Identification of agents causing vibriosis in *Litopenaeus vannamei* shrimp culture in Kendal, Central Java, Indonesia and application of microalgae *Dunaliella salina* and *Tetraselmis chui* as bio-control agents against vibriosis

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**Abstract.** Aquaculture of Vannime shrimp at Kendal, Central Java, Indonesia has been using an intensive system, which could provoke the degradation of water quality in the pond. One of the emerging effects due to such degradation becoming threat for shrimp aquaculture is the loss caused by vibriosis disease. This study aimed to identify the presence of vibriosis-causing bacteria in shrimp pond at Kaliwungu, Kendal, Central Java and use the microalgae *Dunaliella salina* and *Tetraselmis chui* as biocontrol agents against Vibriosis. The Vibriosis-causing bacteria were isolated from the hepatopancreas and tails of ten shrimps showing clinical signs of vibriosis infection, cultured on the Thiosulfate Citrate Bile Salt Sucrose media. Out of 20 isolates, three isolates were selected for further gene sequence identification of 16rRNA using the Repetitive Sequence-based Polymerase Chain Reaction (rep-PCR) and were analyzed with BLAST. Universal primer 27F (5'AGAGTTTGATCMTGGCTCAG -3') and 1492R (5'TACGGTTAACCTTGTTACGACTT-3') were selected as the primers to identify the isolated bacteria. Results demonstrated that the three isolated bacteria were positive as vibriosis-causing agent in shrimp, which were U1H1 AND U4H1, identified as *Vibrio alginolyticus*, and U1E1 as *V. harveyi*. The use of microalgae as biocontrols was performed. Forty-five shrimps infected by vibrio reared during 21 days and feed with *D. salina* and *T. chui* showed a decreased of bacteria amount, which were counted by Total Plate Count method. The result indicated that the microalgae was capable to produce an antibacterial compounds against vibrio.

**Key Words:** aquaculture, disease, pond water quality, bacteriosis, clinical symptom.

**Introduction.** Community trend towards the need of fisheries products is constantly increasing, thus, the aquaculture activities are highly developed, and one of them is the Vannime shrimp culture. This species is a fresh water commodity. The production of Vannime in Central Java reached 411.729 tons in 2014. From 2010 to 2014, the production of this commodity increased with 20.49% (LAKIP-Directory General of Aquaculture Fisheries, 2014). The augmentation of Vannime production at Central Java has been generated by the application of intensive systems characterized by spreading high quantity of breed stocks and the usage of 100% of artificial feeds. The application of intensive systems decreases the water quality in ponds making the Vannime shrimp
susceptible to diseases caused by parasites, virus and bacteria. The mortality of shrimp due to vibriosis infection might reach 80 to 100% in the larval, post larval, juvenile, and adult stages within 1 to 3 days. Case of mass mortality in shrimp aquaculture occurring across every region in Indonesia in the 1900s, including Central Java, had made great loss for most of the companies.

*Vibrio* sp. has been widely reported to be one of the main pathogenic bacteria causing high mortality rate in shrimp aquaculture (Haldar et al 2011; Heenatigala & Fernando 2016; Solidum et al 2016). The presence of this pathogenic bacteria itself is a part of natural microflora contributing to 60% of total bacteria population as mentioned by Heenatigala & Fernando (2016). As opportunistic organism, the infection of *Vibrios* to the cultivated organisms occurs when the environmental condition degrades affecting the health condition of the cultivated organism (Kumaran & Citarasu 2016). Bacterial disease outbreak provoked by *Vibrios* has become constraint on the sustainable production of shrimp (Manilal et al 2010). In order to suppress the outbreak, marine secondary metabolites have been mentioned as promising resources to further develop, especially from microalgae. It has been reported that the extracts of microalgae could prevent the bacterial infections (Cadiz et al 2016; Dash et al 2017).

Despite of the intensive shrimp farming activities in order to fulfill markets demand, infectious disease caused by bacteria remains the main problem that still happens present days. As a consequence, it is crucial to carry out a study that evaluates and identifies the vibriosis-causing bacteria as an update and also serves as basic information prior to further treatment. It is meant to apply the suitable and appropriate treatment to overcome the infection. The objectives of this study were to identify the agents causing vibriosis in the cultivated *Litopenaeus vannamei* and to evaluate the efficacy of microalgae extracts from *Dunaliella salina* and *Tetraselmis chui* in suppressing the vibriosis infection.

**Material and Method**

**Collection of infected Vannamei shrimp.** The sampling location of Vannamei shrimp, infected with vibriosis, was collected from the Panggang Ayom village, Kaliwungu district, Kendal, Central Java, Indonesia. *L. vannamei* samples had an average weight of 4.30 g and length of 8 cm. Samples was chosen having morphological signs of vibriosis infection (Prayitno & Lachtford 1995). The research was conducted in October 2016 – March 2017. Sample was analyzed at Integrated Laboratory, Diponegoro University, Indonesia.

**Isolation of vibriosis-causing bacteria.** Tryptone Soya Agar (TSA) as universal agar media and Thiosulfate Citrate Bile Salt (TCBS) (Merck, Jerman) as vibrios-specific agar media for bacterial culture were selected. Bacteria were isolated from the infected shrimp using Streacking method (Haldar et al 2011).

**DNA extraction, sequencing and identification of isolated bacteria.** DNA extraction was conducted by using chelex method from Walsh et al (2013) as performed in Susilowati et al (2015). Selected colonies were inoculated in 50-100 μL ddH₂O and 1 mL of 0.5% saponin in PBS 1x (saved overnight). The mixture was centrifuged (12,000 RPM, 10 min). Supernatant was discarded. Then 100 μL ddH₂O and 50 μL of 20% chelex 100 (shake up chelex solution and ensure that some of the crystals make it into sample) were added to a final solution and the solution was boiled for 10 min and vortex once after 5 min. The mixture was centrifuged (12,000 RPM, 10 min) and stored at -20°C. The DNA concentrations were quantified and qualified by using NanoDrop 2000 spectrophotometer (Thermo Scientific). The concentration of 1 μL DNA sample was determined by using the NanoDrop 2000 spectrophotometer (Thermo Scientific). The 260/280 and 260/230 nm ratios was calculated by the NanoDrop spectrophotometer and used to evaluate the DNA purity and also the concentration of DNA.

DNA extracts for 16S rRNA genes sequences were amplified by PCR using universal primers 27F (5'AGAGTTTGATCMTGGCTCAG-3') and 1492 R (5'TACGGTTAACCTTGTACGACTT-
The PCR mixture consisted GoTaq® Green Master Mix Promega (25 μL), primer 27F (0.5-5 μL), primer 1492R (0.5-5 μL), DNA extract (1-5 μL), and Nuclease-Free Water (50 μL). The PCR reaction was performed in a MJ Mini Personal Thermal Cycler (BIO RAD) using cycling conditions consisting of an initial denaturation at 95°C for 3 min followed by 30 cycles of denaturation at 95°C for 1 min, annealing at 55°C for 1 min, and extension at 72°C for 1 min. A final extension was performed at 72°C for 7 min (Lee et al 2006). The PCR products were analyzed by agarose 1% gel electrophoresis and the result was showed by using UVIDoc HD5 (UVITEC Cambridge).

Extraction of microalgae. Dry biomass of microalgae T. chui (10 g) and D. salina (9 g) were dissolved in 100 mL methanol and was left for 5 x 24 h. Methanolic extract of T. chui was passed in ultrasound for 15 min and was evaporated to dryness.

Preparation of feeds. Dry extracts of T. chui and D. salina (2 g) were blended with 10 g of commercial pellet and were left to dry in the drying machine for 2 x 24 h. Commercial pellet without microalgae extracts served as negative control. The commercial pellets used in this study contain 30% of protein, 5% of lipid, 4% of fiber and with 12% moisture.

The use of microalgae as bio-control (in vivo assay). Forty-five vibriosis infected L. vannamei samples were taken based on the previous biomolecular test, used for the in vivo assay. Shrimp were reared in aerated aquarium with a density of five individuals in 5 L of water. The water quality was maintained at 26-28°C and salinity 25-30 ppt. Shrimp were then acclimatized for three days before the in vivo assay.

Shrimps were placed in three different tanks. Each tank represented different treatments related to the type of feeds given. Treated shrimps were fed three times a day, i.e. at 08:00, 12:00 and 17:00 WIB (local time). All experiments were performed in three replicates.

Treatment I: commercial pellet mixed with T. chui extract of 200 ppm against L. vannamei infected with vibriosis.
Treatment II: commercial pellet mixed with D. salina extract of 200 ppm against L. vannamei infected with vibriosis.
Treatment II: commercial pellet without addition of microalgae extract to Vannamei shrimp infected with vibriosis.

Microbial enumeration from the hepatopancreas of shrimp. The observed parameters were amount of bacteria in the hepatopancreas of treated and control shrimps counted using the Total Plate Count (TPC) method (Benson 2001). Hepatopancreas of treated and control shrimp were dissolved in 10 mL sterile seawater. The solutions were stirred for 15 min and were diluted three times (10⁻¹, 10⁻² and 10⁻³). The treatment was carried out for 21 days and the bacteria amount was counted at the 7th, 14th, and 21st days.

Results

Morphological signs of the infected shrimp. The clinical signs of the infected shrimps were ‘luminescent pleopoda’, melanocyst on the shrimp’s body, reddish tail, and brownish-red hepatopancreas.

Bacterial isolates. Based on the isolation of vibriosis-causing bacteria, there were 13 isolates that was morphologically characterized based on their colonial edge form, characteristic and color. Out of 13 bacterial isolates, three potential bacteria were selected and results are presented in Table 1.
**Morphological characteristic sand color of 13 bacterial isolates**

<table>
<thead>
<tr>
<th>No</th>
<th>Colonial edge form</th>
<th>Colonial characteristic</th>
<th>Colonial color</th>
<th>Total</th>
<th>Isolates code</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Jagged</td>
<td>Convex</td>
<td>Green</td>
<td>1</td>
<td>U1H1</td>
</tr>
<tr>
<td>2</td>
<td>Smooth</td>
<td>Convex</td>
<td>Yellow</td>
<td>7</td>
<td>U1H2, U1E1, U2H1, U2H1, U3H2, U4H1, U5H1, U1E2, U3H1</td>
</tr>
<tr>
<td>3</td>
<td>Smooth</td>
<td>Convex</td>
<td>Green</td>
<td>2</td>
<td>U1E2, U3H1</td>
</tr>
<tr>
<td>4</td>
<td>Smooth</td>
<td>Convex</td>
<td>Milk white</td>
<td>1</td>
<td>U7E1</td>
</tr>
<tr>
<td>5</td>
<td>Jagged</td>
<td>Convex</td>
<td>Milk white</td>
<td>1</td>
<td>U8E1</td>
</tr>
<tr>
<td>6</td>
<td>Jagged</td>
<td>Convex</td>
<td>Yellow</td>
<td>1</td>
<td>U8E2</td>
</tr>
</tbody>
</table>

**Bacterial identification.** Based on the biomolecular analysis, there were two bacterial isolates that were classified as same species. This analysis managed to identify the presence of *V. alginolyticus* and *V. harveyi* as two bacteria strain being responsible for the vibriosis infection on the treated shrimps. Biomolecular analysis is presented in Table 2.

**Biochemical analysis of three selected bacterial isolates from Vannamei shrimp**

<table>
<thead>
<tr>
<th>No</th>
<th>Code</th>
<th>Universal Primer</th>
<th>Species Homology</th>
<th>Homology</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>U1H1</td>
<td>1.27F 1.1429R</td>
<td><em>V. alginolyticus</em> strain ATCC 17749</td>
<td>97%</td>
</tr>
<tr>
<td>2</td>
<td>U1E1</td>
<td>2.27F 2.1429R</td>
<td><em>V. harveyi</em> strain NCIMB 1280</td>
<td>96%</td>
</tr>
<tr>
<td>3</td>
<td>U4H1</td>
<td>3.27F 3.1429R</td>
<td><em>V. alginolyticus</em> strain ATCC 17749</td>
<td>97%</td>
</tr>
</tbody>
</table>

**In vivo assay.** Forty-five shrimps infected by vibrio reared during 21 days and feed with *D. salina* and *T. chui* showed a decreased of bacteria amount (Figure 1).

![Figure 1](image-url)  
*Figure 1. Number of bacteria observed from infected *Litopenaeus vannamei* during the 21 days experiment.*

**Discussion.** Clinical signs observed on the infected *L. vannamei* collected from Kaliwungu, Kendal were in accordance as infection signs in Vannamei caused by vibrio’s
as reported by Lina et al (2007) and Huang et al (2013). Another features remarked correlates to the behavior of shrimp are passive movement, slow response to feed and fluorescent swimming legs. Prayitno & Lachtford (1995) stated that V. harveyi is one of causative agent that also known as luminescent bacteria. Those clinical signs became the reference in isolating the vibriosis-causing bacteria from the shrimp.

Bacterial isolates demonstrated that there were 13 isolates with diverse morphology. The selected isolate, which was U1H1, had a jagged colonial edge, convex with green color. Meanwhile, the U1E1 AND U4H1 were characterized with smooth colonial edge, convex with yellow color. These three isolates showed similar characteristic as Vibrio bacteria. Sarjito et al (2015) revealed that morphological features of bacteria with smooth colonial edge, convex with yellow color tested biochemically were identical with those of V. harveyi.

DNA amplification results from the rep-PCR exhibited three isolates with similar band profiles as seen on marker 800 bp. Rep-PCR method has been proven as an efficient method to identify bacterial species and the genetic relationship (Lina et al 2007). DNA sequencing of isolate U1H1 and U4H1 had identical nucleotide with V. alginolyticus strain ATCC 17749 with homological level 97%. Isolate U1E1 showed to have similar nucleotide with V. harveyi strain NCIMB 1280 with homological level 96%.

Presence and abundance of V. alginolyticus and V. harveyi are highly correlated with environmental conditions. Asplund (2013) stated that the abundance of bacteria Vibrio sp. usually relates to temperature and salinity and increases in stable environmental condition. The abundance of V. alginolyticus and V. harveyi counted was 21 (isolate U1H1) and 4 (isolate U4H1) colony/plate and 11 (isolate U1E1) colony/plate. Environmental condition in the pond when the sampling was performed in October-November was fluctuated due to the rainy season. In this season, the temperature and salinity in the pond tended to change in short period; thus, it affected the abundance of bacterial colony, either in the pond or on the L. vannamei body.

The result of the in vivo assay shown that the vibrioid infected shrimps feed with mixed T. chui extract and D. salina reared for 21 days shown a decreased of the number of bacterial infection as shown on Figure 1. Widowati et al (2017) observed that Dunalliela sp., T. chui showed antioxidant potential which could be considered for future applications in aquaculture. The result of this study indicated that the microalgae are capable to produce antibacterial compounds against vibrio.

Conclusions. The present study showed that vibriosis-causing bacteria isolated from L. vannamei, which showed clinical signs of vibriosis infection, was identified as V. alginolyticus, and as V. harveyi. The shrimps infected by vibrio and feed with D. salina and T. chui reared during 21 days showed a decreased of bacteria amount. The result indicated that the microalgae were capable to produce antibacterial compounds against vibrio. Based on the present findings, it could be inferred that the secondary metabolites of T. chui and D. salina may be an excellent source in order to develop potent formulations for sustainable shrimp farming.

Acknowledgements. The authors wish to thank the Directorate of Research and Community Service, Directorate General of Research Strengthening and Development, Indonesian Ministry of Research Technology and Higher Education, for funding this research through the Implementation Agreement Assignment of Research Collaboration and International Publication 2017.

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