the course of urbanization.

This study adopted a high-throughput qPCR design using 296 pairs of primers in urban and rural river sediments of the Liaohe River Basin. Comparative analysis of the urban-rural distribution of ARGs in the sediments of the Liaohe River Basin and exploring the diversity and differences of antibiotic resistance genes in the sediments of the region. Combining the characteristics of the urban and rural to explain their relationship with the surrounding environment provides a better understanding of the factors that affect the dynamic changes of ARGs in different environments. 296 ARGs and MGEs and 16S rRNA were detected and compared between urban and rural areas. Using reservoirs as a reference point, it is of great significance to study the distribution of ARGs in urban and rural river sediments, and to establish the relationship between ARGs pollution and urban and rural areas. The relationship between MGEs and ARGs was also explored. Understanding the core genes in the urban sediment environment is essential for formulating effective strategies to alleviate antibiotic resistance. The results of this work provide valuable data for evaluating the current status of ARGs in the study area and contribute to a better understanding of future ARGs pollution in urban and rural areas.

Materials And Methods

**Sample collection and physicochemical property determination**

In June 2019, the sediment sampling stations are set up in Figure 1: D1, H4, and T5 sampling stations are located in the city, and the surrounding population is densely populated. There are many hospitals nearby, and the river is more susceptible to human influence. H1, L4, and L7 are located in rural or mountainous
areas with relatively small populations, and human factors have little influence. H6, which is located at the reservoir point, was selected as the contrast point. Compared with the normal river environment, the reservoir environment may be less susceptible to the pollution of antibiotics and ARGs (Su et al, 2014), and most reservoir systems were located in some relatively primitive areas, with almost no human activities except for river input. The area were an important source of drinking water, which reflects the original pollution state of the river. At each sampling point, a sediment sampler was used to collect sediment samples within 2 cm of the mud-water interface. The sediment samples are collected and placed in aseptic bags and stored in an ice box (4°C), and taken back to the laboratory within 24 hours for storage. Stored in a refrigerator at -20°C for later use. The standard method (GB18668-2002) was used to determine the total organic carbon (TOC) content in the sediment in the laboratory.

**DNA extraction and High-throughput quantitative PCR**

The PowerSoil DNA Isolation Kit (MOBIO, CA) was used to extract DNA from the sediment with a 0.25 g sample according to the instructions of the kit. The concentration and purity of DNA were measured and evaluated on Nano-Drop spectrophotometer(Nano-drop Technologies Inc. Wilmington, DE). The absorbance value of DNA A260/A280 was 1.8 to 2.0, and the extracted DNA stored at -20 °C for subsequent analysis.

The Wafergen smart chip real-time PCR system was used for high-throughput qPCR reactions. The system could be used for large-scale gene expression studies, and could process 5184 nanocell responses per run (Wang et al, 2014). All primers used in this study are shown in Table 1. A total of 296 primer sets were used, including 295 primer sets, targeting almost all major categories of ARGs and mobile genetic elements (MGEs) and a 16S rRNA gene. ARGs are divided into 9 categories of antibiotic resistance genes (such as aminoglycosides, β-lactams, chloramphenicols, Macrolide-Lincosamide-StreptograminB (MLSB) resistance genes, multidrugs, sulfonamides, Tetracyclines, vancomycins and others) and MGEs. These ARGs represent almost all major classes of antibiotics widely used in humans and animals. The PCR reaction mixture was first added to the microwell chip using the (296 assays)×16 (samples) mode of the MSND (MSND), and then the qPCR reaction was performed on the cycler. Reaction system: 1×LightCycler 480 SYBR Gree I Master, 500nM each primer, DNA template 2ng/uL, total reaction volume: 100nL.

After the initial enzyme was activated at 95°C for 10 min, amplification was carried out using 40 cycles of the following procedure: denaturation at 95°C for 30 seconds, and annealing at 60°C for 30 seconds. The melting process is automatically generated by the Wafergen software. SmartChip qPCR software was used to analyze the results, multiple peaks or wells whose amplification efficiency exceeded the range (90%-110%) were excluded and then screened for (1) A threshold cycle (CT) must be ≤31, (2) Positive samples must be repeated three times simultaneously. Calculate the relative copy number (Eq.1) was calculated based on previous studies (Chen et al, 2018). In addition, the method of comparing CT was
also used to calculate the FC value of ARGs between the modified sample and the control (Eq. 2) (Schmittgen et al, 2008).

\[
\text{Gene copy number} = 10^{(31-C_T)/(10/3)}
\]

\[
\Delta C_T = C_{T(\text{ARG})} - C_{T(\text{16S})}
\]

\[
\Delta \Delta C_T = C_{T(\text{Target})} - \Delta C_{T(\text{Ref})}
\]

\[
FC = 2^{(-\Delta \Delta C_T)}
\]

\(CT\) is the threshold cycle, and the detection limit \(CT\) (31) is used to replace the non-amplified gene. ARG is one of 295 genes, 16S is the 16S rRNA gene, Target is the treatment group sample, and Ref is the control sample (Chen et al, 2016).

**Absolute quantification of 16S rRNA**

Amplification with Roche was carried out as follows: The qPCR reaction (20μL) consists of 10μL 2×LightCycler 480 SYBR®Green I Master Mix, 1μM per primer (the same primers used in high-throughput qPCR), 1μL DNA template, and 7μL nucleic acid-free water. Amplify with Roche 480 as follows: Initially preheat at 95 °C for 5 minutes, then perform 40 cycles at 95 °C for 15 s, 60 °C for 1 minute, and 72 °C for 15 s to contain 16S rRNA for clonal sequencing. The standard plasmid of the gene fragment \((1.39 \times 10^{10} \text{copies/L}^{-1})\) was used as the 8-point calibration curve of its 10-fold dilution for the calculation of the external standard \((r=0.96, P<0.01)\). Therefore, the relative copy number of ARGs generated by high-throughput qPCR can be converted to absolute copy number by normalizing to absolute 16S rRNA gene copy numbers.

**Data analysis**

SmartChip qPCR software (V2.7.0.1) was used to analyze the results of high-throughput qPCR, and discard multiple peaks and amplification efficiency beyond the range. The threshold period (CT) 31 is the detection limit. Samples with only three replicates were considered positive, the correlation analysis was conducted using Pearson's correlation coefficient, Wilcoxon signed-rank test was used for the diversity analysis and network construction. Data processing and charts were made with Excel 2016 and R Studio related software packages, TBtools (Chen et al, 2020) was used to make heat maps, and Gephi 0.9.2 was used to make co-occurrence network \((\text{Pearson}, p<0.05, \text{Closeness Centrality}, |\rho| \geq 0.6)\) structure diagrams.

**Results And Discussion**

**Diversity of ARGs and MGEs**
A total of 187 genes were detected in all samples, including 177 ARGs, 10 mobile genetic element (MGEs) genes (including 8 transposon genes and 2 integron genes) and 16S rRNA genes. The detected ARGs almost included all major types of antibiotic resistance genes (aminoglycosides, beta-lactams, chloramphenicols(Chlor), Macrolide-Lincosamide-StreptograminB (MLSB) resistance genes, multidrugs, sulfonamides(Sul), tetracyclines(Tet), vancomycin(Van) and others(Figure 2) and the three main resistance mechanisms including antibiotic inactivation, efflux pump, and cell protection, as well as other mechanisms, transposon and integron genes (Figure 3). In general, the average number of genes detected in urban sites is 89 higher than that in rural sites. 104 ARGs were detected at the urban site H4 with the most and 70 at the rural site L7 were the least. The number of genes detected is not significantly different between urban and rural areas (P=0.109>0.05), which indicated that the rural river sediments in the region may be affected by river flow, leading to insignificant differences between the urban and rural areas. The sediments in the Liaohe River Basin were influenced by a variety of ARGs, which is not only reflected in the diversity of the types of ARGs detected in the sediments, but also affected by ARGs with different resistance mechanisms. Drug-resistant bacteria against sulfonamides, tetracycline, chloramphenicol, quinolones and other antibiotics usually carry one or more resistance mechanisms ARGs, including cell protection mechanism genes such as sul2, tetM, tetT, etc. Drug efflux pump mechanism genes floR, tetH, ttgB, etc. The drug-resistant genes cata1, aadA1, and strB with inactivated drug active sites have been qualitatively detected in humans, livestock and poultry and isolates from multiple environmental media (McKinney et al, 2010). Antibiotic inactivation is the main resistance mechanism, and the total abundance of antibiotic inactivation 6.02*10^9 copies/g accounts for more than half of the detection of total resistance mechanism, which was similar to the detection result of Jiulongjiang (Ouyang, 2014). Compared with other resistance mechanisms, antibiotic inactivation mechanism genes are more conducive to the survival of resistant bacteria under the condition of antibiotic residues in the environment.

Through correlation analysis, it was found that there was no significant difference between the types of resistance genes and the number of resistance mechanisms detected in urban and rural areas (P=0.102>0.05)(Figure 2, Figure3). This may be due to the impact of the exchange effect caused by the river flow. However, based on the results of the cluster analysis of the points, there are differences in the average number and distance between the types of resistance genes and the number of resistance mechanisms detected in urban and rural areas. Compared with rural areas, ARGs induced by urbanization contain more types and mechanisms of drug resistance genes, and the corresponding effects of human factors such as urban hospital discharge and sewage discharge during urban treatment can not be ignored. So that, the number of types and mechanisms detected in urban areas (total 267) is higher than that of detection mechanisms than that checked out in the village (total 245) (Figure 3).

**Abundance of ARGs and MGEs**

**Absolute abundance of ARGs and MGEs**
The total abundance of the major types of ARGs and MGEs was $10^6$ copies/g~$10^9$ copies/g (Figure 4), aminoglycoside resistance gene (Aminoglycoside) $4.39 \times 10^9$ copies/g and multidrug resistance gene (Multidrug) $3.57 \times 10^9$ copies/g have the highest total abundance detected. Chloramphenicol (Chlor) and Vancomycin (Van) were the lowest detected, respectively $3.67 \times 10^7$ copies/g and $2.85 \times 10^7$ copies/g. The total abundance detected by MGEs was $1.76 \times 10^9$ copies/g, indicating that the horizontal gene transfer (Horizontal Gene Transfer, HGT) possibility of ARGs in the sample is higher. This is congruent with a previous report, which suggested that the abundance and diversity of integrons were correlated with the abundance and diversity of ARGs (Chen et al, 2013).

Among all the detected genes, genes such as *sul2, qacEdelta1-01* and *aadA-02* were highly detected in urban and rural areas; genes such as *vanHB, blaSHV-01* and *tetB-02* were relatively low in urban and rural areas (Figure 5). The highest gene is *qacEdelta1-01* with an abundance of $1.50 \times 10^9$ copies/g. The number of total genes detected was the highest at point H4, and the detected abundance was $8.72 \times 10^9$ copies/g, followed by T5, and L4 was the lowest (Figure 6). Since the level of ARGs is closely related to the degree of human influence, human activities have a high degree of impact on urban rivers. Czekalski studied the distribution characteristics of ARGs in the Vidi Bay of Lake Geneva and found that ARGs showed a trend of gradually decreasing from the center of the pollution source to the distance (Czekalski, 2014), while H4 is located in Shenyang city with dense population and numerous hospitals. It was speculated that the higher detection of ARGs at H4 was closely related to human activities. Lu studied the pollution status and fate of sulfa resistance genes in the Daliao River estuary and Liao River estuary, and the results also showed that changes in factors affecting the transmission ability of ARGs in the estuary area may lead to changes in the occurrence status of ARGs (Lu et al, 2015). D1 is located at the estuary of the Daliao River. Due to the interaction between seawater and river water, the detection of ARGs at D1 in cities was lower than that at L7 in rural areas. The lower ARGs detection at point D1 in the estuary area may be reflected to the combined effect of the ocean and rivers in the estuary area, while the human factors may be relatively weak.

Through comparative analysis of the absolute abundance of ARGs detected in urban and rural sites, it was found that there was significant difference between urban and rural ($P=0.039<0.05$), and the abundance level appears to be higher in cities than in rural (Figure 7). Except for vancomycin-type Van ARGs, the detection rates in urban were higher than those in rural. The ARGs detected in cities have the characteristics of high abundance, strong diversity, and diverse structures. The concentration of antibiotics in the rivers is detected at sub-inhibitory levels to produce the flora in the river environment certain pressure selection (Henando et al, 2006, Graham et al, 2010). The selection of antibiotic-resistant bacteria at low antibiotic concentrations is carried out in the presence of other environmental factors such as heavy metals and pesticides (Gullberg et al, 2011). The above conditions may cause the detection of ARGs in urban river sections to show the above characteristics. The increase in the abundance and number of ARGs in urban rivers may be influenced by the selective pressure exerted by antibiotics or other chemicals released by human and veterinary applications (Li et al, 2020).
**FC value of ARGs and MGEs**

The FC value represents the fold relationship between the detected gene and the control group. The same control is used to analyze the relative fold relationship between the study samples under different treatments or influencing factors. When the water passes through the water purification facilities of the reservoir, common pollutants, antibiotics and ARGs in the reservoir flowing from the river are removed, and the reservoir system effectively controls antibiotics and antibiotic resistance to acceptable levels (Li et al, 2020). Therefore, the H6 point of the upstream reservoir was selected as a reference to calculate the FC value of the detected ARGs and MGEs. Of the 177 ARGs and 10 MGEs detected, 55 (29.4%) were detected in urban and rural areas and controls (Figure 8), while there were 20(10.7%), 23(12.3%), and 21(11.2%) uniquely detected genes in urban, rural, and control sites, respectively. Compared with the number of common detections between the control group and the urban and rural areas, the common detection categories of the urban and rural areas overlap more, indicating that the environment after the reservoir treatment in the control group is different from the environment in the normal urban and rural river system. Factors such as river flow and human activities will also affect the occurrence of ARGs in river sediments. For the reservoir control, 21(11.2%) unique genes were detected, which may be caused by the decrease of water flow velocity in the ecological water storage area of the reservoir. The precipitation of suspended solids and organic matter in the pretreatment zone, and the combined influence of aquatic plants and microorganisms (Fang et al. 2017), indicating that the ARGs of the reservoir are affected by the combined effects of natural environmental factors and anthropogenic factor. The observation of 55 (29.4%) ARGs in the reservoir showed that they were resistant to all major antibiotics, and were enriched in rural and urban areas. This supports the existing findings that the background of drug resistance is widespread in the natural environment (D ‘Costa et al. 2011).

According to the distribution map of FC value of each category (Figure 9), it can be seen that the genes detected in sediments in urban and rural were generally higher than those in sediments in reservoirs. Aminoglycoside resistance gene was detected in urban and rural areas with a relatively high multiple of FC value range from 607.57 to 7176.08. It has showed that 83 genes (Figure 2) were detected at point D1 and the FC value at this point was 1.49~607.57, further indicating that ARGs pollution in the estuary area was influenced by different factors than urban and rural and control reservoirs. The FC heat map (Figure 10) shows that the urban (outer circle) is more enriched than the rural (inner circle). Further analysis of the ARGs profile in the control reservoir and urban and rural sediments, the ARGs in the urban sediments may be composed of two types of natural origin and exogenous sources (Li et al, 2020).

**Analysis of ARGs Influencing Factors**

The sum of the concentration of each type of ARGs is the total ARGs of this type of ARGs. The correlation between each type of total ARGs and 16S rRNA, TOC and MGEs was analyzed: Except for Vancomycin (Van) of 9 types of ARGs, the other types were positively correlated with MGEs (P<0.01 or P<0.05), indicating that MGEs play an important role in the transfer and spread of ARGs(Table 1). The absolute abundance of ARGs was related to 16S rRNA. 16S rRNA represents to a certain extent the level of
biomass in the sediments of the study sites. The increase or decrease of biomass may be another way to regulate ARGs (Huang et al, 2015). The low difference in biomass levels in urban and rural sediments in this study may be a reason for the insignificant difference in ARGs. MGEs can be used as the typical molecular characteristics of ARGs; which can try to track antibiotic-resistant pollution in rivers to identify the typical molecular characteristics of ARGs (Storteboom et al, 2010). Through the correlation diagram between sampling points and resistance gene MGEs (Figure 11), it can be seen that the H4, T5, and D1 points are highly correlated with ARGs and MGEs, which means the detected concentrations of ARGs and MGEs in urban were relatively higher than those in rural.

Table 1: Correlation between various types of ARGs and 16S rRNA, TOC and MGEs

|                      | 16S rRNA | MGEs  | TOC  |
|----------------------|----------|-------|------|
| Aminoglycoside       | 0.972**  | 0.999** | -0.299 |
| Beta_Lactamase       | 0.977**  | 0.997** | -0.324 |
| Chlor                | 0.978**  | 0.998** | -0.32 |
| MLSB                 | 0.979**  | 0.995** | -0.333 |
| Multidrug            | 0.789    | 0.891*  | -0.397 |
| Others               | 0.976**  | 0.997** | -0.312 |
| Sul                  | 0.79     | 0.887*  | -0.312 |
| Van                  | 0.395    | 0.433   | -0.409 |
| Tet                  | 0.979**  | 0.995** | -0.333 |

*Indicates a significant correlation P<0.05

** indicates a very significant correlation P<0.01

Constructing a network structure diagram of the co-occurrence of various ARGs and MGEs (Figure 12) shows that the tnpA gene in MGEs plays an important role in the co-occurrence of ARGs. TnpA is a member of the serine recombinase (SR) gene family. It has reported that the full-length TnpA proteins of IS607, IS1535 and ISC1926 bind to the DNA at the ends of their respective transposons, and chemically active TnpA oligomers play a role. The tnpA gene was at the key position of co-occurrence of various types of ARGs. The transfer of ARGs in this region is closely related to tnpA (Chen et al, 2018b). It also found that tnpA-related genes played a key role in the network structure diagram, which showed that tnpA-07 and tnpA-01 were detected at each site. TnpA-01 and tnpA-03 mainly played key node roles in the co-occurrence network of aminoglycosides, sulfonamides, β-lactams and multidrug resistance genes. In short, tnpA-07 plays a key role in the co-occurrence network of macrolides and chloramphenicol resistance genes, and the connection point of the two large modules is tnpA-03. Previous studies have shown that MGEs, as the main promotion unit of horizontal transfer, play an important role in the
occurrence and spread of ARGs (Chen et al, 2018, Zheng et al, 2018). In summary, the occurrence and spread of ARGs in sediments is highly correlated with MGEs, especially the *tnpA* gene plays a key role in the co-occurrence of ARGs. The biomass characterized by 16S rRNA also plays a certain role in ARGs, such as microbial community differences play a key role in the distribution of ARGs (Xiang et al., 2018). Biomass differences in urban and rural sediments, river flow, environmental factors caused by human factors, and environmental factors such as ocean and river exchange under natural conditions have an impact on the occurrence and spread of sediment ARGs.

**Conclusions**

In summary, this study analyzed the distribution and enrichment of 177 antibiotic resistance genes, 16SrRNA, MGEs and the distribution of resistance mechanisms detected in the sediments of the Liaohe River Basin. It shows that Antibiotic resistance gene pollution is widespread in the sediments of the Liaohe River Basin. Through the comparative analysis of urban and rural, it was found that there was significant difference in the detection between urban and rural (P<0.05) and the urban resistance gene pollution is higher than that in the rural areas. MGEs play an important role in the occurrence of ARGs and *tnpA-07, tnpA-01* and *tnpA-03* mainly played key node roles in the co-occurrence network, which indicates that *tnpA* was important to the co-occurrence of ARGs. ARGs in urban and rural may be related to many factors, such as differences in pollution between urban and rural areas, differences in human activities, differences in the original existence of urban and rural natural environments, and river geographic locations such as estuaries etc.

**Declarations**

**Authors Contributions**

GSN: Conceptualization, Methodology, Supervision, Funding Acquisition; KYZ: Conceptualization, Methodology, Investigation, Visualization, Data Curation, Formal Analysis, Writing - Original Draft, Writing - Review & Editing; HG: Methodology, Investigation, Validation, Resources, Writing - Review & Editing; RJL: Investigation, Software, Resources; SCJ: Investigation, Software, Resources; FQZ: Visualization, Investigation, Resources; HBZ: Validation, Software, Resources; SSL: Investigation, Resources

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**Figures**
Figure 1

Sampling stations. Note: The designations employed and the presentation of the material on this map do not imply the expression of any opinion whatsoever on the part of Research Square concerning the legal status of any country, territory, city or area or of its authorities, or concerning the delimitation of its frontiers or boundaries. This map has been provided by the authors.
Figure 2

The number of ARGs and MGEs detected in each category

Figure 3
The number of detected resistance mechanisms in each category

Figure 4

Total absolute abundance of ARGs, MGEs and 16SrRNA
Detection heat map of the absolute abundances of ARGs and MGEs (using the logarithmic value to base 2). Cluster IV means that the detection in rural areas is higher than that in cities; Cluster III means that both urban and rural detections are low or not detected (only in the reference point); Cluster II means that the detections in cities are higher than rural; Cluster I means that both urban and rural detections are high.
Figure 6

Detected absolute abundances of various types of ARGs and MGEs (using the logarithmic value based on 2)

Figure 7

Comparison of the detection of ARGs and MGEs in urban and rural. (P=0.039<0.05)
Figure 8

Venn diagram of ARGs and MGEs detected in urban, rural and control
Figure 9

Distribution of FC values detected by ARGs and MGEs
Figure 10

Heat map of FC values detected by ARGs and MGEs (take the logarithmic value based on 2)
Figure 11

Association diagram of sampling points and resistance genes. The left side of the figure represents each sampling point, and the right side represents different types of antibiotic resistance genes; the left and right sides were connected with each other to indicate the degree of association between the corresponding site and the resistance gene. The value is 0% to 100%, corresponding the higher the value, the higher the concentration of resistance gene at the locus.
Figure 12

Analysis of the network structure of the co-occurrence of MGEs and ARGs, $|p| \geq 0.6$, $P<0.05$. Each node in the figure represents the detected gene. The larger the point, the more important the co-occurrence position. The color of the point indicates different types of ARGs or MGEs.

Supplementary Files

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