The application of pigment-protein fraction from *Nannochloropsis oculata* on β-actin response of *Cromileptes altivelis* infected with viral nervous necrosis

Aplikasi fraksi protein pigmen *Nannochloropsis oculata* terhadap respons β-aktin pada ikan kerapu tikus yang diinfeksi viral nervous necrosis

**ABSTRACT**

β-actin is a prominent protein in the immune system. An outcome from gene transcription, the protein protects against pathogen infection, such as receptor clustering, antigen internalization, and regulating vesicle for antigen processing. β - actin expression will determine the success of immune response of the organism. This study aimed to understand the role of pigment-protein fraction (PPF) from *Nannochloropsis oculata* in the increase of β-actin expression in humpback grouper *Cromileptes altivelis* infected by viral nervous necrosis (VNN). The experiment was performed with one negative control (A: normal fish) and three PPF treatments (B: fish + PPF, C: fish + VNN, and D: fish + PPF + VNN). PPF was applied through the sonde method and analyzed with immunohistochemistry technique and Immunoratio software. The results showed that PPF was able to increase β-actin expression on all treatments: A (34.9%), B (38.1%), C (39.1%), and D (51.6%). It demonstrated PPF ability to induce the increase of β-actin expression indicating an improved defense of humpback grouper *C. altivelis* against VNN infection.

Keywords: β-actin, pigment-protein fraction, *Nannochloropsis oculata*, VNN

**INTRODUCTION**

The humpback grouper is one of the most important commodity in Indonesian aquaculture due to its high economic value, priced up to Rp34,000,000 for 2015 export starting at Rp500,000,-/kg from the farmers (Ariyanti, 2015). The high price drive farmers to start culturing this fish, albeit they are faced with several challenges in the process. The biggest challenge in a humpback grouper culture is the intracellular parasite or virus. Viral nervous necrosis (VNN) contribute a large portion of the failure of humpback grouper culture. It is capable of causing tissue damage, hyperplasia, necrosis, vacuolation and high mortality up to 100%
This virus infected early stages of larvae and juveniles focusing on the eye and brain nerve system with distinctive symptoms such as whirling, sleeping sickness, and abnormal behavior.

Regulation of immune system mechanism is vital for the success against VNN infection. One of the protein with significant function of the immune system is β-actin (Jönsson et al., 2012). This protein is a cytoskeleton component which functions to regulate cell morphology, receptor clustering, antigen internalization, and regulating vesicle for antigen processing. A deficiency in cytoskeleton actin regulation will cause a disturbance in immunological synapse formation, the activation of T and B cells (Yuseff and Reversat, 2011). Furthermore, β-actin is also involved in an important nucleus mechanism e.g. transcription, mRNA export, and chromatin renovation (Zheng et al., 2009; Spencer, 2011). In the nucleus, β-actin binds with RNA polymerase II and III (Tang et al., 2009; Spencer, 2011).

*Nannochloropsis oculata* is a single cell microalgae containing pyrenoid and numerous pigments with anti-bacterial and anti-viral properties, namely chlorophyll a, β-carotene, violaxanthin, and veucherxanthin (Cao et al., 2013). *N. oculata* is also rich on nutritional value of the following: protein 52.11%; carbohydrate 16.00%, and lipid 27.64% consisting of eicosapentaenoic acid (EPA) up to 31.42% and 3.94% arachidonic acid (ARA/AA) (Bentley et al., 2008). *Nannochloropsis oculata* used in this study was acquired from BBAP Situbondo, filtered until 150 g wet paste was obtained. The isolation method referred to Yanuhar (2015), where the cells were homogenized in a mortar for one hour with liquid nitrogen addition. 8 mL from 50 mM glycine and 20 mM KCl (pH 7.5) were added then centrifuged at 12,000 rpm for 60 min at 4 °C temperature. The supernatant was then added to solid ammonium sulphate (SAS) solvent gradually to get a final concentration of 30% v/v and centrifuged was performed at 15,000 rpm, 4 °C for 30 min. A dialysis bag was sterilized by boiling in 0.1 mM Tris-EDTA (pH 7.3) for ten min. Sample of 2,000 mL was treated with dialysis using 20 mM Tris-HCl, pH 8.0 for 24 hours at 4 °C temperatures with continuous mixing. Afterwards, the sample was filtered by amillipore filter (0.22 µm, Sartorius). The dialysis and filtration process were repeated once. Protein content was then measured using a nanodrop spectrophotometer (NanoDrop Technology, Wilmington, USA) at 280 nm. One absorbance corresponds with 1 mg/mL protein.

In this study *Nannochloropsis oculata* was used as an inducer, a compound activating β-actin, specific genes in protein expression. This particular protein played a role in increasing the humpback grouper *C. altivelis* immune system against VNN infection. The inducing effect is highly dependent on the target cell’s ability to recognize the inducer as a protein molecule (Ringo et al., 2011). Therefore, this study was aimed to understand the role of pigment-protein fraction (PPF) in increasing β-actin response on humpback infected VNN.

**MATERIALS AND METHODS**

**Experimental design**

This study used descriptive experimental design dividing the fish into four groups, A: negative control (grouper fed with normal feed/no PPF and viral infection); B (fish + PPF): the grouper was fed and given PPF based on the body weight; C (fish + VNN): the grouper was fed and infected with VNN corresponding to its body weight, infection was conducted by giving feed mixed with VNN positive fish; D (fish + FPP + VNN): the grouper was treated with PPF and infected with VNN.

**Pigment protein fraction (PPF) isolation from *Nannochloropsis oculata***

*N. oculata* used in this study was acquired from BBAP Situbondo, filtered until 150 g wet paste was obtained. The isolation method referred to Yanuhar (2015), where the cells were homogenized in a mortar for one hour with liquid nitrogen addition. 8 mL from 50 mM glycine and 20 mM KCl (pH 7.5) were added then centrifuged at 12,000 rpm for 60 min at 4 °C temperature. The supernatant was then added to solid ammonium sulphate (SAS) solvent gradually to get a final concentration of 30% v/v and centrifuged was performed at 15,000 rpm, 4 °C for 30 min. A dialysis bag was sterilized by boiling in 0.1 mM Tris-EDTA (pH 7.3) for ten min. Sample of 2,000 mL was treated with dialysis using 20 mM Tris-HCl, pH 8.0 for 24 hours at 4 °C temperatures with continuous mixing. Afterwards, the sample was filtered by amillipore filter (0.22 µm, Sartorius). The dialysis and filtration process were repeated once. Protein content was then measured using a nanodrop spectrophotometer (NanoDrop Technology, Wilmington, USA) at 280 nm. One absorbance corresponds with 1 mg/mL protein.

**PPF in-vivo test on humpback grouper**

In this study PPF was given to the fish from the first day of rearing, while VNN infection was conducted on day 14. PPF was given earlier to monitor its activities in enhancing fish immune system. Thus, it is expected that when challenged against VNN the immune system would be ready prepared for the infection.

Prior to treatments, fish were acclimatized for one week. On treatments B, C, and D, PPF was given with sonde method for six times on day 1, 5, 9, 14, 19, and 24; each application dosage was adjusted to fish body weight at 306 µL, 315 µL, 322 µL, 326 µL, 345 µL, and 351 µL. On treatment D, VNN infection was performed using VNN positive fish meat mixed with fresh feed and fed *ad libitum* on day-14, 19, and 24. On day-
27 the target organ, which was the main part of
the central nervous system was removed through
dissection.

Brain isolation of humpback grouper

On the organ isolation process, post in vivo
test fish was sedated with clove oil at 2 mL per 10
L seawater. When the fish was unconscious, it is
dissected and the brain was removed. Then it was
immersed in liquid nitrogen and preserved in the
liquid nitrogen tank.

Protein analysis with sodium dodecyl sulfate
polyacrylamide gel electrophoresis (SDS Page)

Protein analysis was conducted with SDS-
Page Laemmli electrophoresis (Laemmli,
1970). The electrophoresis gel used was sodium
dodecyl sulfate polyacrylamide gel with 12.5%
concentration. Analysis was conducted by adding
the buffer sample into protein sample (1:1 ratio)
in an eppendorf tube and heated at 100 °C
temperature for five min in 5 mm Tris HCl solvent
at pH 6.8; 5% 2-mercapto ethanol; 2.5% w/v
sodium dodecyl sulfate, and 10% v/v glycerol. It
was then cooled and kept at 20 °C until further
analyses. Analysis was performed at a constant
current 20 mA for 40–50 min or the tracking dye
was 0.5 cm from the gel bottom. Then, the sample
was dyed with commsasie brilliant blue. The last
step was de-staining to remove gel colorant and
emphasize the protein band.

Immunohistochemistry

Immunohistochemistry dye was used in
accordance to a study by Khan et al. (2014).
VECTASTAIN® ABC kit (ABC-Elite, Santa
Cruz, Vector Laboratories) was used to detect
biotin using chromogen (diaminobenzidine
tetrachloride). Tissue sample was dehydrated
with alcohol and cleaned with xylene. Peroxidase
dendogenous enzyme was cooled with 3% hydrogen
peroxide and methanol at room temperature for
30 min in no light condition. The microwave
antigen was retrieved with 0.01 mol/L Sodium
citrate buffer (pH 6.0) then incubated in antibody
monoclonal β-actin (AC-15) for 16 hours at 4 °C
with 1:1,000 dilution. Afterwards, the secondary
antibody biotin conjugated anti-IgG was added
for 30 min at room temperature.

Immunohistochemistry analysis

β-actin profile can be visualized using the
immunohistochemistry (IHC) method. In this
study, IHC was used to examine the β-actin
profile on brain related to antigen and antibody
bonds. Indirect method was used with the primary
and secondary antibody. The primary antibody
was the monoclonal β-actin (AC-15) while the
secondary was IgG antimouse conjugated protein.

Immunohistochemistry result was visualized
using Olympus microscope. It was then
analyzed using Immunoratio (IR) online
software (Tuominen et al., 2010) to determine
the percentage of β-actin expression showed as
various colors. ImageJ software was also used
to analyze the target gene (β-actin) intensity
as previously demonstrated by Varghese et al.
(2014). Histogram scoring was based on pixel
intensity, where 0 represent the darkest color
while 255 represent the color white. This range
was divided into four zones: 0–60 strong positive,
61–120 positive, 121–180 weak positive, and
181–255 negative.

Antibody response confirmation with Western
blotting

Further analysis with Western blot method
was performed to confirm β-actin expression on
fish brain. In this study, a specific antibody, anti-
β-actin was used to confirm the presence of the
target gene or protein (β-actin) in the sample. The
Western blotting analysis was performed based
on the Towbin method (1979). Gel was placed on
a nitrocellulose paper using semi-dry blotter from
biorad after the protein analysis with SDS-Page.
The electricity current used in this process was
300 mA for 30 min. Then, the sample was dyed
with ponceau 2% containing 3% trichloracetic
acid (TCA) to detect protein movement to the
nitrocellulose paper and marked to determine its
molecular weight. The nitrocellulose paper was
then cut following the well pattern.

To remove the dye, it was rinsed with dH2O
and blocked with TBE (Tris/Borate/EDTA)
contains 3% albumin) at pH 7.4 added with 1%
bovine serum albumin (BSA) and shaken for two
h. The sample was rinsed again twice at five min
interval using TBE pH 7.4 solvent containing
0.05% Tween 20. Then secondary antibody
anti fish IgG mice was added with 1/1,000
concentration to TBE pH 7.4 and BSA 1%,
which was protected from rays’ influence. The
sample then shaken for two hours and underwent
two five-min rinsing processes with TBE pH 7.4
Tween 20 0.05%. Tablet Cip diluted in H2O 10 mL
was used as staining material. The solution then
poured into nitrocellulose paper and observed for
the appearance of red color. If the reaction was
sufficient with \( H_2O \) rinsing, drying process was continued with filter paper.

**Data analysis**

Data analysis was performed with descriptive quantitative method by describing the results and quantifying immunohistochemistry analysis result with immunoratio software.

**RESULTS AND DISCUSSION**

**Result**

*Nannochloropsis oculata* PPF analysis

PPF isolation result using SDS-Page method is presented in Figure 1. Based on molecular weight identification above, it was indicated that *N. oculata* microalgae contained violaxanthin chlorophyll protein (VCP) and peridinin chlorophyll protein (PCP). The indication was based on several previous microalgae protein researches which concluded protein with molecular weight of 22 kDa was VCP (Sukenik *et al.*, 1992; Basso *et al.*, 2014), and of 14 kDa was PCP (Weis *et al.*, 2002).

**Humpback grouper brain’s protein profile**

Protein profile analysis result using SDS-Page showed that there were protein profile differences in each treatment (Figure 2). Different expressions emerged from each treatments in Figure 2 showed different responses on fish test samples. The focus in this research was β-actin protein. Jovčevska *et al.* (2014) stated that β-actin molecular weight was 42 kDa. According to Figure 2, there was a protein band with molecular weight of 42.4 kDa. That finding became an early indication of β-actin expression existence from fish brain samples in every treatment.

Observation result is presented in Figure 3. According to Figure 3, it was concluded that β-actin was expressed in every treatment. The result also showed that fish expressed β-actin in normal circumstances.

**β-actin profile in control fish**

β-actin profile observation was also performed in the group of control fish. The result would be a standard β-actin profile to be compared to other treatments shown in the Figure 4. On the control fish group, the fish brain was in a normal condition shown by normal tissue cells (Figure 4b). IR analysis on treatments were shown with DAB value of 34.9% (Figure 4a). The percentage showed that 34.9% target gene (β-actin) on the control fish was detected, indicated with an orange color which confirmed antigen and antibody bond.

![Figure 1. SDS-Page pigment-protein fraction result](image)

**Table 1. PPF of *Nannochloropsis oculata* identification based on protein molecular weight**

| Sample’s molecular weight (kDa) | Molecular identification | Reference                      |
|--------------------------------|--------------------------|--------------------------------|
| 88.3                          | -                        | -                              |
| 51.2                          | -                        | -                              |
| 41.5                          | -                        | -                              |
| 22.1                          | violaxanthin chlorophyll protein (VCP) | Sukenik *et al.*, 1992; Basso *et al.*, 2014 |
| 20.4                          | -                        | -                              |
| 14.6                          | peridinin chlorophyll protein (PCP) | Weis *et al.*, 2002; Yanuhar, 2011; Komariyah, 2014 |
| 8.8                           | -                        | -                              |
Figure 4b also showed that β-actin was expressed at cell membrane and cytoplasm. It could be seen that normal fish also express β-actin. Figure 4c is a brain histogram showing the intensity of target gene appearance. Also, it showed that the target gene was categorized as strong positive within 50–60 range point.

**β-actin of PPF induced brain organ**

Upon PPF induction, the brain showed a difference compared to those of the control fish (Figure 5). On Figure 5a the distinct brownish yellow color indicates a bond between antigen and antibody. Image analysis with Immunoratio showed a value of 38.1% (Figure 5b). The number explained that 38.1% β-actin is expressed on fish treated with PPF. The histogram analysis (Figure 5c) also showed that the highest color intensity was 58–65, categorized as positive.

**β-actin of brain organ infected with VNN**

Immunohistochemistry observation in this study confirmed that VNN infection on humpback grouper has a capability to cause cell and tissue damage. Visualization on VNN infected organ was provided in Figure 6. The damage in cell and tissue comprise of vacuolization (black arrow) and necrosis (yellow arrow) (Figure 6a). In Figure 6b, some cells were reduced and shrunk (green arrow) indicating apoptosis, severe cell damage
Figure 4. Control humpback grouper brain profile: (a) immunoratio (34.9%), (b) immunohistochemistry (IHC) result, (c) histogram (ImageJ). Note: arrows indicate β-actin expression.

Figure 5. Humpback grouper brain profile with pigment protein fraction application: (a) immunoratio (38.1%), (b) immunohistochemistry (IHC) result, (c) histogram (ImageJ); note: arrows indicate β-actin expression.
that are no longer repairable. Immunoratio analysis (Figure 6a) showed a \( \beta \)-actin percentage of 39.1\%. \( \beta \)-actin was also expressed at a small intensity on membrane and cytoplasm (Figure 6b), while histogram analysis displayed a positive category in target gene intensity with the highest value of 61–64.

**\( \beta \)-actin of PPF induced and VNN infected brain organ**

The fish died after two weeks when infected with virus but survived until the end of experiment when PPF was applied prior to VNN infection. IHC analysis also showed an increase in \( \beta \)-actin expression compared to control at 51.6\% (Figure 7a). Figure 7b displayed \( \beta \)-actin on almost every cell part, membrane, cytoplasm, and nucleus. Histogram in Figure 7c also showed target gene intensity in the strong positive group at 31–44.

**Discussion**

\( \beta \)-actin is a central component for body cellular mechanism within the organism. Based on gene function, it is regarded as the internal cell regulating gene (Pollard & Cooper, 2009). This gene has a vital role in immune system against pathogen infection, namely receptor clustering, antigen internalization, and regulating vesicle for antigen processing.

One of the natural ingredient that can be used to manage the humpback grouper health is *N. oculata* (Yanuhar, 2009; Yanuhar *et al.*, 2011; Yanuhar *et al.*, 2012) which contain PPF as anti viral and bacterial agents. When induced to *C. altivelis* brain, PPF gave a significant effect on \( \beta \)-actin expression. Observation at \( \beta \)-actin expression on *C. altivelis* brain increased from the negative control (34.9\%), FPP induced (38.1\%), VNN infected (39.1\%), to FPP induced and infected with VNN (51.8\%). The increase indicated an effective immune response from the fish. VNN infection itself has been a serious problem for humpback grouper culture due to its capability to cause cell and tissue damage (Yanuhar *et al.*, 2012; Yuwanita & Yanuhar, 2013; Costa & Thompson, 2016).

The experiment showed an increase in \( \beta \)-actin expression when being induced with PPF. Fish respond to VNN by escalating \( \beta \)-actin expression to organize and proliferate immune system to face virus infection. PPF role as inducer and biocatalyst for immune system development were enhanced with the increasing \( \beta \)-actin expression. Said increase is possible through the violaxanthin chlorophyll protein (VCP) and peridinin chlorophyll protein (PCP) in the microalgae. The proteins are available as enzymes and function as biocatalyst in physiological response development and are categorized as carotenoid (Takaichi, 2011).

\( \beta \)-actin was expressed in every treatment during the experiment. As a promoter gene element, it was constitutively expressed in all the cells. The expression gave a preview of better and faster immune response. This is partly due to \( \beta \)-actin housekeeping characteristic, which in normal condition was also used by the cells for cell growth, migration, and survival (Bunnel *et al.*, 2011).

The increase in gene expression indicate an effective cellular defense mechanism within the fish body. Antigen presenting cell (APC) is one of the key components in fish immune system, covering a wide range of cells including the B cells. On the B cells, \( \beta \)-actin remodeling will change its morphology enabling it to easily capture antigen (Brezski & Monroe, 2007). \( \beta \)-actin also function in regulating MHC (major histocompatibility complex) molecule movement to cell membrane (Khumaidi *et al.*, 2015). MHC is a vital component in T cell activation. Deficiency in \( \beta \)-actin expression will lower MHC presentation which eventually reduce antigen presentation to the T cells (Vascotto *et al.*, 2007). Actin remodeling is also essential for the macrophage and dendritic cells to help in motility, phagocytosis, and antigen presentation.

A rise in \( \beta \)-actin expression also increases the function of cell wall receptor in detecting antigen signal and presentation. A previous study by Komariyah (2014) on PCP application in *C. altivelis* resulted that PCP is capable of increasing toll-like receptor (TLR) expression. TLR has a role in capturing and sending downstream signal to receive a specific response (West *et al.*, 2004). The signal transmitting mechanism is affected by the presence of \( \beta \)-actin. Also, the receptor signal absorption was also affected by actin cytoskeleton bundle (Irving *et al.*, 2012).

PCP has been applied in inducing antivirus gene on *C. altivelis* as P56/ISG56, which is capable of blocking VNN protein synthesis. To maintain its function, P56/ISG56 depends on an effective regulation from cytoskeleton actin. PPF application on humpback grouper specifically regulate TNF and IL-6 expression indicating an increase in fish immune. The expression of antivirus genes and cytokine is
Figure 6. Brain profile of viral nervous necrosis-infected humpback grouper: (a) immunoratio (39.1%), (b) immunohistochemistry (IHC) result, (c) histogram (ImageJ). Note: red arrow: β-actin expression, black arrow: vacuolation, yellow arrow: necrosis, green arrow: apoptosis.

Figure 7. Brain profile of PPF induced and viral nervous necrosis-infected humpback grouper: (a) immunoratio (51.6%), (b) immunohistochemistry (IHC) result, (c) histogram (ImageJ). Note: β-actin expressed throughout the cell (red arrow).
mainly due to β-actin effect (Yanuhar, 2015). β-actin has a vital role in remodeling chromatin group, RNA polymerase complex, and some ribonucleic protein (McDonald et al., 2006). Saha et al. (2006) reported that this gene utilized cytoskeleton actin function to reach the blocking site of virus protein synthesis. 

β-actin increase will also elevate the regulation of gene synthesis and expression. A report by Visa and Percipalle (2010) stated that β-actin has an essential role in gene expression as a component of chromatin modification complex. In gene expression, β-actin function was connected to RNA Polymerase I, II, and III (Almuzzaini et al., 2016; Serebryannyy, 2016); Hu et al., 2004). On the contrary, Ye et al. (2008) stated that actin polymer deficiency will cause actin inability to bind with polymerase I, thus the transcription process is not supported.

When infected with VNN without PPF application, the fish showed cell damage e.g. necrosis, vacuolysis, and apoptosis (Figure 6a and 6b). The increasing β-actin expression was insufficient to prevent infection. Yuwanita and Yanuhar (2013) stated that VNN infection in humpback grouper damage the brain cells causing necrosis, vacuolization, and hypertrophy. The VNN infected fish died on the second week while fish treated with VNN and PPF survived until the last day of the four week rearing period.

CONCLUSION

Pigment-protein fraction (PPF) from the marine microalgae *Nannochloropsis oculata* induced to viral nervous necrosis (VNN) infected humpback grouper *Cromileptes altivelis* increased the fish β-actin expression. The increase indicated a reliable immune response against VNN infection. PPF application on humpback grouper culture can be utilized as immunostimulant by increasing immune response in developing antiviral genes to prevent VNN infection.

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