



Physiological, biochemical and *HSP70* and *HSP90* genes expression profiles of tropical abalone *Haliotis squamata* in response to *Vibrio alginolyticus* infection

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ABSTRACT *Vibrio* spp. have been known responsible for fish diseases in marine and brackish-water systems in the tropical regions. Heat shock proteins are a highly conserved protein sequence that is known for its rapid response to environmental stresses, including infection. This study aims to investigate physiological and biochemical responses tropical abalone *Haliotis squamata* to *Vibrio alginolyticus* infection. Abalones were infected with *V. alginolyticus* by intramuscular injection at a concentration of $10^5, 10^6, 10^7$ cfu/abalone. The expression of *HSP70* and *HSP90* genes, the activity of superoxide dismutase, phenol oxidase and catalase enzymes, histology, falling, and mortality were observed at 12, 24, 48, 72, and 96 hours post-infection (hpi). The different expression of *HSPs* was found in this study. While the expression of *HSP70* was downregulated after infection, the expression of *HSP90* is upregulated at 12 hpi and followed by downregulated after 24 hpi for 10^6 cfu infection, and expressed at a normal level for 10^5 infection treatment. The expression of superoxide dismutase and catalase increased within 12 hpi, expression of phenol oxidase was increased after 24 hpi. *V. alginolyticus* was virulent with LD50 on *H. squamata* with an average weight of 5.13 g is less than 105cfu and it caused enlargement of hemolymph sinus, development of abscess intraepithelial and intramuscular.

KEYWORDS *H. squamata*; *V. alginolyticus*; physiology; *HSP*; histology

1. Introduction

The development of aquaculture on increasing intensification and commercialization of aquatic production will increase occurring major disease problems (Bondad-Reantaso et al. 2005). Abalones or ear shells have a low shell, open spiral structure, and are characterized by several open respiratory pores in a row near the shell's outer edge. Abalone species are economically valuable for fishery production in the temperate or sub-tropical areas. Whereas, the large size of abalones are distributed in temperate seas, the small size abalones are distributed at wide-range geographical distribution in warm water, including Indonesia. The commercial aquaculture of the small size of abalone has already well developed, especially in East Asia (Hsu and Gwo 2017). The aquaculture industry has been overwhelmed with its share of diseases and problems caused by several pathogens (Bondad-Reantaso et al. 2005). In Taiwan, the production of small abalone has been dramatically decreased in the past 15 years due to the lack of suitable diatom feed for larvae, poor water quality,

habitat degradation, genetic problems, disease and infection problems (Hsu and Gwo 2017).

Given that bacteria can survive well in aquatic environments independently of their hosts, bacterial diseases have become major impediments to aquaculture, especially when the water temperature is warm (Pridgeon and Klesius 2012). The most frequently encountered bacterial agents associated with fish diseases in marine and brackish-water systems in the tropical environments are *Vibrio* spp. (Karunasagar et al. 2003). *Vibrio harveyi* is known to be pathogenic in a large range of vertebrates and invertebrates, including molluscs. Abalone diseases due to the pathogen *V. harveyi* have been described in *Haliotis diversicolor*, *H. laevis* and *H. tuberculata* causing septicaemia. The evidence of vibriosis on abalone has outbreak in Taiwan in 2000 which was caused by *V. parahaemolyticus* and made a significant economic loss in abalone (*H. diversicolor supertexta* L.) industry (Cheng et al. 2004; Cai et al. 2006b). The *V. alginolyticus* H-11 strain has been isolated from a mass mortality out-

breaks of small abalone *H. diversicolor* supertexta with abscess/ulcers in the mantle that occurred in 1998 at Kao-Hsiung, Taiwan. This strain and its extracellular products were virulent to small abalones with LD50 values of 3.6×10^5 colony forming units (cfu) and 2.96 μg protein/g body weight, respectively (Liu et al. 2001). *V. alginolyticus* also caused disease on post larvae and small juvenile *H. diversicolor*. This bacterium is virulent with LD50 as 1.0×10^4 cfu/ml on post larvae (Cai et al. 2006a).

Animals have defense mechanisms against the pathogen, which is composed of innate and adaptive immune systems. The innate immune system is the only defense system that existed in invertebrates. This innate immune system is the first line of defense against non-self pathogens and can be divided into physical barriers, cellular, and humoral components. Specifically, humoral parameters include growth inhibitors, various lytic enzymes and components of the complement pathways, agglutinins, precipitins, natural antibodies, cytokines, chemokines, and antibacterial peptides. Furthermore, several external and internal factors can also influence the activity of innate immune parameters. The cellular immune system is performed by several types of cells (Magnadóttir 2006).

Hemocytes in mollusks are consisted of agranular and granular hemocytes, which are involved in phagocytosis, an important process of eliminating microorganisms or foreign particles. During phagocytosis, several types of reactive oxygen intermediates (ROIs) are produced, such as: superoxide anion (O_2^-), hydrogen peroxide (H_2O_2), singlet oxygen, and hydroxyl radical. The release of superoxide anion is known as the respiratory burst, and it plays an essential role in antibacterial activity (Cheng et al. 2004). Acid phosphatase (ACP) and alkaline phosphatase (AKP) are important for innate immune defense in the small size of abalones. Superoxide dismutase (SOD) is a key antioxidant enzyme playing a first-line protective role against reactive oxygen species (ROS) by converting superoxide (O_2^-) into H_2O_2 . The AKP, and SOD activities of diseased abalones were significantly lower than in the healthy group (Di et al. 2016). Several enzymes on abalone have been evaluated in response to pathogen infection on *H. diversicolor* (Yao et al. 2019), the high water temperature stress on greenlip abalone (*H. laevigata*) (Buss et al. 2017).

Heat Shock Proteins (HSPs) are a group of highly conserved chaperone proteins expressed by the cell that respond to unfavorable environmental changes (?). The HSPs are considered as ubiquitous protein and widely preserved in prokaryotic and eukaryotic organisms (Roberts et al. 2010). These proteins have functioned as cellular defenses, prevent protein denaturation, and assist in the re-introduction and removal of denatured protein due to biotic and abiotic pressures (Wang et al. 2004). In aquatic organisms, expression of HSP genes was increased as a response to several stresses, such as heat (Park et al. 2015), organic pollutants (Paulino et al. 2014), correlations between metals (Qian et al. 2012), and *Vibrio* infections (Rungrassamee et al. 2010).

Many studies on physiology and disease have been conducted on abalones from the temperate or subtropical zone (Rungrassamee et al. 2010; Di et al. 2016; Fang et al. 2019; Yao et al. 2019). However, only limited studies have been addressed on tropical abalone. The *H. squamata* is an indigenous species with an excellent taste and has been caught on the southern coast of Bali. This species was started to be cultured, especially in Bali. In this study, we investigated the biological responses of tropical abalone *H. squamata* in the response to *V. alginolyticus* infection. This study is the first investigation of *V. alginolyticus* infection in *H. squamata* in Indonesia with a comprehensive evaluation of mortality, histology, enzymes activity, and HSP genes expression.

2. Materials and Methods

2.1. Animal collection and maintenance

The uniform and high quality abalone seeds are very essential for this study, then this research begins with the hatching of abalone in the Abalone Hatchery Unit, National Broodstock Center for Shrimps and Mollusc in Tigarong, Karangasem, Bali. The abalons with a normal morphological and appearance, agile movements, sticking firmly to the substrate, minimal size of the shell length 4 cm were selected as broodstocks for use in this study. The broodstocks were maintained in fiberglass tubs with PVC pipes as shelters and fed with *Gracillaria* sp. and *Ulva* sp. seaweed at the dosage of 10-20% of biomass/day.

The stress treatment was applied to mature gonad abalones for inducing the spawning. The stress was addressed by lifting the basket of the broodstock from the water tank for one hour then put it back into the water. Then the broodstock was maintained in a tank with a flowing water system until spawning. Eggs produced from spawning abalone were harvested using an egg collector. After 12 to 13 h of incubation, the eggs hatched into first-stage swimming larvae, trochophores. The trochophores within a few hours this becomes a veliger larvae. The veliger larvae were fed with attached diatoms (*Nitzschia* sp. and *Navicula* sp.) which attached on the substrate rearing plate.

After one month rearing, the veliger larvae reached a juvenile stage at size diameter of shell more of than 0.6 cm. The juveniles were reared on the basket in the tank and fed with macroalgae *Gracillaria* sp. and *Ulva* sp. The grading was carried out every two months for continue rearing on relatively same size. After eight months of rearing, the juvenile abalone *H. squamata* with an average shell length of 32.97 ± 1.83 mm and an average weight of 5.13 ± 0.83 g were used for this study. The abalones from the hatchery were acclimatized to laboratory conditions for one week. During acclimatization, abalones were reared on pipe basket in the tank with seawater at a salinity of 34 g/L, the temperature at 29-30 °C and fed with seaweed *Gracillaria* sp. twice a day.

2.2. *Vibrio alginolyticus* infection

A pathogenic strain of *V. alginolyticus* was received from Fish Disease and Environmental Inspection Center at Serang Banten. The bacterium was cultured on nutrient broth and incubated at 35 °C for 48 h. The bacterium was harvested, washed and suspended PBS on at desired concentration for infection treatments.

The *V. alginolyticus* infection was conducted by intramuscular injection on pallial sinus using 25 gauge 1 mL syringe at a concentration of $10^5, 10^6, 10^7$ cfu/abalone with a volume of 100 µL. For the control, abalones were injected with 100 µL of PBS. After injection, the abalones were kept on pipe baskets and observed on the superoxide dismutase (SOD), phenol oxidase (PO) and catalase (CAT) enzymes activity, *Heat Shock Proteins (HSPs)* expression, survival rate, falling rate and histology.

2.3. SOD, PO, and CAT enzymes activity

The evaluation of enzymes activity was performed by sampling at 0, 12, 24, 48, 72, and 96 h post *V. alginolyticus* infection. The hemolymph was collected and pooled from three animals for measuring the SOD, PO and CAT activity. The SOD activity was determined by measuring the ability to inhibit the reduction of photochemical nitroblue tetrazolium chloride (NBT), as described previously (Datkhile et al. 2009) with SOD Kit-WST (water-soluble tetrazolium salt) Access (Dojindo, Japan). Briefly, 40 µL of hemolymph was added into 360 µL buffer phosphate, then centrifuged at 6000 g at 4 °C for 7 min. The supernatant was then heated up at 65 °C for 5 min to obtain the crude extract. Finally, 150 µL of the crude extract was added with 50 µL of nitroblue tetrazolium (NBT) reagent (0.1 Mm EDTA, 13 µM methionine, 0.75 mM NBT and 20 µM riboflavin in 50 mM phosphate buffer, pH 7.8) and incubated for 2 min. Then the optical density was measured at 4500 nm using a spectrophotometer.

Phenol oxidase activity was measured spectrophotometrically by recording the formation of dopachrome produced from L-dihydroxyphenylalanine (L-DOPA) according to (Hooper et al. 2014). One hundred microliters of hemolymph plasma were transferred in duplicate to 96-well microplate wells. The 100µL of L-DOPA (30 mM L-3,4-dihydroxyphenylalanine, Sigma D9628, in HCl 0.2 M, pH 8) was added to each well and mixed for 10 s. The absorbance at 492 nm was recorded every 5 min at 20 °C for over than 30 min, using a microplate reader Heales® MB-580, (Shenzhen Huisong Technology China).

Catalase activity was measured colorimetrically by CAT activity Assay Kit (GeneWay, Biotech) according to the manufacture instruction. The level of H₂O₂ loss was measured by reading absorbance with a microplate reader at 492 nm. One unit of enzyme was defined as the amount of enzyme required to convert 1 mol of H₂O₂ to the product in one min in pH 4.5 at 25°C.

2.4. HSPs expression

The hemolymph was collected from the animal using a syringe at 0, 12, 24, 48, 72 and 96 h post *V. alginolyticus* infection. The hemolymphs from three animals were pooled in microtube then immediately proceed for RNA extraction or kept at -80 °C until ready to be used. Total RNA was extracted from hemolymph using QuickRNA MiniPrepPlus Kit (R1058) (Zymo Research) following manufacturer protocol. The integrity of RNA was assessed by electrophoresis on 1.2% agarose gel. The purity of RNA was verified by measuring absorbance at 260 nm and 280 nm with NDD 2000 (Nano Drop Technologies, USA). The cDNA was synthesized by mixing the 100 µg of RNA with others component of ReverTra Ace® qPCR RT Master Mix (Toyobo, Japan). The mixture was incubated at 37 °C for 15 min and at 50 °C for 5 min, then followed by incubation at 98 °C for 5 min and enzyme inactivation.

HSP gene expression was measured by real-time PCR using Thunderbird SYBR® qPCR kit with Applied Biosystem machine (ABI, USA). The 2 µL cDNA was used in each reaction and analyzed in triplicate. The HSP90 F (CCAGGAAGAATATGCCGAGT) and HSP90 R (CACGGAACTCCAAGTACC) primers were used to evaluate HSP90 expression, while HSP70 F (CCGCTCTAGAACTAGTGGAT) and HSP70 R (CCGCCAAGTGGGTGTCT) primers were used to evaluate HSP90 expression, and β-actin F (GGGTGTGATGGTCCGGTAT) and β-actin R (AGCGAGGGCAGTGATTTC) primer pairs were used to determining the expression of β-actin as an internal control (Farcy et al. 2007). The thermal cycling condition was 95°C for 30 seconds for the initial denaturation stage, followed by 40 cycles of 95°C for 5 seconds, 58°C for 30 seconds, and 72°C for 30 seconds for final extension stage. At the end of reaction, the melting or dissociation curve analysis to ensure reaction specificity. This analysis was applied by increasing temperature from 65°C to 95°C, with rate increasing the temperature at 0.5°C sec⁻¹.

2.5. Falling rate

The abalones were injected intramuscularly with *V. alginolyticus* at a dose of $10^5, 10^6, 10^7$ cfu/abalone Falling rate was conducted to evaluate the changes of adhesion ability of abalones on the PVC substrate. Thirty abalones were attached to vertical PVC pipe substrates in the aquarium. The numbers of fallen abalones from the vertical substrate was recorded every 12 h. This experiment was conducted in triplicate.

2.6. Survival rate

Thirty abalones from each dose infection treatment were transferred to aquaria. The mortality of abalone was recorded daily. The death of abalone was indicated by fallen from the wall, laid at the bottom with upside-down position or the shell at the floor. This experiment was conducted in triplicate.

2.7. Histological analysis

Histology was conducted to observe the effects of *V. alginolyticus* infection on the foot muscle structure. The abalones were collected at 96 hpi, the shells were removed, and the tissues were fixed in Bouin's solution. The tissues were proceeded on standard histology, then sectioned at a thickness of five μm and stained with H&E. The slides were observed under light microscopy (ZEISS Primovert P35-C). The level of histological alterations in the foot was determined descriptively.

3. Results and Discussion

3.1. HSP70 and HSP90 genes expression of *H. squamata*

Heat-shock proteins (HSPs) are a group of highly conserved proteins which responsible for responding to disease infection. The expression profile of the two HSP genes in the abalone hemolymph after *V. alginolyticus* infection was shown in Figure 1. *HSP70* and *HSP90* are expressed in a different pattern. The expression level of *HSP70* was decreased rapidly in the first 12 h after infection (hpi), and remained in a low expression level until at the end of the experiment at 96 hpi (Figure 1). In contrast, the expression of *HSP90* gene was increased in all infection treatments at 12 hpi, with the *HSP90* expression at 10^7

cfu infection treatment reached 4.5 times over the control. Next, the *HSP90* expression was decreased in all infection treatment after 24 hpi. Moreover, the *HSP90* expression at 10^6 and 10^7 cfu infection treatments were still in a low level until 96 hpi. Meanwhile, the expression of *HSP90* at 10^5 cfu infection treatment was expressed at the nearly same level with control (Figure 1B).

3.2. Biochemical responses of *H. squamata* to *V. alginolyticus* infection

Hemocytes are involved in phagocytosis for the elimination of microorganisms or foreign particles. Several enzymes play important roles in phagocytosis process. Therefore, the superoxide dismutase (SOD), phenol oxidase (PO) and catalase (CAT) enzyme activity were measured to determine abalone responses to *V. alginolyticus* infection at 0, 12, 24, 48, 72 and 96 hpi (Figure 2). Result showed that the SOD activity was increased in abalone infected with *V. alginolyticus* ($P < 0.05$) compared to the control after 12 h with the highest increased of SOD activity was observed in abalones with 10^6 cfu infection. Moreover, after 24 h, SOD activity was decreased significantly in the abalone with 10^5 and 10^5 cfu infection treatments

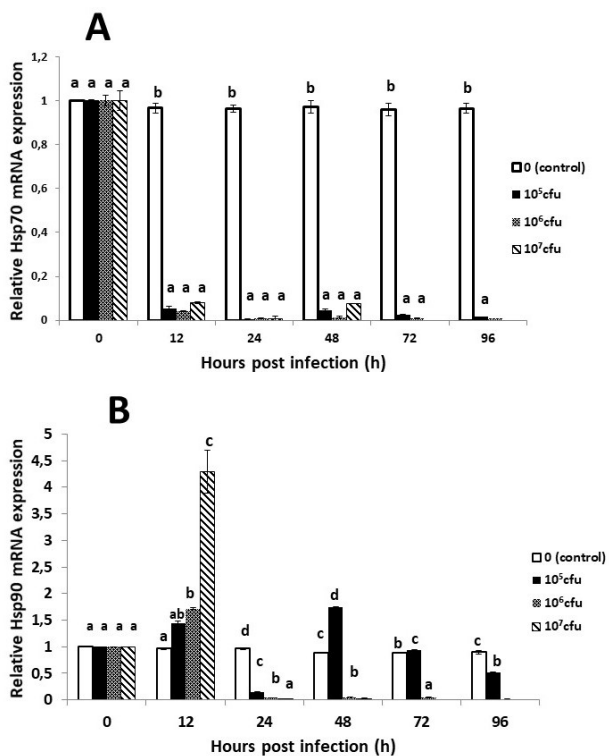


FIGURE 1 The relative expression levels *HSP70* (A) and *HSP90* (B) of abalone *H. squamata* at various *V. alginolyticus* infection as indicated. Different lowercase letters on each observation indicates a significant difference of the expression ($P < 0.05$) compared with the control using Tukey's test.

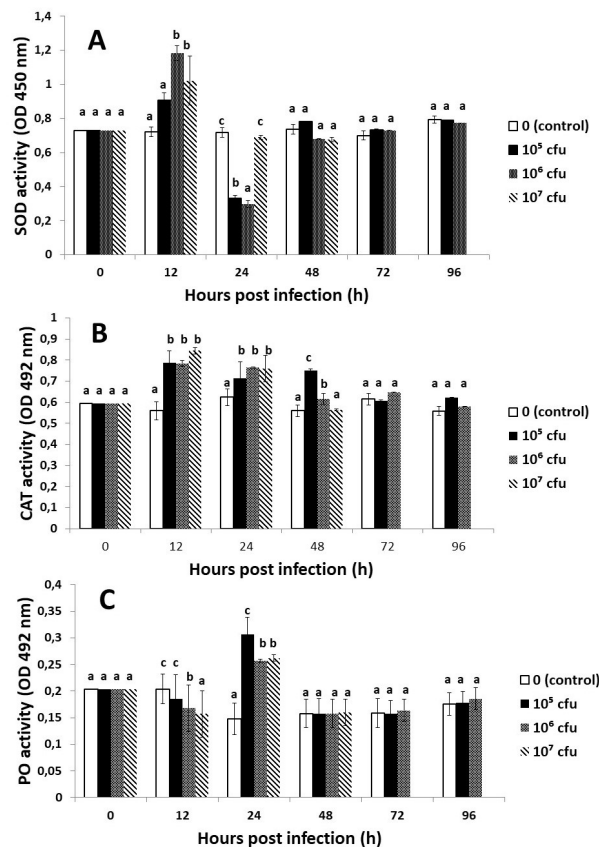


FIGURE 2 Antioxidant enzyme activity of superoxide dismutase (A), catalase (B) and phenol oxidase (B) and of abalone *H. squamata* at various *V. alginolyticus* infection as indicated. Different lowercase letters on each observation indicates a significant difference of the enzyme activity ($P < 0.05$) compared with the control using Tukey's test.

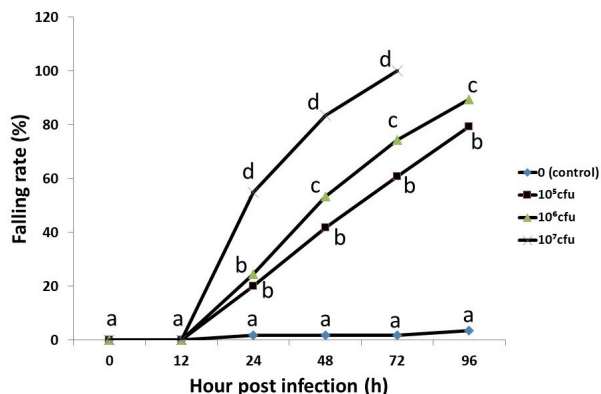


FIGURE 3 Falling rate (%) at different observation of abalones *H. squamata* on various *V. alginolyticus* infection as indicated. Different lowercase letters on each observation indicates a significant difference of the falling rate ($P < 0.05$) compared with the control using Tukey's test

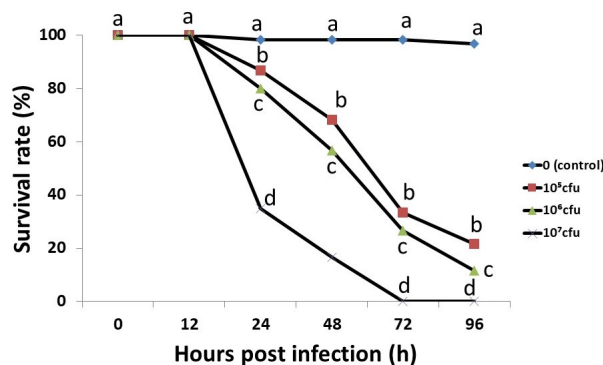


FIGURE 4 Survival rate (%) at different observation of abalones *H. squamata* on various *V. alginolyticus* infection as indicated. Different lowercase letters on each observation indicates a significant difference of the survival rate ($P < 0.05$) compared with the control using Tukey's test.

307 compared to the control. At 96 hpi, the SOD activity
 308 of all infection treatments was the same level with control
 309 (Figure 2A). The CAT enzyme activities significantly increased
 310 in all of the infection treatments from 12 hpi until
 311 48 hpi ($P < 0.05$). Then CAT activity tended to decrease
 312 and reached the normal condition at 72 hpi (Figure 2B).
 313 The PO activity of infection treatments was significantly
 314 decreased at 12 hpi ($P < 0.05$) compared to the control in
 315 24 h. After 24 h, the PO activity tended to decrease and
 316 reached same level of expression with control at 48 hpi
 317 (Figure 2C).

318 **3.3. Physiological responses of *H. squamata* to *V. al-***
 319 ***ginolyticus* infection**

320 The adhesion of abalones to the substrate was an important
 321 end-point of their health and protection from environmental
 322 threats. The falling rate of the substrate was observed
 323 in *H. squamata* after it was exposed to various level of
 324 bacterial densities (Figure 3). On the other hand, there
 325 was no statistically significant difference of the falling rate
 326 abalones *H. squamata*'s substrate in the control group.
 327 However, abalones at the concentration of 10^5 cfu, 10^6 cfu,
 328 and 10^7 cfu treatments at 48 hpi was showed falling rates
 329 as 50%, 70% and 100% respectively. Furthermore, after
 330 72 h, the falling rates were 80% at the concentration of 10^5
 331 cfu and 90% for 10^6 cfu treatments.

332 In this infection experiment of *V. alginolyticus*, at the
 333 concentration ranging from 10^5 to 10^7 cfu, the mortality
 334 was started at 24 hpi in all infected treatments. The survival
 335 rate of 10^7 cfu infection treatment was decreased significantly
 336 and all of the abalones were deceased by 72 hpi. Moreover,
 337 at the concentration of 10^5 and 10^6 cfu infection
 338 treatments, 21.6% and 11.6% of animals still alive at
 339 96 hpi. As shown in Figure 4, in the control group the
 340 mortality of abalones was not observed and a 96.6% of
 341 survival rate was achieved until 96 hpi. Taken together,
 342 all those data suggested that LD50 of this *V. alginolyticus*
 343 in *H. squamata* with an average weight of 5.13 g was

less than 10^5 cfu. This result indicated that *V. alginolyticus*
 is a virulent bacterium against abalone *H. squamata*.
 Liu et al. (2000) reported that LD50 of *V. parahaemolyticus*
 on abalone *H. diversicolor supertexta* weighing 10–14 g
 is 1.6×10^5 cfu, and mortalities occurred within 2 d of
 infection. While Liu et al. (2001) reported that LD50 of
V. alginolyticus strain H-11 on abalone *H. diversicolor*
supertexta is 3.6×10^5 cfu. Therefore, the result of this study
 is in a good agreement with the previous report.

3.4. Histology changes of *H. squamata* in response to
***V. alginolyticus* infection**

The abalone attaches and moves using its foot muscles
 along to the substrate for feeding and other activities. Due
 to its vital role, therefore in this study the effect of *V. al-*
ginolyticus infection on the foot muscle was investigated,
 the histology was conducted in this study. The normal foot
 of *H. squamata* consisted of an epithelial layer (EL), connective
 tissue layer, and muscle layer (ML) in a crosssection view
 (Figure 5A). The epithelial layer included mucous cells,
 eosinophilic granule cells (Egc), and melano granule cells
 (Mgc) (Figure 5A). The muscle layer was broad and
 consisted of muscle fiber bundles (Mfb) and hemolymph
 sinus (Hs) (Figure 5A). Muscle fiber bundles distributed
 evenly to fulfilled the muscle layer as longitudinal fibers.

In the 1×10^5 cfu infection treatment, the structural
 was changed in the abalone foot included small abscess
 (Abs) in the muscle layer, vacuolation, and enlargement
 of hemolymph sinus (Hs) in the muscle layer (Figure 5B).
 Whereas in the 1×10^6 cfu infection treatment, many
 abscesses (Abs) both in the intra epithelial layer and muscle
 layer, and enlargement of hemolymph sinus (Hs) in the
 muscle layer were observed (Figure 5C). In addition, in
 the 1×10^7 cfu infection treatment, the structural changes
 in the abalone foot included intrusion of hemolymph through
 the hemolymph sinus and moving closed to the epithelial
 layer post the enlargement of hemolymph sinus (Hs) and
 decreasing the density of muscle fiber bundles in the mus-

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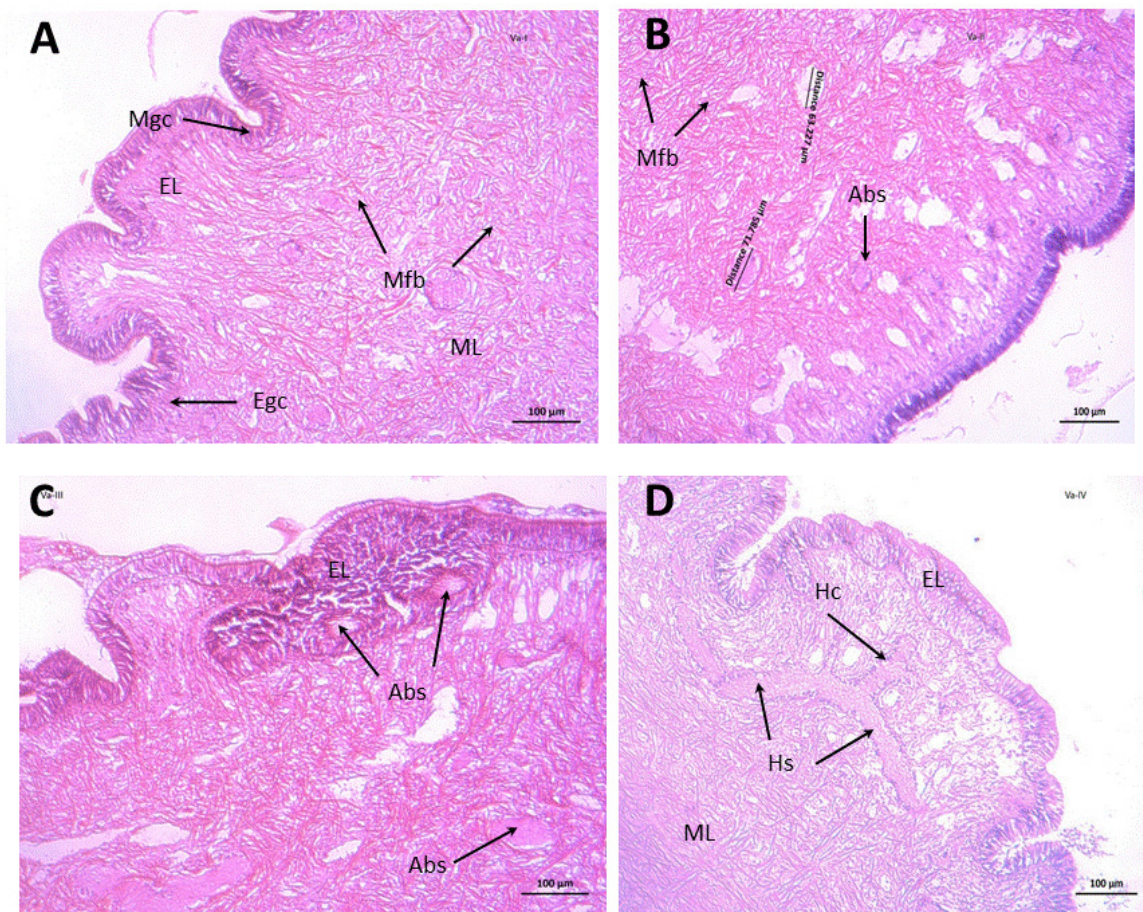


FIGURE 5 Histological section of abalone *H. squamata* foot (100x magnification). Control treatment (A), infected with *V. alginolyticus* at concentrations of 1×10^5 cfu (B) 1×10^6 cfu (C) and 1×10^7 cfu (D). Vacuolations, enlargement of hemolymph sinuses and abscesses were found in infection treatments. ML=muscle layer; Hs= hemolymph sinus, EL=epithelial layer, abscesses (Abs).

cle layer (Figure 5D). Thus, the infection of *V. alginolyticus* induced enlargement of hemolymph sinus, development of abscess intra epithelial and intramuscular and the intrusion of hemolymph closed to epithelial layer due to the disintegration of the epithelial layer and muscle layer of abalone foot tissues. The histological alterations of the foot in abalones more severe with increasing bacterial concentration.

3.5. Discussion

Abalone species are economically valuable for fishery production in the temperate or sub-tropical areas, so that the commercial aquaculture of abalone has developed in many countries. However, this industry has faced several problems, and one of the most important problems is diseases (Hsu and Gwo 2017). *Vibrio* spp. bacteria have been identified as pathogenic bacteria that causes diseases in many species of abalone and it can lead to the economic significant losses (Liu et al. 2001; Cheng et al. 2004; Cai et al. 2006b). Several parameters of disease mechanism have been investigated on disease-related gene expression, enzyme activity of abalone (Rungrasamee et al. 2010; Di et al. 2016; Fang et al. 2019; Yao et al. 2019). However,

the study on those subjects is very limited in Indonesia. This study is the first investigation the effect of *V. alginolyticus* infection on Indonesian abalone *H. squamata* with a comprehensive evaluation of mortality, histology, enzymes activity, and *HSP* genes expression.

The large size of abalones are distributed in temperate seas, while the small size abalones are distributed at wide-range geographical distribution in warm water, including Indonesia. The commercial aquaculture of the small size has developed well, especially in East Asia (Hsu and Gwo 2017). The aquaculture industry has been overwhelmed with its share of diseases and problems caused by several pathogens (Bondad-Reantaso et al. 2005). In Taiwan, the production of small abalone has dramatically decreased in the past 15 years which caused by lack of suitable diatom feed for larvae, poor water quality, habitat degradation, and genetic problems, disease and infection problems (Hsu and Gwo 2017).

Heat shock protein (HSP) family forms the most ancient defense system in all living organisms, from bacteria to humans. Heat shock proteins are classified into six major families: small *HSPs*, *HSP40*, *HSP60*, *HSP70*, *HSP90*, and *HSP110* according to their molecular weight. Among

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these HSPs, HSP70 and HSP90 are common and widely studied heat-related proteins (Wang et al. 2004; Xie et al. 2015). On abalone *Haliotis diversicolor*, the HSP was expressed in the mantle, mucous gland, muscle, gills, digestive tract, hemocytes, and hepatopancreas tissues. However, the expression level of HSP differed among tissues with a significantly higher expression level being in hepatopancreas, followed by hemocytes (Fang et al. 2019). In this experiment, we studied the expression HSP on abalone form hemocytes. Taking samples as hemocytes has advantages, due to its easiness to get hemocytes from animals and its low risk to abalone conditions.

In channel catfish (*Ictalurus punctatus*), Heat Shock Protein (HSP) genes were differentially expressed after *Edwardsiella ictaluri* or *Flavobacterium columnare* bacterial infections. The expression of those genes exhibited both temporal and spatial regulation. The induction of HSP genes was observed soon after bacterial infection, suggesting their distinct roles in immune responses and disease defenses (Xie et al. 2015). The expression level of HSP70 and HSP90 of *Penaeus monodon* genes have been reported significantly increased after a 3-hour exposure to *V. harveyi* (Rungrassamee et al. 2010). In this study, we evaluate the expression of HSP90 and HSP70 of abalone *Haliotis squamata* in response to *Vibrio alginolyticus* infection at 12 hpi.

In this study, the expression of HSP90 was upregulated in 12 hpi in all doses infection and reached the highest up-regulation more than four times at the treatment of 10^7 cfu infection compare to the control. The HSP90 expression was then down-regulated after 24 hpi to one-seventh to one seventeenth for infection treatments compared to the control (Figure 1). This expression pattern was similar to study of Wang et al. (2011) that transcription of HSP90 of disk abalone (*H. discus*) gene in response to bacterial LPS challenge significantly increased within 2 h and reached highest transcription at 4 hpi, then recovered to the normal level of transcription in 24 h finally. The low expression of HSP90 on high density infection (10^7 cfu) may occur due to the severe condition abalone, which leads to mortality.

Heat-inducible forms of HSP70 play a central role in stress tolerance by the promotion of growth at moderately high temperature and/or protecting organisms from death at extreme temperature (Cheng et al. 2007). HSP70 has been reported exhibits physiological and ecological importance in response to pathogen infection and environmental stress. For example, heat shock in fish was the most effective stress stimuli to induce HSP70 response compared to other stressors including hypoxia and air exposure. In mollusks, HSP70 transcripts increased significantly after acute heat stress. Up-regulation of HSP70 was observed after *V. parahaemolyticus* infection in adult bay scallops *Argopecten irradians*. The expression of HSP70 in the zebra mussel *Dreissena polymorpha* showed a time-dependent increase after lipopolysaccharide (LPS) stimulation (Fang et al. 2019).

The expression levels of HSP70 and HSP90 of *Penaeus monodon* significantly increased after a 3-h expo-

sure to *V. harveyi* (Rungrassamee et al. 2010). The bacterial challenge of *V. anguillarum* on Pacific abalone (*H. discus hannai*) showed a time-dependent expression of the HSP gene with a significant increase in the expression of HSP70 mRNA and reach the highest at 124 h and expression level of HSP70 returned to about control levels following a 96-hour recovery period (Cheng et al. 2007). Different from the result of Cheng et al. (2007), the relative expression level of HSP70 in this study decreased rapidly in 12 h after *V. alginolyticus* infection. This result may be due to quick expression of the HSP70 and the peak of the expression was less than 12 h. Wang et al. (2011) noted that in response to the LPS challenge, the transcription of disk abalone HSP90 gene significantly increased within 2 hpi and approached maximum induction at 4 hpi. Due to the earliest analysis of HSP in this study was at 12 hpi, the expression of the HSP at this time was already decreased.

In this study, the superoxide dismutase (SOD) activity of *H. squamata* in response to *V. alginolyticus* infection was increased at 12 hpi and then decreases at 24 hpi followed with normal expression started on 48 hpi (Figure 2A). Di et al. (2016) found activity SOD of *H. diversicolor* with the withering syndrome was significantly lower than in the healthy abalone. Catalase activities of infected abalone was started from 12 hpi then it was likely to decrease and there was no significant different among treatment after 72 hpi (Figure 2B). Buss et al. (2017) found that catalase CAT activity of greenlip abalone (*H. laevigata*) significantly higher when reared at 25 °C. Different from the expression of SOD and CAT which showed a significant increase within 12 hpi, the expression of phenol oxidase was increased after 24 hpi (Figure 2C). The increasing phenol oxidase (PO) activity in *H. diversicolor* was stimulated by a viral infection (Yao et al. 2019).

Several cases of mass mortality of abalone have been recorded from several countries. Mortality of Japanese abalone *Sulculus (Haliotis) diversicolor supertexta* in Kanawaga, Japan in June to October 1997 is caused by *Vibrio carchariae (V. harveyi)* (Nishimori et al. 1998). At the nearly same time, mass mortality of the abalone *Haliotis tuberculata* L. has occurred in the natural environment along the south coast of Brittany, French in 1997 also caused by *V. carchariae (V. harveyi)* (Nicolas et al. 2002). Mass mortality among cultured small abalone *H. diversicolor supertexta* with abscess/ulcers in the mantle in 1998 at Kao-Hsiung Taiwan was caused by *V. alginolyticus* (Liu et al. 2001). In China, *V. alginolyticus* and *V. parahaemolyticus* were associated with a severe epidemic in farmed *H. diversicolor supertexta* in Fujian Province (Zhang et al. 2001), and a *Vibrio harveyi*-related species was linked with the mass mortality of farmed adult *H. diversicolor* in Fujian (Jiang et al. 2013). Those data supported that *Vibrio* spp caused diseases on abalones. In this study, we reported that *V. alginolyticus* caused disease on tropical abalone (*H. squamata*) (Figure 4). This is the first report on confirmation of the pathogenicity of *V. alginolyticus* on *H. squamata*. *Vibrio* spp. has been reported as virulent bacteria to abalone. Liu et al. (2000) reported

542 that LD50 of *V. parahaemolyticus* on abalone *H. diversicolor*
 543 *supertexta* weighing 10–14 g is 1.6×10^5 cfu, and
 544 mortalities occurred within 2 days of infection. Cai et al.
 545 (2006a) reported that *V. alginolyticus* Strain 19 was vir-
 546 ulent to abalone postlarvae with an LD50 value of 1.00
 547 $\times 10^4$ cfu. Liu et al. (2001) reported that LD50 of *V. al-*
 548 *ginolyticus* strain H-11 on abalone *H. diversicolor super-*
 549 *texta* is 3.6×10^5 cfu. In this study, we also confirmed
 550 that *V. alginolyticus* was virulent with LD50 on *H. squa-*
 551 *mata* with an average weight of 5.13 g is less than 10^5 cfu
 552 (Figure 4).

553 *Vibrio* spp. produced and released toxins from the cells
 554 as an extra cellular product (ECP). The LD50 of ECP of *V.*
 555 *alginolyticus* strain H-11 on small *H. diversicolor super-*
 556 *texta* is 2.96 µg protein/g body weight (Liu et al. 2001),
 557 while *Vibrio* strain B4 has LD50 of CPS as 7.58 µg pro-
 558 tein g⁻¹ body-weight. This toxin caused several changes
 559 in abalone organs and tissues and led to mortality. In this
 560 study, the infection of *V. alginolyticus* caused histologi-
 561 cal changes as enlargement of hemolymph sinus, develop-
 562 ment of abscess intra epithelial and intramuscular, and the
 563 intrusion of hemolymph closed to epithelial layer (Figure
 564 5).

565 In this study, we showed that infection *V. alginolyticus*
 566 will be responded by *H. squamata* with the rapid increas-
 567 ing level of *HSP70* and *HSP90* expression, then it was fol-
 568 lowed by decreasing level of *HSP70* and *HSP90* expres-
 569 sion. Similar responses were occurred on anti-oxidant ac-
 570 tivity of SOD and CAT enzymes with delay time. Those
 571 conditions caused histological change in the tissues and
 572 led to mortality.

573 4. Conclusions

574 We conclude that Pisang Kepok from Bandung (West
 575 Java), Denpasar (Bali), Flores (East Nusa Tenggara), and
 576 Sorong (Papua) are the same cultivar as indicated by the
 577 identical ITS2 length, structure, and secondary structure.
 578 This also suggests that the distribution of Pisang Kepok
 579 to these four islands in Indonesia took place a short time
 580 ago and possibly was introduced by humans for agricul-
 581 tural purposes. Furthermore, it indicates that they might
 582 have the identical superior traits of *Fusarium* and drought
 583 tolerance, which are useful for the improvement of other
 584 banana cultivars. ITS2 has been illustrated in many studies
 585 as a robust molecular marker to some degree, particularly
 586 in this research, in clustering the bananas based on their
 587 genome group. Nevertheless, the use of ITS2 in the minor
 588 taxonomical groups such as cultivar could be meticulous
 589 and needs extra details. Provided that cultivar 'Kepok'
 590 might possess desirable traits such as tolerance to drought
 591 and disease such as in Foc TR1, with more research, it
 592 assumed that these superior traits might provide the im-
 593 provement of other banana cultivars.

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 599

Authors' contributions

600 NSY, M, AI, NSN designed the study. NSY conducted
 601 animal handlings and physiological studies. LA carried
 602 out the molecular laboratory work. NSY, GT analyzed the
 603 data. NSY, M, AI, NSN wrote the manuscript. All authors
 604 read and approved the final version of the manuscript.
 605

Competing interests

606 The authors declare no competing interest.
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