NORFLOXACIN RESISTANT BACTERIAL COMPOSITIONS IN SEDIMENTS OF CHINESE SUBTROPICAL FISH PONDS


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Abstract. Antibiotic resistant bacteria are widely spread in environments. However, the compositions of bacteria with antibiotic resistance in subtropical fish pond sediment are still unclear. In present study, to analyze the compositions of norfloxacin resistant bacteria in subtropical fish pond sediments, we selectively cultured the sediment bacteria collected from six subtropical fish ponds located in southern China using the basic medium with 50 µg/ml of norfloxacin, and analyzed their compositions using MiSeq high-throughput sequencing of 16S rRNA gene amplicons. Our results showed that various norfloxacin resistant bacteria existed in the fish pond sediments. And most of them were Gram-negative bacteria. Their spatial distribution was mainly influenced by environmental heterogeneity of the ponds. These results implied that norfloxacin resistance was widespread in Gram-negative bacteria in the fish pond sediments located in southern China. These finding provided reference information to prevent fish infected diseases and to assess the risk of antibiotic resistant bacteria in the fish pond sediments.

Keywords: antibiotic, bacterial community, environmental heterogeneity, Gram-negative bacteria, high-throughput sequencing

Introduction

Bacterial pathogens are the major cause of infectious diseases and mortality in wild and farmed fish (Sudheesh et al., 2012). Disease problems constitute that the largest single cause of economic losses in aquaculture (Meyer, 1991). With the rapid growth and intensification of aquaculture, the list of new pathogenic bacterial species isolated from fish has been steadily increasing (Harvell et al., 1999). To control the pathogenic bacteria and enhance fish growth, antibiotics have been widely used in agricultural production over the past decades (Ma et al., 2001; Neela et al., 2015). However, extensive use of antibiotics, especially the overuse of antibiotics in agricultural production, has caused a serious threat of antibiotic resistance (Levy and Marshall, 2004; Goossens et al., 2005; Mathew et al., 2007; Kumarasamy et al., 2010; Laxminarayan et al., 2013). Although the usage of antibiotics in agricultural production is strictly restricted presently, residues of antibiotics in environment and illegal use of antibiotics still cause prevalence of antibiotic resistance, especially in the aquatic systems, such as lakes and fish ponds (Wahid et al., 2014; Moore et al., 2014; Far et al., 2015; Neela et al., 2015; Patil et al., 2016; Yang et al., 2016). In addition, there are many antibiotics used in aquaculture, such as terramycin, also used in clinical trials in China (Ma et al., 2001).

Pond culture is one of the most prevailing culture patterns in China, which is the largest producing area of aquatic products around the world (FAO, 2014; Lu et al., 2015). Sewage discharge and agricultural production cause residues of antibiotics in ponds (Lin et al., 2010; Zhang et al., 2011; Wei et al., 2011; Song et al., 2016; Cheng et
al., 2016). These residues of antibiotics increased the risk of antibiotic resistance. Although the genes of antibiotic resistance in the ponds have been widely investigated (Cheng et al., 2016), and a lot of antibiotic resistance bacteria have been isolated from many ponds (Zhang et al., 2011; Wei et al., 2011; Neela et al., 2015), the compositions of bacteria with antibiotic resistance in the ponds are still unclear.

Screening antibiotic resistant bacteria through culturing by mediums containing antibiotics is commonly used method to study antibiotic resistant bacteria (Kumarasamy et al., 2010; Neela et al., 2015; Li et al., 2017). However, the previous studies mostly focused on signal or few bacterial species. Although high-throughput sequencing technology has been widely used in microbial community analysis (Huang et al., 2018; Ni et al., 2018; Xiang et al., 2018), it is still not to use to investigate composition of antibiotic resistant bacteria in pond sediment.

Norfloxacin is commonly used antibiotic in aquaculture and is frequently detected from various water environments and aquatic animals (Ortiz et al., 1999; Guo and Zhang, 2009; Spongberg et al., 2011; Zhang et al., 2017). To analyze the compositions of antibiotic resistant bacteria in pond sediment, we selectively cultured the sediment bacteria collected from six subtropical fish ponds in southern China using the basic medium containing with norfloxacin and analyzed their compositions using MiSeq high-throughput sequencing of 16S rRNA gene amplicons. Results of the present study provided reference information to prevent fish infective diseases and assess the risk of antibiotic resistance in Chinese subtropical fish ponds.

Materials and methods

Sampling collection and treatment

Sediment samples were collected from 5 fish ponds (GraA, GraF, LatF, GruF, and OreD) located in Yuanzhou Town (113°57’ E, 23°07’ N) and 1 fish pond (ChiP) located in Huicheng District (114°23’ E, 23°05’ N) of Huizhou, a subtropical city in southern China on January 22, 2016 (Fig. 1). Fish species farmed in the ponds were showed in Table 1. Three sediment samples were parallelly collected from each pond. Five-point sampling method was used to collect each sediment sample (i.e., five sub-samples were collected at area within 1 m of diameter, and then pooled together with an equal wet weight as one sample).

Table 1. Fish species farmed in the ponds. All fish ponds located in Huizhou, a subtropical city in southern China

<table>
<thead>
<tr>
<th>Pond name</th>
<th>Pond location</th>
<th>Pond surface area (m²)</th>
<th>Fish species</th>
</tr>
</thead>
<tbody>
<tr>
<td>GraA</td>
<td>Yuanzhou Town</td>
<td>4,330</td>
<td>Grass carp (Ctenopharyngodon idellus) and tilapia (Tilapia mossambica)</td>
</tr>
<tr>
<td>GraF</td>
<td>Yuanzhou Town</td>
<td>3,000</td>
<td>Fries of grass carp</td>
</tr>
<tr>
<td>LatF</td>
<td>Yuanzhou Town</td>
<td>3,330</td>
<td>Weever (Lateolabrax japonicus)</td>
</tr>
<tr>
<td>GruF</td>
<td>Yuanzhou Town</td>
<td>2,330</td>
<td>Crucian carp (Carassius auratus)</td>
</tr>
<tr>
<td>OreD</td>
<td>Yuanzhou Town</td>
<td>4,670</td>
<td>Tilapia (Tilapia mossambica)</td>
</tr>
<tr>
<td>ChiP</td>
<td>Huicheng District</td>
<td>5,330</td>
<td>A mixture of multiple fishes</td>
</tr>
</tbody>
</table>

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Triplicate 1 g of sediments (wet weight) of each sample were added into conical flask containing 100 ml of the basic medium with 50 µg/ml of norfloxacin referenced a previous report (Ma et al., 2012) and were cultured 24 h at 30 °C in an incubator with 150 r/min. Then the inoculums were centrifugated 10 min at 1200 rpm to concentrate and the suspensions were transferred into 2 ml centrifuge tubes for microbial DNA extraction. The microbial DNA were extracted according to previous description (Fang et al., 2015) and purified using a gel extraction kit (Dingguo, China). DNA concentrations were measured using a Nanodrop 2000 spectrophotometer and diluted to 10 ng/µl for PCR amplification.

**PCR amplification and sequencing**

V4-V5 hypervariable region of 16S rRNA gene was amplified and sequenced using a MiSeq system, as described previously (Ni et al., 2017; Huang et al., 2018; Xiang et al., 2018). Briefly, the V4-V5 region was amplified using the universal primer pair 515F (5’ – GTG YCA GCM GCC GCG GTA – 3’) and 909R (5’ – CCC CGY CAA TTC MTT TRA GT – 3’) with a 12-nucleotide sample-specific barcode included at the 5’-end of the 515F sequence to distinguish samples. PCR was conducted in duplicate with 25-µl reaction mix containing 1 × buffer, 0.25 U of Taq DNA polymerase (Transgen, China), 0.2 mM of each dNTP, 1.0 µM of each primer and 10 ng DNA. The PCR procedure consisted of a pre-denaturation step at 94 °C for 10 min, followed by 30 cycles of 94 °C for 30 s, 56 °C for 30 s and 72 °C for 30 s and a final extension at 72 °C for 10 min. After amplification, the two products of each sample were mixed together and subjected to electrophoresis using a 1.5% agarose gel. The correct band was excised, purified using a gel extraction kit (Axygen, USA), and quantified with the Nanodrop 2000 spectrophotometer. All amplicons were pooled together with an equal molar amount for each sample and sequencing using an Illumina Miseq system at Guangdong Meilikang Bio-Science, Ltd., China.

**Data analysis**

The raw sequencing reads were merged using FLASH-1.2.8 software (Magoc and Salzberg, 2011) and per-processed for removing low-quality sequences using QIIME.
pipeline version 1.9.0 (Caporaso et al., 2010) as previously described (Ni et al., 2017; Huang et al., 2018; Xiang et al., 2018). Chimera sequences were identified and removed before further analysis using the Uchime algorithm (Edgar et al., 2011). The high-quality sequences were clustered into operational taxonomic units (OTUs) at 97% identity using UPARSE software (Edgar, 2013). Then all samples were randomly resampled to obtain the same number of sequences to overcome the influence of sequencing depth on results. Each OTU was assigned taxonomic information using the RDP classifier (Wang et al., 2007) with Greengenes database gg_13_8.

Principal component analysis (PCA) and non-parametric multivariate analysis of variance (MANOVA) (Anderson, 2001) were used to analyze the difference of microbiota among different groups and were conducted using R 3.5.1 with vegan package (Dixon, 2003). The box plots were drawn to show the relative abundances of significantly different OTUs among groups using STAMP software (Parks et al., 2014). And the statistically significant markers were added to the box plots using Adobe Illustrator CS5 software according to the post-hoc test results.

All DNA datasets have been submitted to the NCBI Sequence Read Archive database under accession number SRP160514.

Results and discussion

Composition of norfloxacin resistant bacteria in pond sediments

Removing low-quality and chimera sequences, 346,643 high-quality sequences were obtained from the 18 samples. Finally, each sample was randomly resampled 11,000 sequences to further analysis. Except for little sequences (0.97 ± 0.38%, mean ± S.E.) could not classified into an explicit phylum, other sequences were classified into 2 Archaea and 65 Bacteria phyla. However, according to previous definition by other researchers (Huang et al., 2018; Xiang et al., 2018), only 15 phyla – Crenarchaeota, Euryarchaeota, Acidobacteria, Actinobacteria, Bacteroidetes, Chlorobi, Chloroflexi, Cyanobacteria, Firmicutes, Nitrospira, OP8, Planctomycetes, Proteobacteria, Spirochaetes, and Verrucomicrobia – dominated the sediment microbiotas (their relative abundances were more than 1% in at least one sample; Fig. 2). They contained up to 98.19 ± 0.72% of the sequences.

Total of 35,194 OTUs were detected from the 18 sediment microbiotas. However, only 48 OTUs dominated the microbiotas (their relative abundances were more than 1% in at least one sample; Fig. 3). They contained up to 67.16 ± 5.79% of the sequences. Lots of the dominant OTUs were reported as fish and mammal potentially pathogenic bacterial species, such as Acinetobacter sp., Lactococcus sp., Escherichia sp., Burkholderia sp., Streptomyces lanatus, Bacteroides sp., and Ruminococcus sp. (Slots and Listgarten, 1988; Kofteridis et al., 2007; Anandham et al., 2010; Fishbain and Peleg, 2010; Sudheesh et al., 2012; Woods and Sokol, 2006; Titécat et al., 2014; Li et al., 2017). And some of them were reported with multi-drug resistance (Li et al., 2017). In addition, potential plant bacterial pathogen Erwinia sp. and Ralstonia sp. (Swanson et al., 2005; Amin et al., 2011; Kube et al., 2010) also dominated the microbiota. Their norfloxacin resistance increased the bacterial infected risk of fish and plants.

Although norfloxacin is a broad-spectrum antibiotic, especially has a strong bactericidal effect to Gram-negative bacteria, most dominant OTUs were Gram-negative bacteria. This result implied that norfloxacin resistance was widespread in Gram-negative bacteria in the fish pond sediments located at southern China.
Figure 2. Dominant phyla of norfloxacin resistant bacteria in Chinese subtropical pond sediments. The sediment samples were collected from 5 fish ponds (GraA, GraF, LatF, GruF, and OreD) located at Yuanyou Town (113°57' E, 23°07' N) and 1 fish pond (ChiP) located at Huicheng District (114°23' E, 23°05' N), in Huizhou, a subtropical city in southern China on January 22, 2016. The pond GraA mainly farmed grass carp (Ctenopharyngodon idellus) and tilapia (Tilapia mossambica). The pond GraF farmed fries of grass carp. The pond LatF farmed weever (Lateolabrax japonicus). The pond GruF farmed fries of crucian carp (Carassius auratus). The pond OreD farmed tilapia (Tilapia mossambica). The pond ChiP farmed a mixture of multiple fishes.

Figure 3. Heat map profile showed dominant OTUs of norfloxacin resistant bacteria in pond sediments. The data was transformed according to the equation $\log_{10}(\text{relative abundance} \times 100 + 1)$ to reduce the magnitude of extremum. The sediment samples were collected from 5 fish ponds (GraA, GraF, LatF, GruF, and OreD) located at Yuanyou Town (113°57' E, 23°07' N) and 1 fish pond (ChiP) located at Huicheng District (114°23' E, 23°05' N), in Huizhou, a subtropical city in southern China on January 22, 2016. The pond GraF farmed fries of grass carp. The pond LatF farmed weever (Lateolabrax japonicus). The pond GruF farmed fries of crucian carp (Carassius auratus). The pond OreD farmed tilapia (Tilapia mossambica). The pond ChiP farmed a mixture of multiple fishes.
Different of norfloxacin resistant bacteria among different ponds

Mechanisms forming and maintaining bacterial bio-diversity are the basic issue of microbial ecology (Ni et al., 2014, 2016; Wu et al., 2017).

Figure 4. PCA profile based on OTUs (A) and based on dominant OTUs (B), and LEfSe profile based on dominant OTUs (C) showed differences of norfloxacin resistant bacteria in pond sediments. The sediment samples were collected from 5 fish ponds (GraA, GraF, LatF, GruF, and OreD) located at Yuanzhou Town (113°57’ E, 23°07’ N) and 1 fish pond (ChiP) located at Huicheng District (114°23’ E, 23°05’ N), in Huizhou, a subtropical city in southern China on January 22, 2016. The pond GraF farmed fries of grass carp. The pond LatF farmed weever (Lateolabrax japonicus). The pond GruF farmed fries of crucian carp (Carassius auratus). The pond OreD farmed tilapia (Tilapia mossambica). The pond ChiP farmed a mixture of multiple fishes.
Geographical isolation is one of the primary factors that restrict microbial distribution and emerge a significant distance-decay relationship regarding microbial community similarity (Bell, 2010). However, protesters claim microorganisms are cosmopolitan, and spatial patterns of microbial diversity are driven by environmental heterogeneity (Green and Bohannan, 2006). Our results showed that although significant differences of sediment microbiotas were detected among different ponds both based on the OTU compositions (MANOVA, $F = 1.851, p = 0.002$; Fig. 4A) and based on the dominant OTU compositions (MANOVA, $F = 1.926, p = 0.016$; Fig. 4B). And *Bacteroides* sp. was significantly enhanced at pond GruF, *Alicyclobacillus* sp. and an unidentified OTUs in family Methanoregulaceae was significantly enhanced at pond OreD, and *Escherichia* sp. and an unidentified OTUs in family Enterobacteriaceae was significantly enhanced at pond Chip (Fig. 4C). However, the microbiotas from the same pond did not cluster together except for those from pond GraA (Fig. 4A and B). These results showed that environmental heterogeneity was the major factor that casted the spatial patterns of microbial diversity.

Conclusion

In conclusion, we firstly investigated the norfloxacin resistant bacteria in fish pond sediments using the method that connected selective culture and high-throughput sequencing. Our results showed norfloxacin resistance was widespread in Gram-negative bacteria, such as *Acinetobacter* sp. and *Burkholderia* sp., in fish pond sediments located at southern China. This increased bacterial infected risk of farmed fish. In addition, mechanisms of these bacteria to resistant to norfloxacin and how to reduce the norfloxacin resistant bacteria are still needed to further study.

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