

Original Paper

APPLICATION OF REPETITIVE SEQUENCE-BASED PCR ON THE RICHNESS OF VIBRIO ON THE TIGER SHRIMP (*Penaeus monodon* Fab.)

Sarjito¹, Nestin E. W. Ningrum², Ocky Karna Radjasa³ and Slamet Budi Prayitno¹

¹Aquaculture Study Program, Fisheries Department, Fisheries and Marine Science Faculty, Diponegoro University, Semarang 50275, Central Java, Indonesia

²Mandiri Bank, Jakarta Office Branch

³Marine Science Study Program, Fisheries Department, Fisheries and Marine Science Faculty, Diponegoro University, Semarang 50275, Central Java, Indonesia

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ABSTRACT

A molecular-based study was conducted to estimate the richness of the *Vibrio* on tiger shrimp (*Penaeus monodon* Fab.) from brackish water pond of Mororejo, Kendal. Tiger shrimps were collected from the extensive brackish water pond and a total of 22 isolates were obtained from hepatopancreas and telson of tiger shrimp. Based on the repetitive sequence-based polymerase chain reaction (rep-PCR), it was found that two groups of *Vibrio*. To investigate the effectiveness of rep-PCR in estimating the richness of *Vibrio* on tiger shrimps, three isolates (JTW 01, JTW 03 and JTW 06) were chosen for further investigation. On the basis of sequence analysis, the result showed that the JTW 01, JTW 03 and JTW 06 were closely related to *Vibrio* sp. Absa7 clone 423.1, *Vibrio splendidus* and *Vibrio splendidus*, respectively. The result proved that two associated of *Vibrio* on tiger shrimp were *Vibrio* sp. Absa7 clone 423.1 and *Vibrio splendidus*. Therefore the present study highlights the effectiveness of rep-PCR in rapid grouping and estimating the richness of *Vibrio* on tiger shrimp.

Keywords: Rep-PCR; Vibriosis; Causative Agent; *Penaeus monodon* Fab

*Correspondence: Phone: +62-24-7474698; E-mail: sarjito_msdp@yahoo.com

INTRODUCTION

Tiger shrimp (*Penaeus monodon* Fab.) is a potential fishery commodity which has high economic value in domestic and international markets. While demanding of shrimp in the world is increasing, shrimp production is decreasing every year. Declining of shrimp production partly due to disease caused by bacterial disease like Vibriosis.

Vibriosis is a serious problem in the majority of penaeid shrimp culture operations. *Vibrio* species are a normal part of the bacterial flora in aquatic environments and formerly considered to be mostly opportunistic pathogens (Lightner, 1996; Myers, *et al.*, 2006; Thompson, *et al.*, 2003). However, some more occurring disease syndromes of penaeid shrimp have been caused by *Vibrio* species which behave more like true pathogens than opportunistic invaders

(Gomez-Gil, *et al.*, 2004; Kannaripan, *et al.*, 2008). Vibriosis is the main cause of production loss due to bacterial disease in penaeid shrimp farms (Kannaripan *et al.*, 2008). Vibriosis causes mortality in larvae, postlarvae, juveniles, sub-adults and also adults of shrimps. Outbreaks of the disease cause mortality up to nearly 100% of affected population (Sunaryanto and Mariyam, 1987). The gross signs of localized infection in the cuticle or sub-cuticle are called shell disease or black or brown spot disease and these superficial infections can develop into systemic infections under some circumstances. It is the systemic infections that cause mortality (Chen *et al.*, 1992; Myers, *et al.*, 2003; Suddesh and Xu, 2001.).

Recently, many reports have also been published on the 16S rRNA gene sequences of

bacteria and the phylogenetic relationships deduced from analysis of these sequences (Radjasa, *et al.*, 2001; Radjasa, *et al.*, 2007a; Sabdono, 2001; Sarjito, *et al.*, 2009). Most of the results indicate that phylogenetic relationships based on 16S rRNA sequences support the distinction of species among eubacteria, archaeobacteria, and eukaryotes (Radjasa, *et al.*, 2001; Sabdono, 2001). Because of this feature, Cunningham (2002) suggested to use molecular methods for diagnostic of fish diseases. The molecular methods using 16S rDNA for detection of causative agent of Vibriosis have been conducted in Turbot, *Colistium nudipinnis*, (Diggles, *et al.*, 2000; Montes, *et al.*, 2006), Brill, *C. Guntheri*, (Diggles, *et al.*, 2000), Spotted Rose Snapper, *Lutjanus guttatus*, Steindachner, 1869 (Gomez-Gil, *et al.*, 2004), groupers (Sarjito, *et al.*, 2008; Sarjito, *et al.*, 2009) and white shrimps, *Litopeneus vannamei*, (Sarjito *et al.*, 2011).

Various molecular methods have been applied on *Vibrio*, such as: RAPD (Somarny *et al.*, 2002) and rep-PCR (repetitive sequence based polychain reaction) (Sarjito, 2011). Rep-PCR was conducted in order to group a bacterial number isolates that produced complex fingerprint profiles from gram negative bacteria. Furthermore, rep-PCR has been applied on various diversity of associated sponge bacteria (Radjasa, *et al.*, 2007a.b), psychrotrophic bacteria from Makasar straight (Radjasa, *et al.*, 2007c) and causative agent of Vibriosis (Sarjito, *et al.*, 2009). However, to best our knowledge, there has limited report been documented so far on describing the application of rep-PCR on the richness of *Vibrio* on tiger shrimp from Indonesian extensive brackish water pond using molecular tools. The research regarding the diversity of *Vibrio* as the causative agent of vibriosis is important for creating health management of tiger shrimp culture. In this study, we reported the richness of *Vibrio* bacteria on tiger shrimp from extensive brackish water pond of Kendal, Central Java assessed by 16S rDNA approach.

MATERIALS AND METHODS

Sampling of Tiger Shrimp

The shrimps were collected from extensive culture in extensive brackish water pond of Mororejo village, Kendal Regency, Central Java,

Indonesia and was identified as tiger shrimp (*Penaeus monodon* Fab.). After collection, tiger shrimps were put into the plastic containers and immediately brought to the integrated Marine Science Laboratory of Fisheries and Marine Science Faculty, Diponegoro University in Semarang, Central Java for bacterial isolation.

Bacterial Isolation

Bacteria *Vibrio* were isolated directly from hepatopancreas and telson of tiger shrimps by streak method on TCBS medium. Bacterial isolation was also conducted from the inner part of hepatopancreas and telson, which were scraped off with a sterile knife. The resultant tissues were serially diluted, spread on TCBS agar medium and were incubated at room temperature for 24-48 hours. On the morphological features, colonies were randomly picked and purified by making streak plate (Brock and Madigan, 1991).

Repetitive – PCR

The procedure was carried out according a method previously described by Radjasa *et al.* (2007b). In the rep-PCR, BOX AIR (5'-CTACggCAAaggCgACgCTgACg-3') (Versalovic *et al.*, 1994) was used. The REP 1R-I and REP 2-I primers contain the nucleotide inosine (I) at ambiguous positions in the REP consensus. PCR reaction contained of 1 µL DNA template (diluted 100x), 1 µL primer, 7.5 µL Megamix Royal and sterile water up to total volume of 15 µL.

Amplifications were performed with a thermal cycler model Gene Amp PCR system 9700 with the following temperature conditions: initial denaturation at 95°C for 5 minutes; 30 cycled of denaturation (92°C for 1 minutes), annealing (50°C for 1,5 minutes), extension (68°C for 8 minutes) and final extension at 68°C for 10 minutes. Five microliter aliquot PCR products were run using electrophoresis on 1 % ethidium bromide gel by using 1X TBE buffer.

Grouping of Isolates

Grouping was carried out according to a method of Radjasa, *et al.*, (2007c) by making matrixes from the position of bands on the gel which were there analyzed by using Free Tree program by using UPGMA method for constructing the tree.

Resampling was performed by bootstrapping with 1000 replications.

PCR Amplification and Sequencing of 16s rRNA Gene Fragments

PCR amplification was carried out according to method of Radjasa *et al.*,(2007a). Two primers, GM3F (5'AGAGTTTGATCMTGGC-3') and GM4R (5'-TACCTTGTTACGACTT-3') were used to amplify nearly complete 16S rRNA gene. Genomic DNA of causative agent of vibriosis strains for PCR analysis were obtained from cell materials taken from agar plate, suspended in sterile water (Sigma, Germany) and subjected to five cycles of freeze (-80°C) and thaw (95°C). PCR amplification of partial 16S rRNA gene of bacteria, purification of PCR products and subsequent sequencing analysis were performed according to the method of Radjasa, *et al.* (2007b). The determined DNA sequences of

strains were then compared for homology to the BLAST database (Atschul, *et al.*, 1997).

RESULTS AND DISCUSSION

Results

Characteristic of the Bacterial Isolates

The clinical signs of tiger shrimps affected by Vibriosis from extensive brackish water pond of Kendal Regency, Indonesia were body melanosis, white spot, pale abdomen, dark mouth and red color at telson and tail.

Bacterial isolation resulted in total of 22 Vibrio isolates (JTW 1 – JTW 22) obtained from hepatopancreas of tiger shrimp (**Table 1**).

Repetitive-PCR Analysis

Based on the repetitive-PCR result and constructed dendrogram of the Vibrio as causative agent of vibriosis on tiger shrimp, two groups were formed (**Fig.1**).

Table 1. Characteristic of isolates associated with tiger shrimp (*Penaeus monodon*) from extensive brackish water ponds of Kendal Regency, Indonesia.

No	Isolate Code	Isolate Source	Colony Color on TCBS	Colony Form
1	JTW 1	Hepatopankreas	Green	Round
2	JTW 2	Hepatopankreas	Yellow	Round
3	JTW 3	Hepatopankreas	Black	Round
4	JTW 4	Hepatopankreas	Yellow	Round
5	JTW 5	Hepatopankreas	Shiny yellow	Round
6	JTW 6	Hepatopankreas	Yellow	Round
7	JTW 7	Hepatopankreas	Dark yellow	Round
8	JTW 8	Hepatopankreas	Dark yellow	Round
9	JTW 9	Hepatopankreas	Dark green	Round
10	JTW10	Hepatopankreas	Black	Round
11	JTW11	Hepatopankreas	Yellow	Round
12	JTW12	Hepatopankreas	Shiny green	Round
13	JTW13	Hepatopankreas	Dark yellow	Round
14	JTW14	Hepatopankreas	Black	Round
15	JTW15	Hepatopankreas	Green	Round
16	JTW16	Hepatopankreas	Yellow	Round
17	JTW17	Hepatopankreas	Black	Round
18	JTW18	Hepatopankreas	Yellow	Round
19	JTW19	Hepatopankreas	Shiny yellow	Round
20	JTW20	Hepatopankreas	Yellow	Round
21	JTW21	Hepatopankreas	Black	Round
22	JTW22	Hepatopankreas	Shiny yellow	Round

Repetitive-PCR Analysis

causative agent of vibriosis on tiger shrimp, two groups were formed (**Fig.1**).

Based on the repetitive-PCR result and constructed dendrogram of the *Vibrio* as

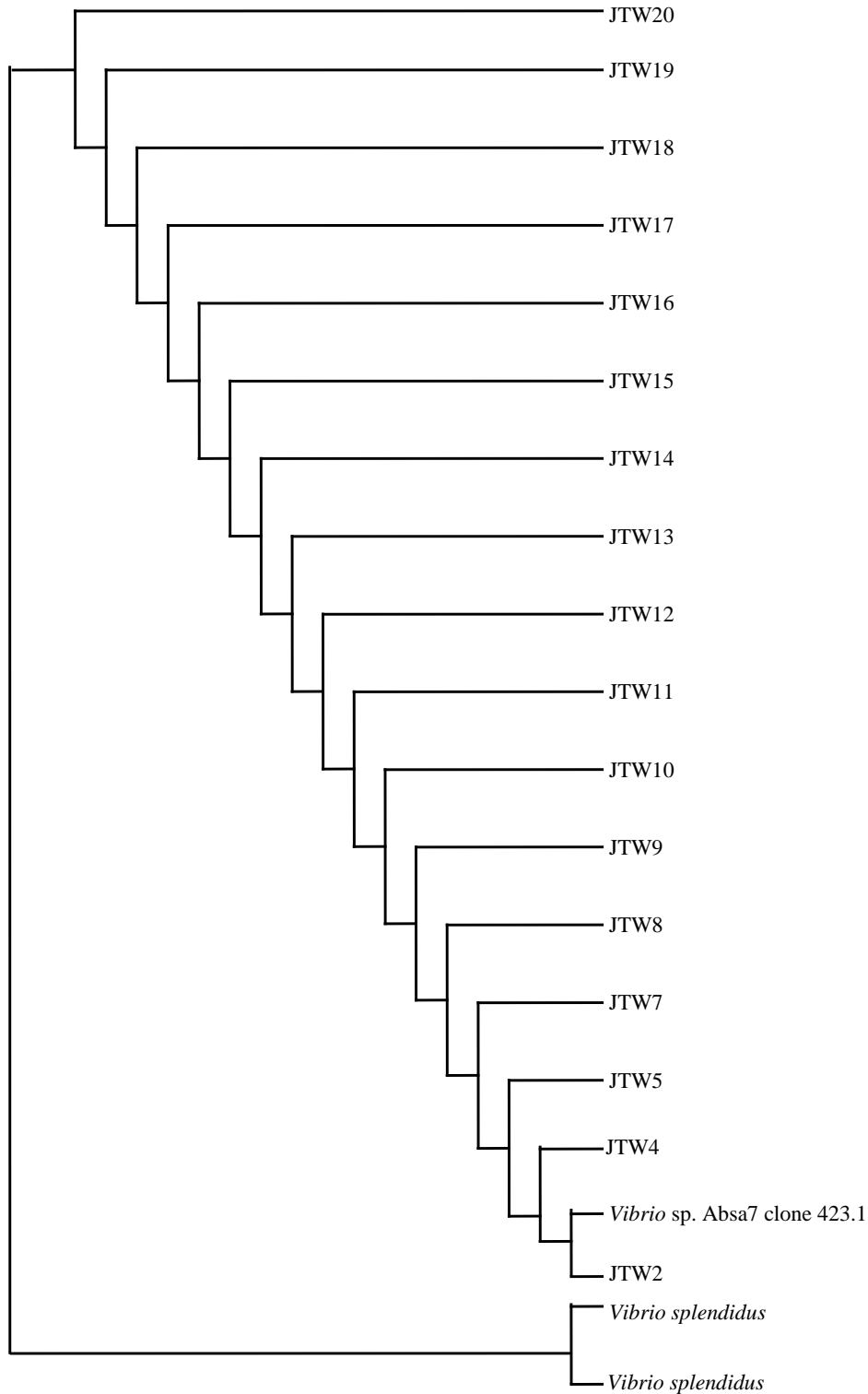


Fig.1. Diagram of *Vibrio* group based on the Repetive Sequence-Based Polymerase Chain Reaction and 16S rDNA on the Tiger Shrimps.

Sequencing of representative Vibrio as causative agent on tiger shrimps

showed that all isolates were the members of genus *Vibrio* as presented in **Table 2**

Based on molecular characterization it was

Table 2. Molecular characterization of 3 representative of causative agents

No.	Isolates	Close Relative	Homology (%)	Acc. Number
1	JTW 1	<i>Vibrio</i> sp. Absa7 clone 423.1	96	DQ357813.1
2.	JTW 3	<i>Vibrio splendidus</i>	100	GQ254509.1
3.	JTW 06	<i>Vibrio splendidus</i>	100	GQ254509.1

Based on the **Fig. 2** and **Table 2** showed that vibrio on tiger shrimp from extensive brackish water pond of Kendal regency was *Vibrio* sp. Absa7 clone 423.1 (JTW 01 - Groups I) and *Vibrio splendidus* (JTW 03; JTW 06 -groups II) with a homology of 96 % and 100 % respectively.

Discussion

It is widely known that the characterization and identification of bacterial isolates have traditionally been based on phenotypic traits, which are derived from biochemical test reactions. It is also noted that one disadvantage of these tests has been the requirement of strains to grow in order to produce a detectable reaction. The recent and rapid development of molecular biology techniques have been regarded as the solution to the problem of bacterial identification based on phenotypic approach.

In this work, a molecular biology approach based on 16S rDNA coupled with rapid grouping technique, rep-PCR was applied to estimate the richness of pathogens *Vibrio* associated with tiger shrimp from brackish waters of Kendal, Central Java.

Vibrio species are natural habitants of seawater and brackish water widely distributed throughout the world (Myers *et al.*, 2003). However, some species have exhibited clinical significance for aquatic animal and are recognized as potential pathogens (Myers *et al.*, 2006). The large number of *Vibrio* shrimp pathogens causing epizootic outbreaks in aquaculture has made it necessary to develop efficient, fast and sensitive methods for their detection. Both detection and identification of vibrios have been traditionally depend on their growth on Thio-Sulphate Citrate Bile Salt

Sucrose (TCBSA) selective medium and subsequent characterization by biochemical test (Diggles *et al.*, 2000).

Vibriosis in the tiger srimps was characterized by melanosis of body, white spot, pale abdomen, dark mouth, red color at tail and lesion of the tail.

The result of this study revealed that the application of Rep-PCR has been a reliable tool for strain rapid grouping and differentiation / richness among the vibrio on the tiger shrimp in extensive brackish water pond of Kendal Regency. This molecular approach may be used for the analysis of other *Vibrio* species related to aquaculture disease. Molecular identification of *Vibrio* of on shows that the identified strains were nicely in accordance with the dendrogram constructed from the Rep-PCR analysis. The identify strains were *Vibrio* sp. Absa7 clone 423.1 (JTW 01) and *Vibrio splendidus* (JTW 03 and JTW 06). *Vibrio* sp. Absa 7 clone 42y 3.1. was identified by Schulze *et al.*, (2006) on water of marine hatchery-abalone larvae from Vancouver Island waters, Canada. Whereas *V. splendidus* was reported by Panicker *et al.* (2004) on shell fish and water of Mexico Gulf.

In conclusion, the application of the Rep-PCR method is useful for rapid grouping and estimating of richness of vibrio as causative agents of vibriosis on tiger shrimp with high power and offers an alternative technique for grouping of numerous of marine bacterial isolates.

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