# The White Spot Syndrome Virus (WSSV) Load in Dendronereis spp.

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Original Paper

# THE WHITE SPOT SYNDROME VIRUS (WSSV) LOAD IN Dendronereis spp.

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### ABSTRACT

The white spot syndrome virus (WSSV), the causative agent of White Spot Syndrome Disease (WSSD), is a major shrimp pathogen in Indonesia. Dendronereis set is a ubiquitous Polychaetes and natural food of shrimp raised in brackishwater pond in Indonesia. The objective of this research is to determine the occurrence of WSSV and the viral load in Dendronereis spp. obtained from the shrimp pond. Dendronereis spp. was obtained with PVC (10 cm in diameter) from a traditional shrimp pond in Semarang vicinity. As a comparison, healthy looking Penaeus monodon was also obtained from the same pond. The occurrence of WSSV in Dendronereis spp. was determined with 1-step and nested PCR using primer for WSSV major envelope protein, VP 28. The viral load was counted with 1-step Real Time PCR. The WSSV was detected in Dendronereis spp. with 1-step and nested PCR. The point prevalence of WSSV infection in Dendronereis spp. is 90 %. The viral load ranged from 0 to 1.9 x 10<sup>4</sup> copy of DNA/µg total DNA. The viral load in Dendronereis is comparable with that of naturally infected and at carrier state P.monodon from the same pond. This is the first report of WSSV load in naturally infected Dendronereis spp.

Key words: WSSV; Dendronereis spp.; viral load

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# Introduction

Shrimp farming is an important aquaculture industry in Indonesia. For the last two decades, this industry has become source of export grnings for Indonesia that lead Indonesia to be one of top shrimp producing counties in the world (Lem, 2006; FAO, 2010). White spot syndrome disease (WSSD) is a major shrimp disease that hampered cultured shrimp production in Indonesia. It was reported for the first time in Java island in 1994 and has caused serious impact on the shrimp culture (Sunarto et al., 2004), since then WSSD has become endemic in Indonesia. In the pond, the onset of the diseases usually occurs during the second month of cultu26 (PL are 42-50). Once the disease occurs, it can cause high mortality in one week.

The causative agent is the white spot syndrome virus (WSSV), a large DNA virus belong to genus Whispoviru 13 and family Nimaviridae (Lo et al., 2012). WSSV is very

pathogenic and highly virulent on penaeid shrimp, which the most affected hosts of this virus (Hamed et al., 2005; Flegel, 2006). Moreover, it is a broad host range in addition to penaeid shrimp. WSSV DNA detected by 1-step as well as nested PCR in various benthic invertebarate cariers and vectors such as crabs (Supamattaya et al., 1998; Kanchanaphum et al., 1998; Chen et al., 2000; Meng et al., 2009; Liu, et al., 2011), rotifers (Yan et al., 2004), polychaetes (Vijayan et al., 2005), copepods (Zhang et al., 2008), mache microalgae (Liu, et al., 2007) plankton (Esparza-Leal et al., 2009), mollusk (Chang et al., 2011).

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Qualitatively, WSSV load has been classified heavy infection when WSSV DNA was detected with 1-step PCR and light infection when it was detected with nested PCR (OIE, 2006). Nowadays, Real-Time PCR was has been developed to quantitatively

determined the number of viral copy in the host (Schmitt and Anderson, 2005).

Polychaetes is a common benthic fauna in the shrimp pond. Among Nereidids polychaetes ubiquitous in shrimp pond in Indonesia is Denronereis spp.. Dendronereis spp. is potential to be infected by WSSV because it lives in the sediment, and detritofeeder. WSSV infects tissues of ectodermal and messodermal origin especially gills, the fore and hind-gut, hemopoie 10 tissue, antennal gland and lymphoid gland (Chang et al., 1996, Chen et al., 2000, Rahman et al., 2008, 133 obedo-Bonilla 2007). In the previous study we were able to detect the presence of WSSV in Dendronereis spp. from shrimp pond in Semarang vicinity with one step and nested PCR (Desrina et al., 2011). This research is to determine the WSSV load in Dendronereis spp. with natural infection with 1-step, nested and 1step RT-PCR.

### MATERIAL AND METHODS

Dendronereis spp.

The Dendronereis spp. was obtained from a traditional shrimp pond located in Semarang vicinity. The main cultivant raised is Penaeus monodon monoculture or polyculture with tilapia (Tilapia nilotica) and milkfish (Chanos chanos) with shrimp density was 2 shrimp/ m<sup>2</sup>. The pond has experienced reoccurring of WSSV infection based on farmers report and clinical signs such as lethargic, low appetite, white discoloration on the body and carapace and low number of shrimp mortality that occurred over extended period of time, usually started about one months after stocking. Three months prior to sampling, the pond experienced mass mortality and has been fallow since. The sediment was obtained using a PVC (diameter 10 cm and high 40 cm) and sieved thorough a series of sieve shaker (mesh: 2 mm,0,6 dan 0,3 mm). Dendronereis spp. was easily recognized by its bright red color and the gills. The Dendronereis spp. retained was cleaned with sterile PBS (pH 7.4) and stored in the deep freezer (-80 °C) for PCR analysis. Eight worms were randomly picked to be used in this study. Along with the worm, we also obtained three P.monodon (average weight 25 g/piece) that survived and remained in the pond, treated and stored as done on Dendronereis spp.

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Template DNA preparation for PCR analysis

A piece of the head section (first 15-20 segments, 25 mg) of frozen *Dendronereis* spp. was cut with sterile scalpel and placed in a sterile microcentrifuge tube. DN 12 xtraction and purification was done using DNeasy Blood Tissue Kit (Qiagen) according to the instruction manual. In the case of *P.monodon*, the gills was used as the source of DNA and processed as for *Dendronereis* spp..

1-step and Nested-PCR

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PCR was performed with 1  $\mu$ l of DNA termalate using Taq Polymerase, the same amount of the first-step product was used as template for nested-PCR.. WSSV DNA obtained from known infected shrimp served as positive control of the PCR process. The DNA was amplified using VP 28 (Marks et al.,, 2003) and VP28 nested primer. The PCR was done with Gene Amp PCR System 9700 (Applied Biosystem) for 30 cycles for 1-step PCR and 25 cycles for nested PCR. Primer sequence is VP28-F1 CACAACACTGTGACCAAG (Forward) and VP28-R1 TTTACTCGGTCTCAGTGCCAG (Reverse) produce amplimer 529 bp. The sequence of the second primer used in nested- PCR is VP28-F1 CATTCCTGTGACTGCTGAGG nested VP28-R1nested (Forward) and CCACACACAAAGGTGCCAAC (Reverse) produce amplimer 383 bp. Both primer used same annealing temperature 50 °C (50 sec). WSSV DNA obtained from known infected shrimp served as positive control of the PCR process. The result was visualized using UV illuminator Gel Doc XR System (Biorad).

Real Time PCR to quantify viral load

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WSSV load was quantify by ABI PRISM 7300 Real Time-PCR system (Applied Biossystem) using TAQMAN master mix following instruction of manufacturer. WSSV DNA copy was quantified by analysing the cycle threshold value (Ct) using Step One software v2.1 (Applied Biosystem). ROX dye was used to monitored background fluorescence. Standard quantification was based on a series of WSSV-

recombinant plasmid dilution with known copy number.

# RESULTS AND DISCUSSION

WSSV was detected in *Dendronereis* spp. with 1 step and nested PCR (**Fig. 1**).

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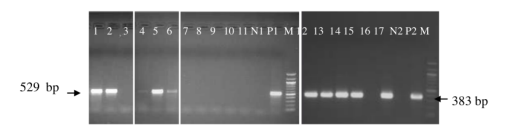


Fig. 1. 1-step and nested- PCR of *Dendronereis* spp. and *P. monodon* from shrimp pond. Lane 1-11 are result of 1-step PCR and lane 12-17are result of nested PCR. Lane 1-3: *P. monodon*, Lane 4-11: *Dendronereis* spp., Lane 12-16 *Dendronereis* spp.., lane 17 *P. monodon*. P1: Positive control for the 1-step PCR, P2: Positive control for nested- PCR, N1= Negative Control for 1-step PCR, P2= Positive control for nested- PCR. M= Marker (100 bp DNA ladder).

WSSV was detected in 7 out of 8 *Dendronereis* spp. tested with various degree of infection. Three out of 8 worm tested were positive with 1-step PCR (lane 4, 5 and 6) and 4 others were positive with nested PCR. 2 out of 3 of the shrimp tested were positive with 1 step PCR. Result of Real time PCR confirmed the findings of conventional PCR, indicating that WSSV load

in the *Dendronereis* spp. that shows positive response with 1-step PCR is quite high and comparable with that in the naturally infected shrimp taken from the same pond. In contrast, those that only gave positive signal with nested PCR, the WSSV DNA is undetectable with RT PCR (**Table 1**).

 Table 1. PCR analysis of WSSV infection in Dendronereisspp. detected with 1-step, nested and Real Time-PCR

Specimen	One step PCR	Nested -PCR	RT-PCR (WSSV copy / µg total DNA ND	
Dendronereis spp.	N	P		
Dendronereis spp.	N	P	NT	
Dendronereis spp.	P	NA	$1.9 \times 10^4$	
Dendronereis spp.	P	NA	$1.2 \times 10^2$	
Dendronereis spp.	P	NA	$1.5 \times 10^2$	
Dendronereis spp.	N	P	ND	
Dendronereis spp.	N	P	NT	
Dendronereis spp.	N	N	NT	
P. monodon	P	NA	$3.0 \times 10^4$	
P. monodon	P	NA	$1.57 \times 10^5$	
P. monodon	N	P	NT	

ND= Not detected, NA= Not applicable because specimen was positive with 1-step PCR; NT= Not tested, N= Negative, P= Positive

### DISCUSSION

OIE (2009) recommended 1 step and nested PCR to detect WSSV infection in shrimp,

because this method is highly specific and accurate. When tested shrimp give positive signal with 1-step PCR, it considers having heavy infection. On the other hand, since nested PCR can detect DNA in a much lower copy

number than 1-step PCR, specimen that gave positive signal with nested PCR is considered to have light infection (OIE 2706) and considered at carrier state (de la Pen 7 et al., 2007). WSSV has been detected with 1-step and nested PCR in broodstock 15 f P. (Withyachumnarnkul et al., 2003, Shahadat Hossain et al., 2004, de la Fina et al., 2007), broodstock and post larvae of fleshy shrimp Fenneropenaeus chinensis (Jang et al., 2009), as well as cultured shrimp (Shahadat Hossain et al., 2004). Unlike conventional PCR, RT-PCR can detest the viral DNA concentrartion in the tissue. Viral load in infected shrimp varied according to intensity of infation. Moribound penaeid shrimp ranged  $2.0 ext{ } 10^4 ext{ to } 9.0 ext{ } \cdot 10^{10}$ WSSV copies g-1 of total DNA (Durand and Lightner 2002) and slightly higher in shrimp with acute infection (Durand et al., 2003).

The WSSV load in the Dendronereis spp. and shrimp observed in this study is quite high, because it was detected with 1-step PCR in some specimen. Moreover, result of RT-PCR this study within the range reported by Jang et (2009), but lower than reported on the experimentally infected shrimp (Durand and Lightner, 2002, Durand et al., 2003). The degree of infection is varied among the Dendronereis spp. examined and in some specimen it is comparable with that of the naturally infected shrimp from the same pond that look healthy with no clinical sign of WSSV. It can be implied that the WSSV load in Dendronereis spp. is comparable with that in P.monodon at the carrier state. It has been suggested that polychaetes is a mechanical vector of WSSV and the virus is accumulated in the digestive tract by artificial infection (Vijayan et al., 2005). The part of Dendronereis spp. that we used included the front gut. However, whether WSSV merely accumulated or replicate in this organ need further study with immunohistochemistry or probe specific for WSSV.

## Conclussion

In conclusion, WSSV DNA load in infected *Dendronereis* spp obtained from shrimp pond is varied, and in some specimens is comparable with that in the infected shrimp.

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### REFERENCES

- Chang, P., Lo, C., Wang, Y., Kou, G. 1996. Identification of white spot syndrome associated baculovirus (WSBV) target organs in the shrimp *Penaeus Monodon* by *in situ* hybridization. *Dis. Aquat. Org.* 27: 131-139.
- Chang, Y., Chen, T., Liu, W., Hwang, J., Lo, C. 2011. Assessment of the roles of copepod *Apocyclops royi* and bivalve mollusk *Meretrix lusoria* in white spot syndrome virus transmission. Marine Biotechnology 1-9 (article in press).
- Chen, L. Lo, C., Chiu, Y., Chang, C. Kou, G. 2000. Natural and experimental infection of White Spot Syndrome virus (WSSV) in benthic larvae of mud crab *Scylla serrata*. Diseases of Aquatic Organisms 40: 157-161
- de la Peña, L.D., Lavilla-Pitogo, C.R., Villar, C.B.R., Paner, M.G., Sombito, C.D., Capulos, G.C. 2007. Prevalence of white spot syndrome virus (WSSV) in wild shrimp Penaeus monodon in the Philippines. *Dis. Aquat. Org.* 77: 175-179.
- Desrina, Haditomo, A.H.C., Prayitno, S.B. 2011.

  Survei keberadaan virus White spot syndrome (WSS) pada cacing polychaeta di tambak udang: studi kasus di Kendal. *Jurnal LITBANG Provinsi Jawa Tengah* 9(2): 124-129.
- Durand, S.V and Lightner, D.V. 2002. Quantitative real time PCR for the measurement of white spot syndrome virus in shrimp. *J. Fish. Dis.* 25: 381– 389.
- Durand V., Redman, R.M., Mohney, L.L., Tang-Nelson, K., Bonami, J.R.

- Lightner, D.V. 2003. Qualitative and quantitative studies on the relative virus load of tails and heads of shrimp acutely infected with WSSV. *Aqua*. 216: 9–18
- Escobedo-Bonilla, C.M., Alday-Sanz, V., Wille, M., Sorgeloos, P., Pensaert, M.B. and Nauwynck, H.J. 2008. A review on the morphology, molecular characterization, morphogenesis and pathogenesis of white spot syndrome virus. *J. Fish Dis.* 31: 1–18
- Esparza-Leal, H.M., Escobedo-Bonilla, C.M., Casillas-Hernández, R., Álvarez-Ruíz, P., Portillo-Clark, G., Valerio-García, R.C., Hernández-López, J., Méndez-Lozano, J., Vibanco-Pérez, N., Magallón-Barajas, F.J. 2009. Detection of white spot syndrome virus in filtered shrimp-farm water fractions and experimental evaluation of its infectivity in *Penaeus (Litopenaeus) vannamei. Aquaculture* 292: 16-22.
- FAO. The State of World Fisheries and Aquaculture 2010. Rome, FAO. 2010. 197 p.
- Flegel, T.W. 2006. Detection of major penaeid shrimp viruses in Asia, a major historical perspective with emphasis on Thailand. *Aquaculture* 258: 1-33.
- Hameed, A.S.S., Parameswaran, V., Mustaq, S.S., Sudhakaran, R., Balasubramanian, G. and Yoganandhan, K. 2005. A simple PCR procedure to detect white spot syndrome virus (WSSV) of shrimp, Penaus monodon (Fabricious). Aqua. Int.13: 441-450
- Jang, I., Meng, X., Seo, H., Cho, Y., Kim, B., Ayyaru, G and Kim, J. 2009. A TaqMan real-time PCR assay for quantifying white spot syndrome virus (WSSV) infections in wild brood stock and hazo ery-reared post larvae of fleshy shrimp Fenneropenaeus chinensis. Aquaculture 287: 40-45.

Kanchanapum, ., Wongteerasupaya, C., Sitidilokratana, N., Boonsaeng, V., Panyim, S., Tassanajakon, A., Withyachumnarnkul, B., Flegel, T.W. 1998. Experimental transmission of White Spot Syndrome Virus (WSSV) from crabs to shrimp *Penaeus monodon. Dis. Aquat. Org.* 34: 1-7.

ISSN: 1410-5217 Acrredited: 83/Dikti/Kep/2009

- Lem, A. 2006. An overview of global shrimp markets and trade. *In*: P. Leung and C. Engle (Editors). Shrimp culture: Economics, Market and Trade. Blackwell Publishing, Iowa.
- Ligtner, D.V. 1996. A handbook of shrimp pathology and diagnostic procedures for diseases of cultured penaeid shrimp. The World Aquaculture Society, Baton Rouge, LA.
- Liu, B., Yu, Z., Song, X., Guan, Y. 2007. Studies on the transmission of WSSV (white spot syndrome virus) in juvenile Marsupenaeus japonicus via marine microalgae. *J. Inv. Path.*
- Liu, W., Qian, D., and Yan, X.J. 2011. Studies on pathogenicity and prevalence of white spot syndrome virus in mud crab, *Scylla serrata* (Forskal), in Zhejiang Province, China. *J.Fish.Dis*. 34: 131-138.
- Marks, 27, Mennen, M., Vlak, J.M & Van Hulten, M.C.W. 2003.

  Transcriptional analysis of the White spot syndrome virus major virion protein genes. *J. Gen Vir.* 84: 1517-1523
- Meng, X., Jang, I., Seo, H., Cho, Y. 2009. White spot syndrome virus quantification in blue crab *Portunus trituberculatus* hatchery-produced larvae and wild populations by TaqMan real-time PCR, with an

- emphasis on the relationship between viral infection and crab health. *Aquacultur* 291: 18-22
- OIE. 2009. Manual of diagnostic test for aquatic animal. Downloaded from http://
  www.oie.int/fileadmin/Home/eng/health\_standards/aahm/2010/2.2.05
  WSD.pdf, 20 April 2010.
- Rahman, M.M., Corteel,M., Escobedo-Bonilla, C.M., Wille, M., Alday-Sanz, V., Pensaert, M.B., Sorgeloos, P., Nauwynck, H.J. 2008. Virulence of white spot syndrome virus (WSSV) isolates may be correlated with degree of replication in gills of *Penaeus vannamei* juveniles. *Dis. Aquat. Org.* 79:191-198.
- Schmitt, B. & Henderson, L. 2005.
  Diagnostic tools for animal diseases.
  Review of Science and technical of
  Office of International de Epizootics
  24 (1): 243-250.
- Shahadat Hossain, Md., Otta, S.K., Chakraborty, A., Sanath Kumar, H., Karunasgar, I., Karunasgar, I. 2004. Detection of WSSV in cultured shrimps, captured brooders, shrimp postlarvae and water samples in Bangladesh by PCR using different primers. *Aquaculture* 237: 59-71.
- Sunarto, Α., Widodo, Taukhid, Koesharyani, I., Supriyadi, H., Gardenia, L., and Sugianti, B. 2004. Current status of transboundary fish diseases in Indonesia: Occurrence, surveillance, research and training:Country report. In Lavilla-Pitogo, C.R., Nagasawa, Proceeding of The meeting on Transboundary Fish Diseases in South East Asia.: Occurrence, Research Surveillance, Training. Manila, The Philipines 23-

24 June 2004. Aquaculture Department Southeast Asian Fisheries Development Center Iloilo, Philippines (downloaded from

ISSN: 1410-5217 Acrredited: 83/Dikti/Kep/2009

- http://rfdp.seafdec.org.ph/publication/manual/trans/frame.htm on 12 March 2011.
- Supamattaya, K., Hoffman, R.W., Bonyaratpalin, S., Kanchanaphum, P. 1998. Experimental transmission of white spot syndrome virus (WSSV) from black tiger shrimp Penaeus monodon to the sand crab Portnus pelagicus mud crab Scylla serrata and krill Acetes sp. Dis. Aquat. Org. 32: 79-85.
- Vijayan, K., Raj, V.S., Balasubramanian, C.P., Alavandi, S.V., Sekhar, V.T., Santiago, T.C. 2005. Polychaete worms- a vector for white spot syndrome virus (wssv). *Dis. Aquat. Org.* 63: 107-111.
- Withyachumnarnkul, B., Boonsaeng, V., Chomsoong, R., Flegel, T.W. Muangsin, S., Nash, G.L. 2003. Seasonal variation in white spot syndrome virus-positive samples in broodstock and post-larvae of Penaeus monodon in Thailand. *Dis. Aquat. Org.* 53: 167-171.
- Yan, D., Dong, S., Huang, J., Yu, X, Feng, M., and Liu, X. 2004. White Spot syndrome virus (WSSV) detected by PCR in rotifers and rotifer resting eggs from shrimp pond sediments. Dis. Aquat. Org. 59: 69-73.
- Zhang, J., Dong, S., Dong, Y., Tian, X., Hou, C. 2008. Bioassay evidence for the transmission of WSSV by the harpacticoid copepod *Nitocra* sp. *J. Inv.Path.* 97: 33-39.

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- RW Chapman, CL Browdy, S Savin, S Prior, E Wenner. "Sampling and evaluation of white spot syndrome virus in commercially important Atlantic penaeid shrimp stocks", Diseases of Aquatic Organisms, 2004
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