Bacterial abundance and community composition in green, brown and red water from intensive catfish (Clarias sp.) culture ponds in Yogyakarta, Indonesia

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Abstract. Wijayanti KAN, Istiqomah I, Murwantoko. 2021. Bacterial abundance and community composition in green, brown and red water from intensive Catfish (Clarias sp.) culture ponds in Yogyakarta, Indonesia. Biodiversitas 22: 3677-3684. Catfish (Clarias sp.) is an important aquaculture commodity in Indonesia and cultured in an intensive system. Microorganisms play an important role in maintaining water quality of aquaculture system. The objective of this study was to determine the bacterial abundance and community composition of green, brown and red water collected from intensive catfish culture ponds in Yogyakarta using next-generation sequencing method. The water samples were collected from intensive catfish culture ponds with different colors, namely green, brown and red ponds located in Yogyakarta. The DNA from water samples was extracted using DNA extraction kit and used as template for 16S rRNA amplification. The V3-V4 hypervariable regions of the 16S rRNA gene were amplified apply for next-generation sequencing technology. This study could explore effectively the bacterial community in water samples. The bacterial communities in this catfish culture water showed higher bacterial richness compared to the other aquaculture system. The diversity of the green, brown and red catfish culture water ponds was similar with the number OTUs of the green, brown and red water samples, which were 1269; 1387 and 1323 OTUs respectively. The 694 OTUs (34.42%) were common core microbiomes in all catfish culture ponds, the 212 OTUs (10.51%) are present on green and brown water ponds, the 182 OTUs (9.02%) were on green and red water ponds, and the 183 OTUs (9.07%) were present on green and brown water ponds. However, the composition of the bacterial community was different. The most dominant phylum in green and brown water ponds was Proteobacteria with relative abundance in green water and brown water 71.6% and 47.0% respectively, whereas, the most dominant phylum in red water was Firmicutes (29.5%). The dominance of Firmicutes phylum in red water ponds may be caused by application of probiotic bacteria, the high organic content, and low oxygen concentration.

Keywords: bacteria; composition; diversity; high-throughput sequencing, catfish culture

INTRODUCTION

The fisheries and aquaculture sector significantly expanded in the past few decades. Since the 1990s, a rise of 527% in global aquaculture production has been observed from 1990 to 2018. While capture fisheries production has been relatively stable, with some growth essentially concerning inland capture with a rise in global capture fisheries production from 1990 to 2018 is 14%. World aquaculture production of farmed aquatic animals has been dominated by Asia, with 89% share in the last two decades with major producing countries is China, India, Indonesia, and Vietnam (FAO 2020).

Microorganisms play an important role in the aquaculture system. The microbial communities associated with the fish and their environment can improve water quality, reduce the abundance of fish pathogenic bacteria and ultimately improve fish survival (Beniston-Tilla et al. 2016). The importance of microbial diversity and its role in maintaining fish health in aquaculture systems has been increasingly recognized in recent years. However, there is still a major knowledge gap regarding the ecology, composition and dynamics of microbial plankton assemblages during fish production. Besides having direct effects on fish health and quality in aquaculture settings, microbial communities also influence fundamental processes such as nutrient cycling and water purification (Duarte et al. 2019). The healthy microbiome is mainly a resistant, resilient, and healthy functional core composed of a blend of articular microbial gene families, metabolic modules, and regulatory pathways capable of resisting any agitation due to internal or external factors (Rajeev et al. 2021).

Controlling the microorganisms associated with aquaculture systems (i.e. the aquaculture microbiome) has always been essential in high-intensity rearing of fish (Ditttman et al. 2017). The microbial-based systems represent one of the most viable strategies to achieve sustainable aquaculture. These systems are based on the promotion of microbial proliferation, either heterotrophic or autotrophic microorganisms. These microbes are expected to use, recycle and transform the excess of nutrients from feces, dead organisms, un Consumed food, and diverse metabolites into biomass, which the cultured organisms would further consume. (Martínez-Córdova 2014) The implementation of microbiome-based products practiced by the specific functional groups of the...
The aquaculture microbiome is enriched by adding, for example, carbon-rich substrates. These applications of targeted microbiome-based products containing a seeding microbial assemblage to aid the heterotrophic assimilation of inorganic nitrogen and/or the nitrification process are now a common practice in intensive tropical pond-based aquaculture systems (Dittman et al. 2017). The ability of organisms to adapt to a particular habitat or living conditions mainly depends on their phenotypic plasticity. Therefore, the treatment approaches are now targeted towards modulating the aquatic species’ microbiome and using them as beneficial partners to overcome the challenges confronted by the aquaculture sector (Rajeev et al. 2021).

Microbial communities in natural aquatic environments respond rapidly to changes in their immediate environment. These changes may be subtle and may manifest themselves as regulating certain metabolic pathways, or they may cause changes to the overall microbial community composition and functionality (Benzton-Tilla et al. 2016). The fundamental baseline information concerning the microbial dynamics of these systems and how ecological interactions can be used to modulate microbial assemblages (Duarte et al. 2019). The high-throughput sequencing (HTS) technologies have been applied as tool microbial ecology of these systems. The HTS has been used to characterize both the healthy and diseased aquaculture microbiome, to investigate the microbial community composition in recirculating aquaculture systems (Benzton-Tilla et al. 2016), investigated temporal variations of bacterioplankton and the seasonal dynamics of bacterial and microeukaryotic plankton communities e.g. in a semi-intensive European seabass aquaculture system (Martins et al. 2018; Duarte et al. 2019). The seasonal studies showed that changes in environmental variables influence the overall microbial communities. (Roquigny et al. 2021).

Catfish is one of the important aquaculture commodities in Indonesia. In 2014, national catfish production was 679,379 tons which increased to 1,280,099 tons in 2018. This means that national catfish production from 2014 to 2018 increased by 88.4% (DJPB, 2020). The catfish have been intensively cultured in the Yogyakarta area. The farmers found the condition of water in ponds showed different colors. They believed that this different color gave different productivity. So, the microbiome from those different color ponds is important to be explored. The present study aimed to determine the bacterial abundance and community composition from green, brown and red water from intensive catfish culture ponds using the next-generation sequencing method. This is perhaps, the first study using high-throughput sequencing to analyze the bacterial community on freshwater aquaculture ponds in Indonesia.

**MATERIALS AND METHODS**

**Sample collection**
This research was conducted in intensive catfish (Clarias sp) culture ponds which have different colors, namely green, brown and red ponds in Sleman Regency of Yogyakarta Indonesia. The green water catfish culture pond is located in Kalasan District as a rectangular earthen pond with an area of 120 m² with a stocking density of 60 fish/m². The brown water pond is located in Prambanan District as concrete wall construction with earthen bottom on a rectangular shape with an area of 30 m² with a stocking density of 100 fish/m². The red culture pond is located in Berbah as a round tarpaulin pond with an area of 7 m² stocked with catfish at a density of 200 fish/m². The water from the ponds was collected by taking one liter of water from the middle of the water column of pond. All the water samples were placed in a cool box, transported to the Fish Health and Environment Laboratory at Universitas Gadjah Mada (Yogyakarta, Indonesia), and stored in a refrigerator at 4°C for further analysis.

**DNA extraction**
Total 30 mL water samples were centrifuged at 3000xg for five minutes and settled pellets were then collected. The pellets were extracted their DNA using Genomic DNA Extraction Mini Kit (Favorprep™ Favorgen) following the manual protocol. The quality and concentration of the total DNA were estimated through electrophoresis on 1% agarose gel and NanoDrop MN913A MaestroNano Pro (MaestroGen Inc., Hsinchu, Taiwan). The genomic DNA with a concentration of 50 ng/µL and DNA purity (A260/280) within the range of 1.8-2.0 were used for amplicon sequencing.

**Sequencing preparation and bioinformatics analysis**
Equal volumes of the total genomic DNA from green, brown and red water ponds proceeded for NGS. The V3-V4 hypervariable regions of the 16S rRNA gene were amplified using the forward primer 341F (5’- CCTAYGGGRBGCASCAG-3’) and the reverse primer 806R (5’-GGACTACNNGGGGTATCTAAT-3’) (Yu et al. 2005).

The DNA amplifications were performed using Phusion® High-Fidelity PCR Master Mix (New England Biolabs) in accordance with the manufacturer’s instructions. The PCR products were then mixed with 1x loading buffer (containing SYB green) at the same volume, detected on 2% agarose gel, and purified with a GeneJETTM gel extraction kit (Thermo Fisher Scientific). Sequencing libraries were generated using Ion Plus Fragment Library Kit (Thermo Fisher Scientific) in accordance with the manufacturer’s recommendations. Their quality was assessed with a Qubit 2.0 fluorometer (Thermo Fisher Scientific). The libraries were then sequenced on an Ion S5TM XL platform (Thermo Fisher Scientific), and 400 bp to 600 bp single-end reads were generated. This PCR amplification and amplicon sequencing were carried out by a service company (NovogeneAIT) in Singapore.

Single-end reads were further assigned to the samples based on their unique barcodes and truncated by cutting off the barcodes and primer sequences. Raw reads were filtered under specific filtering conditions to obtain high-quality clean reads in accordance with the Cutadapt
quality control process (Martin 2011). Then, chimera was removed by comparing the reads with the Silva 132 reference (Quast et al. 2013; https://www.arb-silva.de/) via the UCHIME algorithm (Edgar et al. 2011), and clean reads were finally obtained. Sequences with more than 97% similarity were clustered into the same operational taxonomic units (OTUs) by using the QIIME software version 1.9.1 (Caporaso et al. 2010) in accordance with the UCLUST method (Edgar 2010). The representative sequence of each OTU was screened for further annotation. Each representative sequence was then assigned using the Silva 132 reference based on the Mothur algorithm to annotate taxonomic information.

Alignment of the OTU representative sequences was performed using the MUSCLE software (version 3.8.31) to construct phylogenetic metrics such as Unifrac, which aimed to study phylogenetic relationship of different OTUs and the difference in the dominant species in various samples (Edgar 2004). Data normalization of OTUs abundance was done using a standard sequence number corresponding to the sample with the least sequences. This output normalized data were further used for rarefaction, rank abundance, bacterial community bar plot, bacterial community heatmap, principal coordinate analysis (PCoA), and Weighted Unifrac distance were determined and performed with QIIME (version 1.9.1) and then displayed with R software version 2.15.3 (Ihaka and Gentleman 1996). Alpha diversity is applied in analyzing complexity of species diversity for a sample through Chao1, Shannon, Simpson indices. All these indices in our samples were calculated with QIIME (Version 1.7.0) and displayed with R software (Version 2.15.3).

RESULTS AND DISCUSSION

Sequence analysis

High-throughput sequencing was performed to analyze the 16S rRNA gene of bacteria present in the different watercolor from pond of catfish (Clarias sp) culture. In the 16S rRNA gene sequencing, 440,433 effective sequences were obtained from three samples after chimera filtration. The number of effective sequences in the green, brown and red water samples were 145,588; 142,077 and 152,768 (126,518; 121,109 and 127,650) respectively. The rarefaction curve which tended to reach flat curve (Figure 1a) indicated that the bacteria were effectively explored and only the scarce bacteria remained. The rarefaction curve of each sample revealed that the number of the observed species in equal, which is also supported by the rank abundance curve (Figure 1b).

All the effective sequences with 97% similarity were then grouped into the same, and this study could identify 2016 OTU. The number of OTUs was equal among three different color pond water, with the number OTUs of the green, brown and red water samples were 1269; 1387, and 1323 OTUs respectively. The bacterial community among three water samples was relatively similar, as indicated by the range or difference from the maximum-minimum value indexes are small/narrow. The bacterial/OTU richness as indicated by Chao1 were ranged from 1475.26 to 1331.71. The evenness bacterial/OTU as indicated by Shannon index was ranged from 7.35 to 6.50. Whereas the bacterial/OTU dominance by Simpson’s index were ranged from 0.981 to 0.958 (Table 1).

Table 1. Alpha diversity indices of the bacteria from the green water pond (green12), brown water pond (brown23) and red water pond (red33).

<table>
<thead>
<tr>
<th>Sample</th>
<th>Chao1</th>
<th>Shannon</th>
<th>Simpson</th>
</tr>
</thead>
<tbody>
<tr>
<td>Green12</td>
<td>1331.71</td>
<td>6.50</td>
<td>0.958</td>
</tr>
<tr>
<td>Brown23</td>
<td>1475.26</td>
<td>7.64</td>
<td>0.987</td>
</tr>
<tr>
<td>Red33</td>
<td>1378.07</td>
<td>7.35</td>
<td>0.981</td>
</tr>
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</table>

Figure 1. Rarefaction curves (A) and rank abundance curve (B) resulted from the 16S rRNA gene sequencing of the green (green12), brown (brown23) and red (red33) water pond.
Among 2016 of identified OTUs, the 694 OTUs (34.42%) were common core microbiomes in the catfish culture pond as overlapped to all three color samples. The 212 OTUs (10.51%) are present on green and brown water ponds, the 182 OTUs (9.02%) are present on green and red water ponds, and the 183 OTUs (9.07%) are present on green and brown water ponds. The rest OTUs were only present in the green, brown and red water samples were 181; 298 and 182 OTUs respectively (Figure 2).

**Bacterial diversity and abundance analysis**

All the sequences obtained from the three samples were assigned from phylum to genus levels. In particular, the three most dominant phyla were identified as Proteobacteria, Firmicutes and Bacteroidetes possessed the relative abundance as 88.6%, 82.2%, and 75.1% of the green, brown and red water respectively. The most dominant phylum was different among samples with the green water was dominated by Proteobacteria (71.6%), Bacteroidetes (10.6%), Firmicutes (6.4%), brown water dominated by Proteobacteria (47.0%), Bacteroidetes (24.1%), Firmicutes (11.0%), whereas red water dominated by Firmicutes (29.5%), Proteobacteria (26.9%), Bacteroidetes (18.6%) (Figure 3).

In green water, the Betaproteobacteria was the most abundant class with the relative abundance of 41.9% followed by gammaproteobacteria with a relative abundance of 21.6%. In brown water, the Betaproteobacteria was the most abundant class with the relative abundance of 15.9% followed by alphaproteobacteria with a relative abundance of 14.3% The Clostridia is most abundant class in redwater with relative abundance of 25.5% followed by bacteroidia with a relative abundance of 14.0% (Figure 4).

At the family level, the most abundant genus of the green, brown and red water was Rhodocyclaceae, Comamonadaceae, and Christensenellaceae respectively. The Rhodocyclaceae was found in the green, brown and red water at relative abundance of 31.7%, 6.4%, and 1.3% respectively. The Comamonadaceae and Saprospiraceae were found in significant amounts in the green and brown water. The Comamonadaceae was found at relative abundance of 7.7% and 6.3% on the brown and green water respectively, while Saprospiraceae was found at relative abundance of 5.5% and 5.5% respectively. The Methylococcaceae was found in red water found at relative abundance of 9.7% and in other water only found at relative abundance of less than 1% (Figure 4).

The heatmap analysis revealed that the bacterial community composition in the water ponds differed among different color water, as indicated by different dominant genera. The genera of Thiothrix, Heliscomenobacter, Peredibacter, Competibacter, Thaurea, Methyloparacoccus, Dechloromonas were found dominant in green water. The genera of Bacterriodes, Rhodobacter, Propionivibrio, Desulfovomicrobium, Novosphingobium, and Acidmanibacter were dominant in brown water. The genera of Paludibacter, Calidithrix, Microbacter, Christensenellaceae, Desulfobulbus were found dominant in redwater (Figure 5).
Figure 4. Relative abundance of bacterial communities at the ordo level of the green water pond (green12), brown water pond (brown23) and red water pond (red33)

Figure 5. Abundance heatmap of bacterial communities at the genus level in the green water pond (green12), brown water pond (brown23) and red water pond (red33). The absolute value of the scale represents the distance between the raw score and the mean of the standard deviation. The value is negative when the raw score is below the mean, and vice versa.

Similarity analysis of bacterial composition

The similarity between bacterial communities from different water was further analyzed through PCoA. The PCoA plot showed that the green, brown and red water were separated from each other (Figure 6). The relationship analysis using the unweighted pair group method with arithmetic mean (UPGMA) method showed that the green and brown water was closer than the red water (Figure 7).

Discussion

Microorganisms play important role in aquaculture system. The microbial communities can improve water quality, reduce the abundance of fish pathogenic bacteria, improve fish survival (Benzon-Tilla et al. 2016), and play fundamental processes such as nutrient cycling and water purification (Duarte et al. 2019). Based on the study on the bacterial community structure and functional changes due to Chilean salmon aquaculture, the bacterial shifts can be used as indicators of aquaculture perturbations (Hornick and Buschmann 2018). However, there is still a major knowledge gap regarding the ecology, composition, and dynamics of microbial plankton assemblages during fish production (Duarte et al. 2019). The fish farmers in Yogyakarta have found different colors of water media of intensive catfish culture. They believed that this watercolor pond has different fish productivity. This study explores the bacterial communities from the green, brown and red water from catfish culture ponds using high throughput sequencing (HTS) or new generation sequencing (NGS) technology.

This study’s bacterial communities in catfish culture water showed higher bacterial richness than the other aquaculture system. The bacterial richness as Chaol in this study was ranged from 1475.26 to 1331.71 (average 1394.9). In contrast, the bacterial community in pond of mina padi integrated culture was 1169.9 (Herlambang et al. 2021), the semi-intensive aquaculture for European sea bass (Dicentrarchus labrax) was ranged from 216.53 to 1148.86 (Duarte et al. 2019), the sediments from Chilean salmon aquaculture sites were ranged from 275.4 to 331.9 (Hornick and Buschmann 2018). The intensive catfish culture has put a lot of input such as the high nutrient of feed, so that the nutrient content of water is also high, giving possibility for many microorganisms to grow. The catfish culture in Yogyakarta is land-based aquaculture so that only limited water and nutrients flow out to the open system.

Figure 6. Principal coordinate analysis (PCoA) of the green water pond (green12), brown water pond (brown23), and red water pond (red33) based on Weighted Unifrac distance
Studies showed different dominant phyla characterized by fermentation, assimilation, and metabolism of sugar/hemicellulose levels, available moisture, and nutrient content, such as a rapid response to dissolved organic carbon (Newton et al. 2011). Ecologically, Gammaproteobacteria are an extremely polyphyletic class of Firmicutes, lack aerobic respiration, obligate anaerobes, most of their species are Gram-positive. They can be found in decaying vegetation, marine sediment, soil, and intestinal tract of humans and other vertebrates. The spores of sulfite-reducing anaerobes (Clostridia) are widespread in the environment. The dominance of this Firmicutes phylum may be caused by application of probiotic bacteria, the high organic content, and low oxygen concentration in red water ponds.

At the class level, Betaproteobacteria was the most abundant in green and brown water at relative abundance of 41.9% and 15.9% respectively. The second abundant green and brown water class was different as the Gammaproteobacteria (21.6%) and Alphaproteobacteria (14.3%), respectively. The minapadi system’s water study showed that those three classes as Betaproteobacteria, Alphaproteobacteria, and Gammaproteobacteria were the three most dominant classes (Herlambang et al. 2021). However, the other studies showed different dominant classes in several aquaculture systems. The Alphaproteobacteria (42.60±14.83%), Flavobacteria (23.70±10.04%), Gammaproteobacteria (15.06±8.09%) were the most abundant classes a semi-intensive European seabass (Dicentrarchus labrax) aquaculture system in Ria de Aveiro estuarine lagoon, Portugal during 2012 (Martin et al. 2018). The sampling events throughout the year of 2014 from aquaculture system showed that the most abundant bacterial classes were Gammaproteobacteria (41.73±4.71%), Flavobacteria (20.79±7.26%), Alphaproteobacteria, (13.55±5.26%) (Duarte 2019). The seasonal survey in a sea bass farm in France showed that Gammaproteobacteria, Alphaproteobacteria, and Deltaproteobacteria were the three main classes composing the Proteobacteria phylum with 55.88%, 37.20% and 6.82%, respectively (Roquigny et al. 2020). Those results indicated that at class level, the bacterial composition was more dynamic.

Betaproteobacteria is a typical class of bacteria found in various freshwater habitats (Hahn 2006). The presence and abundance of Betaproteobacteria in freshwater are associated with its ability to respond quickly to changes in nutrient content, such as a rapid response to dissolved organic carbon (Newton et al. 2006). Several Alphaproteobacteria members are involved in the nitrogen cycle (Newton et al. 2011). Ecologically, Gammaproteobacteria helps to modulate excess nitrate (Fernandes et al. 2014).

The present study has effectively explored the bacterial community of the green, brown, and red water from catfish intensive culture ponds. Those three types of ponds have similar bacterial richness but different in bacterial composition from phyla level to genera level. The anthropogenic activity affected the composition of bacteria.
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