The Larval Density of Mosquitos (Diptera: Culicidae) in Jiaxiang County, Shandong Province, China: Influence of Bacterial Diversity, Richness, and Physicochemical Factors

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As Jiaxiang County of Shandong province is an area with complex mosquito vector composition, it is necessary to investigate the relationship between bacterial diversity, physicochemical factors, and larval density. Therefore, the physicochemical properties of 46 breeding sites for six kinds of habitat types (small puddles, small water containers, paddy fields, large water containers, irrigation channels, and drainage ditches) were investigated by a multiparameter analyzer; the water’s bacterial diversity was analyzed by the 16S rRNA full-length sequencing method. Spearman correlation and multiple linear regression were used to analyze the correlation between larval density and variables. The variables analyzed were dissolved oxygen, pH, hardness, turbidity, conductivity, temperature, ammonia nitrogen, water depth, and distance from the nearest house. One-Way ANOVA was used to understand whether there are differences in bacterial diversity in different habitats. Pearson linear correlation model was used to analyze the effects of bacterial diversity and richness on mosquito densities in breeding sites. A total of 3291 larvae were captured, and a total of 6 species of 4 genera were identified. The identified species were *Culex p. pallens*, *Aedes albopictus*, *Anopheles sinensis*, *Culex t. tritaeniorhynchus*, *Culex b. bitaeniorhynchus*, and *Mansonia uniformis*. The density and species can be jointly affected by physicochemical properties and bacterial diversity, especially Shannon index and distance from the nearest house. In general, the physicochemical parameters and bacterial diversity of different habitats were significantly different. Even for the same habitat type, the physicochemical parameters varied greatly due to different environments.

Keywords: full-length sequencing, habitat types, density, mosquito, bacterial diversity
INTRODUCTION

Mosquito is an important medical arthropod transmitting various mosquito-borne diseases such as malaria, dengue fever, and Japanese encephalitis. The larva and pupa stages of various mosquitoes are in water all the time, and the water body with positive larvae is regarded as their breeding sites (El-Ghani et al., 2012; Guo et al., 2017; Ferreira-de-Lima and Lima-Camara, 2018). Jiaxiang County is located in the Southwest Plain of Shandong Province, China. It has a temperate monsoon climate, numerous rivers and lakes, and a developed water system. All these conditions are suitable for breeding a variety of mosquitoes. *Culex pipiens pallens* and *Aedes albopictus* are the dominant mosquito species (Wang et al., 2020). The former is a potential vector of the West Nile virus (WNV) in Shandong Province (Jiang et al., 2014). The latter caused an outbreak of the dengue virus in Jiaxiang County in August 2017. There were 79 cases, which made the area a focus of attention for epidemic prevention personnel (Shi et al., 2019; Liu et al., 2020). In addition, *Anopheles sinensis* is the most important plasmodium vector in Shandong Province. In 2010, Shandong Province reported 117 malaria cases, among which 70% were imported cases. This may be related to the large rice and lotus cultivation area in southwest Shandong Province, which created a favorable breeding environment for Anopheles (Liu et al., 2015).

It is widely believed that larval source management (LSM) can effectively prevent mosquitoes' breeding and development. Researches show that LSM has an inhibitory effect on transmitting mosquito-borne diseases, whether in the countryside or a city. However, Jiaxiang County is rich in water resources, is densely populated, and its residents have a weak awareness of prevention, applying unreasonable mosquito control methods, and inadequate sanitation management. Hence, it is difficult to manage the larval breeding sites in this area (Wang et al., 2020). The Breteau Index (BI), an indicator of the density of *Aedes* mosquitoes in an area, can be used to assess the risk of dengue epidemics in that area. According to reports, the average BI of Jiaxiang County in August 2017 hit a stunning 107.27, whereas, for 2017 and 2018, it was 45.30 and 15.95, respectively (Liu et al., 2020). Areas where BI>20 will be regarded as high-risk means that any importation of external cases could cause a dengue epidemic in this area (Aryaprema and Xue, 2019). These data indirectly indicate that *Ae. albopictus* density in Jiaxiang County is high, and there is a potential risk of mosquito-borne virus transmission. In addition, in 2018, the resistance ratio of *C. p. pallens* to DDVP, propoxur, cypermethrin and deltamethrin was 19.76, 4.6, 438, and 351, respectively. Also, over the last decade, the resistance of *Cx. p. pallens* to different types of insecticides has risen to varying degrees (Liu et al., 2019; Wang et al., 2020). Therefore, scientific guidance is urgently needed to carry out LSM in this county. To some extent, understanding the larval density and distribution in this area, as well as the environment of breeding sites, is the first step in this plan.

While choosing the spawning site, female mosquitoes follow specific cues such as physicochemical factors including dissolved oxygen (DO), pH, hardness, turbidity, conductivity, temperature, ammonia nitrogen (AN), water depth (WD), and several biological factors (e.g., the composition of microorganism in water) (Juliano et al., 2004). They try to avoid water bodies where competitors and/or predators are already present and prefer to choose safe water bodies with specific physicochemical properties and microorganisms (Bentley and Day, 1989; Lin et al., 2008). In addition, the distance from the nearest house (DNH) is an important factor that needed to be considered to estimate the significance of the study to public health (Killeen et al., 2001). The research demonstrated that the distance between breeding sites and the nearest house determined the level of blood uptake by a female mosquito, which in turn affected the density, emergence rate of larvae, growth cycle, and malaria transmission dynamics (Killeen et al., 2001). However, with the expansion of urban sprawl to rural areas and frequent personnel exchanges, significant changes have taken place in the environment of breeding sites (Jacob et al., 2005). This affects the density and distribution of mosquito larvae, which, to a certain degree, increases the risk of transmission of mosquito-borne diseases and boosts the pressure of its prevention and control as well.

Bacteria in the breeding sites and other biological factors jointly create the external environment in which the larvae can live, exchanging materials directly with larval intestines (Dada et al., 2018). This can directly affect the survival status and even the density of the larvae (Mukhopadhyay and Chatterjee, 2016). Hence, it is necessary to investigate the composition of bacteria of water in breeding sites. This information will enable us to estimate whether water bacterial diversity was related to larval density. To our knowledge, habitat analysis on breeding sites is very limited, and reports on bacterial diversity and richness.

Selection of the Study Area

Jiaxiang County (35°11’N, 116°06’E) is located in the southwest of Shandong Province. Jiaxiang County is hot and rainy from June to August, with an average rainfall of 398.5 mm in summer. Zhu Zhaoxin River passes through, and the water system is developed, providing many favorable conditions for mosquito breeding and reproduction. We selected the Jintun town as the center, which had a dengue fever outbreak in August 2017, and radiated out to the surrounding towns and villages to survey the breeding sites. We invited several epidemic prevention personnel from the local health center to assist us in sampling. Water bodies with larva or pupa activity or eggs were selected as sampling sites (Nambunga et al., 2020). Then, GPS was used to locate breeding sites. The distribution of breeding sites is shown in Figure 1.

MATERIALS AND METHODS

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Larva Collection and Identification
Mosquito larva was collected from each breeding site using a standard 350 mL long handle dipper. Each breeding site was dipped four times within a collection period. When the sampling site could not be immersed, siphoning or dumping was used. According to the planned date, each sampling site was sampled five times: July 7, 2019, July 14, 2019, July 23, 2019, July 31, 2019, and August 7, 2019. In the case of thunderstorms or other exceptional circumstances, sampling was postponed. If the water at the sampling site was dry, we located a same-type breeding site nearby. Larvae collected from each breeding site were brought to the Shandong Institute of Parasitic Diseases and classified into species level using standard identification keys (Wilkerson et al., 1990). Quantification of larval density (Villarreal-Treviño et al., 2015) was as follows: D (per dipper) = \[
\frac{\text{the total number of larvae}}{\text{total number of dips}}
\]
where D = mosquito density. The calculation method of mosquito species distribution was based on Kabirul’s method (Bashar et al., 2016): C = \[\frac{n}{N}\times 100\%\], where C = mosquito species distribution, n = number of positive habitats, and N = the total number of habitats.

Identification of the Water Physicochemical Properties
At the site, we measured the water physicochemical properties of each sampling site, including pH, turbidity (NTU: Nephelometric turbidity unit), conductivity (\(\mu S/cm\)), DO (mg/L), AN (mg/L), hardness (mg/L), and temperature (\(^\circ\)C) using a portable multiparameter tester (Palintest Potacheck 2, United Kingdom). WD (cm) and DNH (m) were measured by a graduated string with a lead weight and a tapeline, respectively. After the measurement, at least 200 mL of water was taken from each sampling site and placed into a sterile conical flask. Then, the flask was placed into an icebox and brought back to the laboratory to analyze the water body’s bacterial diversity and richness. The collection time, locations, environmental status, and habitat types were recorded timely and in detail.

Analysis of Bacterial Diversity and Richness
The water samples of each breeding site were centrifuged at 12000 rpm for 10 min in 4°C, and the supernatant was discarded to obtain bacterial precipitation. A Power Soil DNA Isolation kit was used to extract the total bacterial DNA, and the primer sequence 27F (AGRGTITGTYNTGGCTCAG) and 1492R (TASGGHTACCTTGTTASGACTT) were used to amplify the full-length 16S rRNA gene. PCR reaction conditions were 95°C for 5 min, 95°C for 30 s, 50°C for 30 s, and 72°C for 1 min, with a total of 30 cycles and a reaction system of 10\(\mu\)L. After the construction of magnetic beads (MagicPure Size Selection DNA Beads) purification, preparing library, and target gene sequencing, in turn, original reads were corrected to obtain Circular Consensus Sequencing (CCS) (SMRT Link, Version 8.0). The Lima software (version 1.7.0) was used to identify the CCS sequences of different samples by a barcode sequence and a remove chimera (UCHIME, version 8.1), resulting in high-quality CCS sequences. The sequences were clustered at a
similarity level of 97% (USEARCH, version 10.0) (Quast et al., 2013) and the OTU (Operational Taxonomic Units) were filtered with 0.005% of all numbers sequenced as the threshold (Kõljalg et al., 2013). Bacteria chose 16S: Silva databases (Release 132)\(^1\) (Wang et al., 2007) and RDP Classifier (version 2.2)\(^2\) (Larkin et al., 2007), and species were annotated with a confidence threshold of 0.8. Finally, the Alpha index analysis software Mothur (Version V.1.30)\(^3\) was used to analyze the microbial diversity and richness of different habitat samples.

**Statistical Analysis**

All variables were tested by the Shapiro–Wilk test for normality and then analyzed for homogeneity of variance. We used a simple correlation coefficient to analyze the correlation between continuous variables of physicochemical factors. DHN was not considered because there was no logical relationship between variables. Results showed that the variables of physical and chemical factors were not normally distributed and had different variances, and they were not independent of each other. Therefore, it was more reasonable to use a Spearman correlation and multiple linear regression to analyze the correlation between mosquito density and physical and chemical factors (Nikookar et al., 2017). Canonical correspondence analysis (CCA) has been performed to further explore the associations between larval densities and physicochemical factors (Bashar et al., 2016). One-way ANOVA was used to explain whether there were differences in bacterial diversity and richness among different habitat types. Pearson’s correlation model was used to analyze the effects of bacterial diversity and richness on larval densities in breeding sites.

All statistical analyzes were performed using Microsoft Excel 2019 for Windows and SPSS 16 (SPSS Inc., Chicago, IL, United States 2007), with a significance level of 0.05.

**RESULTS**

**Habitat Types**

A total of 62 representative potential habitats were investigated. Out of these, 46 bred larvae. We classified these 46 breeding sites into different habitat types according to the situation on the spot, such as small puddles (6), small water containers (12), paddy fields (7), large water containers (9), irrigation channels (6), and drainage ditches (6). Figures 1, 2 show the locations and habitat types.

**Larval Density and Species Distribution**

A total of 3,291 larvae were collected in all breeding sites, and a total of four genera and six species were identified, including *Cx. p. pallens* (2026/61.56%), *Ae. albopictus* (732/22.24%), *An. sinensis* (147/4.47%), *Cx. tritaeniorhynchus* (319/9.69%), *Cx. bitaeniorhynchus* (13/0.40%), and *Ma. uniformis* (54/1.64%). The larval densities of each habitat type were: small puddles (3.38), small water containers (4.55), paddy fields (2.31), large water containers (4.42), irrigation channels (2.96), and drainage ditches (2.66), respectively (see Table 1). In terms of the distribution of mosquito species, *Cx. p. pallens* was the highest (C = 98.70%), which could be found in all habitat types. It was followed by *Ae. albopictus* (C = 67.83%), *Cx. tritaeniorhynchus* (C = 55.65%), *An. sinensis* (C = 25.65%), and *Ma. uniformis* (C = 10.43%). The distribution of *Cx. bitaeniorhynchus* was the smallest (C = 4.35%). *An. sinensis* was only present in paddy fields and irrigation channels, *Cx. bitaeniorhynchus* was found only in large water containers, and *Ma. uniformis* was found only in irrigation channels.

**The Physical and Chemical Parameters and Bacterial Diversity of Breeding Sites**

Table 2 shows the means and standard deviations of various variables for each habitat. Figure 3 shows the mean and standard error of the OTU, bacterial diversity index (Shannon index), and the richness index (Chao 1) of breeding sites. The Shannon diversity index dilution curve (Figure 4A) showed each sample’s microbial diversity at different sequencing quantities. The curve was flat, indicating that the sequencing quantity was large enough, and the results of OTU clustering accurately reflected the samples’ real situation. The breeding sites with the highest Shannon index were drainage ditches, and the lowest was small water containers. Figure 4B shows the top-ten species at the richness level in each habitat. The bacteria with the highest richness in small puddles was *Hydrogenophaga*. The bacteria with the highest richness in small water containers, paddy fields, and irrigation channels were all *Curvibacter*. *Polyneucleobacter* microbes were most abundant in large water containers. Results of one-way ANOVA showed that the Shannon index (df = 5, F = 5.841, p = 0.001) and Chao 1 (df = 5, F = 10.859, p < 0.01) were different between habitat types.

**Correlations Between Variables**

Table 3 shows the correlation coefficients between the variables of each physicochemical factor. Among the 36 correlation coefficients, 7 (19.44%) were statistically significant, indicating a non-random correlation between the variables with physicochemical properties. pH was negatively correlated with turbidity (r = −0.442, p = 0.002) and conductivity (r = −0.502, p < 0.01). A significant negative and positive correlation existed between hardness with turbidity (r = −0.325, p = 0.028) and with AN (r = 0.339, p = 0.021), respectively. Turbidity was positively correlated with conductivity (r = 0.357, p = 0.015) and negatively correlated with AN (r = −0.375, p = 0.01). Temperature was positively correlated with WD (r = −0.407, p = 0.005).

**Relationship Between Larval Density and Variables**

Spearman correlation analysis (Table 4) shows the correlations between larval densities and physical and chemical parameters. In the multiple linear regression model analysis, the larval densities of various types were taken as dependent variables, and then 9 physical and chemical factors, as independent variables.

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\(^1\)http://www.arb-silva.de
\(^2\)http://sourceforge.net/projects/rdpclassifier/
\(^3\)http://www.mothur.org/
respectively, were incorporated into the model. The variables were fitted into the regression equation according to the stepwise selection method, and the independent variables with a large contribution to the regression equation were reserved ($p < 0.05$) were retained. According to the results of multiple linear regression analyzes (Table 5), Cx. p. pallens ($F = 18.30, p < 0.001$) required low DNH, whereas An. sinensis ($F = 72.388, p < 0.001$) required high DNH. Cx. tritaeniorhynchus ($F = 33.498, p < 0.001$). Our results showed that Cx. bitaeniorhynchus did not fit the regression equation as the $P$ value was greater than 0.05.

Table 6 showed that the density of Cx. p. pallens negatively correlated with the Shannon index ($r = -0.430, p = 0.018$)
and the richness index \( r = -0.644, p < 0.001 \). The density of \textit{Ae. albopictus} negatively correlated with the bacterial diversity index \( r = -0.411, p = 0.024 \) and the richness index \( r = -0.572, p = 0.001 \). \textit{Cx. tritaeniorhynchus} positively correlated with the Shannon index \( r = 0.493, p = 0.006 \), while other mosquito species did not significantly correlate with bacterial diversity. Figure 5 visually showed the correlation between larval density and variables based on the Canonical correspondence analysis (CCA).

## DISCUSSION

Mosquito problems have been categorized as medical care problems and environmental issues with the progress of human society, urban sprawl, and intensification of environmental changes (Ahmed et al., 2007). Sole dependence on insecticides can no longer completely eliminate the hazards that mosquitoes had brought to us. Luckily, LSM is environment friendly and does not lead governments and inhabitants to too much economic burden and time cost. These could be rapidly realized utilizing eliminating breeding sites or using chemical or biological methods. However, developing efficient and safe strategies required a scientific understanding of the control area’s larval density and environmental factors.

In recent years, due to the vigorous development in modern agriculture, the mode of agriculture in Jiaxiang County has changed. For example, paddy fields have been replaced by citrus orchards, which directly resulted in the change of some breeding sites’ environment. We suspect that this may be a factor that led to the failure to obtain mosquitoes such as \textit{An. dirus}, which prefer open water (Service, 1991; Norris, 2004). In addition, agricultural activities such as the use of pesticides and fertilizers changed the richness of microbota and algae in the water, which are the main food sources for larvae (Noori et al., 2016). This disturbed selecting suitable sites for oviposition and incubation by specific mosquitoes.

Most of the microbiota in the breeding sites form parasitic or symbiotic relationships with the larvae, and their impact on the larvae can be classified in the following three aspects: regulating the internal and external environment and immune response of the mosquito host, promoting the developments and growth of larvae, and causing pathogenicity to larvae such as infection of the filamentous fungi \textit{Beauveria bassiana} and \textit{Bacillus thuringiensis} (Bti). In addition, studies have shown that bacterial diversity might be related to larval density (Dida et al., 2018). Therefore, we used molecular biology methods, namely the 16S rRNA sequencing technology, to sequence and analyze the bacterial diversity and richness of breeding sites. Results showed that different habitats have large differences in bacterial diversity and richness, even in the same habitat, because bacterial diversity and richness in water are always dynamic and affected by their environment, such as light, temperature, vegetation cover, and human activities. In general, \textit{Cx. p. pallens} and \textit{Ae. albopictus}, the dominant species in Jiaxiang County, both showed lower tolerance to bacteria, which also seems to explain the lower diversity of intestinal symbiotic bacteria in these two species (Coon et al., 2016). There was no statistical correlation between the density of other species and the bacterial diversity, which we hypothesized might be due to the small
FIGURE 4 | Alpha diversity analysis: (A) Shannon diversity index dilution curve. It shows the microbial diversity of each sample at different sequencing quantities. The curve is flat, indicating that the sequencing quantity is large enough, and the results of Shannon index accurately reflected the real situation of the samples. (B) The top-ten species at the richness level in each habitat. DD, drainage ditches; IC, irrigation channels; LC, large water containers; PF, paddy fields; SC, small water containers; SP, small puddles.

number of samples we tested from water bodies or the large variation in the diversity index of different habitat types. *Cx. p. pallens* has a wide range of fitness for physical and chemical properties such as pH, hardness, turbidity, and AN, which is consistent with the results of Nikookar et al., (Nikookar et al., 2017). In other words, *Cx. p. pallens* have no strict requirements
regarding the environment they breed in, and they can even breed in less hospitable environments. This seems to explain the dominance of Cx. p. pallens in the less sanitary areas of northern China. In addition, Gabriel et al., have shown that Culex (especially Cx. p. pallens and Cx. tritaeniorhynchus) are widely distributed in a variety of water bodies, providing intuitive evidence that they are highly adaptable to different habitats. Urbanization, human activities, and other factors have affected the breeding environment's physicochemical properties and water bodies (Ndaruga et al., 2004). Despite this, the density of Culex in Jiaxiang County remains high. Previous reports and our study give a reasonable explanation for this phenomenon.

**Ae. albopictus** is the main vector of the dengue virus. Our study showed that **Ae. albopictus** density was negatively correlated

### TABLE 3 | Correlation between variables with physicochemical property.

|      | DO     | pH     | Hardness | Turbidity | Conductivity | Temperature | AN   | WD   |
|------|--------|--------|----------|-----------|--------------|-------------|------|------|
| Y    |      -0.286 | 0.213  | 1        | -0.212    | 0.357*       | 1            |      |      |

**Ma. uniformis**

|      |      0.118 | -0.442* | -0.325* | 0.018     | 0.065        | 5.494       | 1    |      |

**Cx. tritaeniorhynchus**

|      |      0.030 | -0.502** | -0.212   | 0.357*     | 1            | 0.079       |      |      |

**An. sinensis**

|      |      0.104 | -0.113   | -0.198   | 0.179      | 0.088        | 0.285       | 0.231| 0.024|

**Ae. albopictus**

|      |      0.270 | 0.339*   | -0.375*  | -0.179     | -0.188       | 0.265       | 0.075| 0.394**|

**Table 4** Spearman correlation coefficient between larval density and physicochemical characteristics of larval habitats.

| Cx. p. pallen | Ae. albopictus | An. sinensis | Cx. tritaeniorhynchus | Cx. bitaeniorhynchus | Ma. uniformis |
|---------------|----------------|--------------|-----------------------|----------------------|---------------|
| r             | 0.033          | 0.034        | -0.062                | -0.061               | 0.147         |
| p             | <0.001         | <0.001       | <0.001                | <0.001               | 0.001         |

**Table 5** The multiple regression equation shows the contribution rate of each variable to mosquito density.

**Table 6** Pearson correlation coefficient between larval density and bacterial diversity of larval habitats.
with WD. This may be because they prefer to lay eggs in small water containers such as junked tires, plastic bottles, and surface-gathered water and have strong adaptability to low water levels (Li et al., 2014; Petric et al., 2014). There is a negative correlation between \textit{Ae. albopictus} and DNH. In other words, the farther the breeding site is from the house, the lower the density of \textit{Ae. albopictus}, and vice versa. This characteristic is observed in all species except \textit{An. sinensis}. WHO has reported that \textit{Ae. albopictus} prefers to lay eggs in small containers having stagnant water in people's yards or houses (World Health Organization, 1983). Based on such characteristics, dumping stagnant water to reduce the incidence of dengue fever by reducing the BI could be an effective and environment-friendly method. Moreover, \textit{Ae. albopictus}, just like \textit{Cx. p. pallens} showed a positive correlation with turbidity and temperature, while a negative correlation with pH and AN. They showed a wide range of adaptation to the abovementioned physical and chemical properties. As Thavara and Bashar reported, \textit{Ae. albopictus} preferred to inhabit human-made habitats inside and outside houses, where the habitats' physical and chemical properties were more fluctuating (Thavara et al., 2004).

The increase in salt compounds and turbidity had a significant effect on the density of \textit{An. sinensis}, and they showed low tolerance to AN levels. Kenawa found a strong positive correlation between its density and dissolved oxygen (El-Ghani et al., 2012), which is consistent with our results ($r = 0.312$, $p = 0.035$). In comparison with \textit{Aedes} and \textit{Culex}, \textit{Anopheles} mosquitoes were more sensitive to other physical and chemical changes in water (El-Ghani et al., 2012). Minakawa et al. (2005) suggested that \textit{Anopheles} preferred sunny open waters such as lotus ponds, paddy fields, and irrigation channels where breeding sites are far from houses. Our results also confirm that \textit{Anopheles} mosquito density is positively correlated with DNH. We suspect that this may be related to the strong long-range flight ability of \textit{An. sinensis}. The rice-growing areas and lotus ponds in Jiaxiang County are widely distributed, which provides the conditions for the breeding of \textit{An. sinensis} in the area. Applying agricultural insecticides seem to be the most effective way to control \textit{An. sinensis}. However, the epidemic prevention department should instruct farmers in the region to spray insecticides scientifically and rationally to slow the development of mosquito resistance. On the basis of the existing rice-planting pattern in Jiaxiang County, humid irrigation replacing the current irrigation regime can effectively eliminate the breeding sites for irrigation channels, which is also an effective method to control \textit{An. Sinensis} (Tusting et al., 2013). Minakawa found that \textit{Anopheles} in western Kenya were not found in sunny open water and preferred lakeside swamps. This difference suggests that even mosquitoes from

![FIGURE 5](image-url)
the same genus may show differences in habitat selection. The primary reason for this phenomenon was geographical isolation and ecological differences (Varela and Yadav, 2020), suggesting that we should first understand their habits and adapt to local conditions to prevent and control larvae.

*Culex tritaeniorhynchus* is the primary medium carrying the Japanese encephalitis virus and the dominant mosquito species in Shandong Province, especially near the South-to-North Water Diversion Project. In terms of distribution alone, they are similar to *Cx. p. pallens*, as both of them, were found in all habitat types. However, the density of *Culex tritaeniorhynchus* that we collected was significantly lower than that of *Cx. p. pallens*. They can survive in deep water with low oxygen content, so the density and WD showed a strong positive correlation. From the results of the regression equation, it is evident that AN and WD had the highest density contribution to *Culex tritaeniorhynchus*, which is the same as the research results of Wang et al., (Wang et al., 2020). *Cx. bitaeniorhynchus* and *Ma. uniformis* showed a low distribution, and they were only found in large water containers and irrigation channels, respectively. Combined with previous surveys, we found that the density of these two species had been low in Jiaxiang County, which did not bring trouble or danger to humans. Therefore, this is not discussed in this article.

Although the CCA results were consistent with the Spearman, it may be able to predict larval density in a much more intuitive way. We can intuitively receive such results from the biplot: the larval density of *Ae. albopictus* and *Cx. p. pallens* can be better predicted by low WD, DNH, and high temperature. Alternatively, high DNH and DO may be more useful in predicting larval density of *An. sinensis* than other variables. The larval density of *Ma. uniforis* can be affected by high WD and low temperature, while the density of *Cx. bitaeniorhynchus* and *Cx. tritaeniorhynchus* can hardly be predicted from the results of CCA. The CCA biplot indicated that *Anopheles, Aedes, Culex*, and *Mansonii* were plotted in different axis which may explain different requirement for different variables (Bashar et al., 2016).

There are several aspects that need to be improved in this study. We did not conduct a survey of algae, protists, and rotifers in breeding grounds. However, protists are the main source of nutrition for larvae (Blaustein and Chase, 2007; Addicott, 2008), and *Coelastrum, Scenedesmus, Selenastrum*, and *Tetraallanto* that are algae for *Aedes and Culex* are difficult to digest, which can reduce the survival rate of larvae (Howland, 1930). Microbial diversity and physicochemical parameters of water bodies change dynamically, and a large number of survey results are needed to develop a scientific and rigorous model to predict larval density. Thus, our sample number is far from enough.

### CONCLUSION

This research showed that the physicochemical parameters and bacterial diversity of different habitat types were significantly different. Even for the same habitat type, the physicochemical parameters varied greatly due to different environments. The variable that contributed the most to the density of *Cx. p. pallens* was DNH. Parameters such as DNH, AN, pH, and temperature contributed significantly to the density of *Ae. albopictus*. These two kinds of mosquitoes are distributed in almost all kinds of habitats in Jiaxiang County, and their density is also the highest. This suggests that we should be vigilant against the control of mosquito-borne viruses in this area, especially the dengue virus carried by *Ae. albopictus*.

### DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/supplementary material, further inquiries can be directed to the corresponding author.

### AUTHOR CONTRIBUTIONS

MQG and HYW designed the study. HFW and HWW participated in field surveys with support from PC. HYW extracted the data and carried out the data analysis with support from YW. HYW and YW wrote and reviewed the article. All authors read and approved the final manuscript.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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