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**Vibrio harveyi** and the ‘green water culture’ of *Penaeus monodon*

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**ABSTRACT**

The “greenwater culture” of the tiger shrimp, *Penaeus monodon*, is an innovative culture technique for the grow-out rearing of shrimps. This culture method involves the use of rearing water of tilapia for the rearing of tiger shrimp in grow-out ponds and or the polyculture of shrimp with tilapia. This culture technique was reported to prevent disease outbreaks attributed to luminescent *Vibrio*. To understand the possible mechanisms of luminous *Vibrio* control in the green water culture system several studies were conducted. This review summarizes the highlights obtained so far from these studies consisting of a) effect of rearing waters from tilapia culture and shrimp cultured with tilapia on *Vibrio harveyi*; b) estimation and preliminary identification of cultivable bacteria, fungi and phytoplankton flora associated with the “green water culture” system and c) detection of anti-*Vibrio harveyi* metabolites from bacteria, yeast, filamentous fungi and phytoplankton indigenous to the “green water culture” system.

**INTRODUCTION**

The giant tiger shrimp, *Penaeus monodon*, comprise 56% of total shrimp production in the world (Fig. 1). As a significant dollar earner for exported shrimp in the Philippines, its production significantly increased through the years until 1995 (Fig.2). By 1996, shrimp production in the country drastically declined. These severe losses in grow-out cultured shrimps were attributed to outbreaks of luminous vibriosis caused by *Vibrio harveyi* (Lavilla-Pitogo et al., 1998a; Leaño et al., 1998, Lio-Po, 1998; de la Peña et al., 2001). Similarly, luminous vibriosis in *P. monodon* juveniles was also reported in Thailand (Jiravanichpaisal, et al., 1994).
Figure 1. World production of shrimps by species (FAO, 2000)

Figure 2. Shrimp production in the Philippines, 1982-2000 (BAS, 1983–2001)
To prevent the multiplication of luminous *Vibrio* in the shrimp grow-out ponds and to improve survival and production, innovations in shrimp culture techniques were developed (NPPMCI, 2000). Among these is the “green water culture technique” also known as “Tilapia integration to shrimp culture technique” (TIPS) for the grow-out culture of the tiger shrimp, *P. monodon*, initiated in 1999. The application of this rearing technique was reported to prevent the increase of *V. harveyi* in the ponds thereby preventing the onset of *V. harveyi* infections in the cultured shrimps (Fig. 3) (Platon, 1997; Corre *et al.* 2000). Briefly, this method utilizes water from tilapia culture ponds as the major source of rearing water for shrimp culture. In addition, tilapia may be stocked in cages in polyculture with the shrimp (Fig. 4).

**Figure 3.** Luminous bacterial counts vs. salinity in a grow-out pond with *Penaeus monodon* applying the “greenwater culture” system (NPPMCI, 2000)

**Figure 4.** The ‘green water culture” system in the grow-out culture of *Peneaus monodon* with Tilapia at the central cage
Because of the successful production runs attributed to the ‘green water culture’ system for the grow-out culture of *P. monodon*, this culture technique has gradually gained acceptance among shrimp farmers. However, the basis for its success needs to be investigated scientifically. This report summarizes the results of studies on the microbial flora associated with the “green water culture system” conducted at SEAFDEC-AQD namely: a) effect of the green water culture rearing waters on *V. harveyi*; b) estimation and preliminary identification of the cultivable bacterial, fungal and phytoplankton flora associated with the green water culture system; and c) detection of anti-*V. harveyi* metabolites from these microbial flora.

**VIBRIO HARVEYI AND SHRIMP/TILAPIA CULTURE WATERS**

Pure cultures of *V. harveyi* were exposed to rearing waters of shrimp grow-out culture, shrimp with tilapia culture, tilapia culture and their respective seawater sources in triplicates. Results did not show a consistent correlation of *V. harveyi* inhibition when treated with rearing waters stocked with tilapia or shrimp with tilapia. It is likely that the varying conditions in the sampled ponds at different sites, influenced the results. Therefore, the experiment needs repetition under more controlled culture conditions.

**GREEN WATER CULTURE: BACTERIAL, FUNGAL AND ALGAL POPULATION/FLORA**

The microbial and phytoplankton flora associated with the green water culture system of *P. monodon* were estimated and isolated. Bacterial counts using the spread plate method were done on nutrient agar (NA) supplemented with NaCl, thiosulfate citrate bile salts (TCBS) agar and Pseudomonas Aeromonas Selective Agar (GSP). Identification was based on Gram staining characteristics and biochemical tests following Baumann *et al.* (1984), Krieg (1984) and Popoff (1984). Fungal enumeration, isolation and identification followed the methods of Barnett and Hunter (1987), Kohlmeyer and Volkman-Kohlmeyer (1991) and Leaño (2002). Phytoplankton estimates were made using standard methods on a Burker Turk hemacytometer and a Sedgwick-Rafter counting chamber with characterization. Identification of the algae followed the key of Yamaji (1991).

The total bacterial counts (TBC) of cultivable bacteria in rearing water from tilapia pond, shrimp with tilapia and seawater sources in Negros Occidental did not vary significantly. The estimated TBCs were $10^2-10^3$, $10^2-10^4$ and $10^2-10^5$ cfu/ml at days of culture (DOC) 15, 30, 45 and 60, respectively. The highest bacterial count of $10^4$ cfu/ml was only observed in the shrimp with tilapia culture waters. The hepatopancreas of shrimps cultured with tilapia had TBCs of $10^3$ to $10^5$ cfu/g. In the tilapia mucus, an increasing trend in TBCs of $10^3$, $10^4$, $10^5$ and $10^6$ cfu/5 cm$^2$ at DOC 15, 30, 45 and 60, respectively, were observed. The tilapia gut, on the otherhand, yielded TBCs of $10^5-10^6$ cfu/5 cm.

Likewise, the presumptive *Vibrio* counts (PVC) of water sources and the green water rearing waters did not vary significantly at $10-10^3$ cfu/ml. The PVCs of the hepatopancreas of shrimps cultured with tilapia were $10-10^3$ cfu/g. The mucus of tilapia contained PVCs of $10^2-10^4$ cfu/5 cm$^2$ while the fish gut harbored PVCs of $10^4-10^6$ cfu/5 cm.
The luminous bacteria counts (LBC) were detectable at a range of 10-10^2 cfu/ml in the water sources but not in the rearing waters. All sampled hepatopancreas of shrimp cultured with tilapia had no detectable luminous bacteria. There were also no luminous bacteria in the tilapia mucus but the tilapia gut had LBCs of 10-10^5 cfu/5 cm.

By and large, the bacterial flora associated with the “green water culture” rearing waters consisted of non-luminous Vibrio spp., Pseudomonas spp., Aeromonas spp, Flavobacteria spp., members of the Enterobacteriaceae family and some unidentified bacteria at varying percentages (Lio-Po et al., 2001).

On the other hand, total fungal flora showed the highest load on the fish gut at 10^2-10^3 cfu/5 cm and mucus samples at 48-10^2 cfu/5 cm^2. Lower fungal load, however, were observed in shrimp hepatopancreas and water samples yielding 2-37 cfu/g and 5-44 cfu/ml, respectively. The yeast population was dominated by Rhodotorula sp., Saccharomyces sp. and Candida sp. Among the genera of filamentous fungi commonly observed were Aspergillus sp., Penicillium sp., Fusarium sp., and Curvularia sp.

The phytoplankton flora of the green water culture system predominantly consisted of Chlorella at 10^5-10^6 cells/ml followed by an alga tentatively identified as Leptolyngbia sp. at 10^4-10^5 cells/ml. Other species like, Thalassiosira sp., Skeletonema sp., Nitzchia sp., Navicula sp., Chaetoceros sp. and Anabaena sp. were also detected at lesser concentrations.

**GREEN WATER CULTURE: ANTI-VIBRIO HARVEYI METABOLITES**

Bacterial, fungal and algal isolates obtained from the “green water culture” system were screened for the presence of extracellular and intracellular metabolites affecting growth and viability of luminous Vibrio. Pure cultures of each isolate were used. In-vitro co-cultivation of selected bacterial and algal isolates with V. harveyi were conducted and the survival of V. harveyi was quantified after 24 and 48 h exposure. Fungal assays for extracellular metabolites were conducted using the modified disc-agar-diffusion method.

Results showed that of the 111 bacterial isolates tested, 31, 18 and 18 isolates exerted marked, moderate and slight inhibition of V. harveyi, respectively, after 24 h culture. Among fungal isolates, no zones of inhibition were observed with the extracellular metabolites of 20 yeast isolates. However, the intracellular extracts of four yeast isolates caused growth inhibition of V. harveyi. Among the 41 filamentous fungal isolates, the extracellular metabolites of three isolates caused slight inhibition of the test bacterial pathogen while the intracellular extracts of five isolates showed slight inhibition of the V. harveyi. With the algae, Skeletonema sp. induced a one-log reduction of V. harveyi after 48-72 h incubation with its extracellular metabolites.

**DISCUSSION**

In this study, the total bacterial counts observed in the hepatopancreas of shrimps in the “green water culture” system at DOC 30 was about two logs less than that observed in P. monodon with luminous vibriosis in the Philippines in an earlier study and in intensively cultured
shrimps in Thailand (Leaño et al., 1998; Ruangpan et al., 1994). The bacterial flora associated with the hepatopancreas of shrimps in the “green water culture” system at DOC 15, 30, 45 and 60 was predominated by non-luminous *Vibrio*. In contrast, Ruangpan et al. (1994) reported that *Pseudomonas* spp. predominated over *Vibrio* spp. in the hepatopancreas of intensively cultured *P. monodon* in Thailand. It was also reported that the hepatopancreas of pond-reared *P. monodon* juveniles associated with luminous *Vibrio* outbreaks may harbor *V. harveyi* of as much as $10^6$ cfu/g (Leaño et al., 1998). These findings indicate that the “green water culture” favorably affects the bacterial population of the rearing waters and of the cultured shrimps.

The present study has shown that some bacteria, fungi and phytoplankton associated with the “green water culture” system secrete growth inhibitory metabolites against *V. harveyi*. The undetectable levels of luminous bacteria in the hepatopancreas of shrimps cultured with tilapia and in the mucus of tilapia is a further positive indicator that the “green water culture” system does prevent the multiplication of the luminous *Vibrio*. This may be attributed to the strong antibacterial activity of protein extracts found in the fish mucus (Ebran et al., 1999). Moreover, Chong et al. (2002) reported that among tilapia, snakehead and carp mucus, tilapia mucus had the highest level of proteins. Earlier, Torrento and Torres (1996) and Ruangpan et al., (1998) reported the *in-vitro* inhibition of *V. harveyi* by *Pseudomonas* spp. and *Vibrio alginolyticus*, respectively.

The presence of filamentous fungi and yeasts especially in the gut and mucus of tilapia in the “green water culture” of *P. monodon*, may also contribute to the inhibition of the growth of *V. harveyi* through the production of intra- and extracellular metabolites, as shown in the metabolite assays in this study. Many marine fungi produce novel compounds and enzymes (Biabani and Laatsch, 1998). Furthermore, fungi are good sources of glucans, a proven immunostimulant for crustacean species (Sung et al. 1994; Devaraja et al., 1998, Chang et al., 2000). Thus, their role or importance in many marine habitats, including aquaculture ponds, need further investigation.

Microalgae as absorbers of carbon dioxide and providers of oxygen, improve water quality in the aquatic environment through oxygenation and filtration. These processes help prevent the occurrence of diseases caused by pathogenic bacteria and fungi (Round, 1973). Moreover, phytoplankton are potential sources of anti-microbial compounds. *Chlorella*, for instance, is used in the preparation of the antibiotic, Chlorellin, against gram positive and gram negative bacteria (Sharma, 1986). *Nitzschia perlea* produces a growth inhibiting antibiotic against *Escherichia coli* (Round, 1973). In a mixed culture experiment of $10^2$ cfu/ml *V. harveyi* with either $10^5$ cells/ml *Skeletonema costatum* or $10^6$ cells/ml *Chaetoceros calcitrans*, the population of the bacterial pathogen was reduced by at least one log but not in the corresponding algal-free filtrates (Lavilla-Pitogo et al., 1998b). Furthermore, Naviner et al. (1999) partly purified the bioactive compound found in *S. costatum*.

In summary, the effectiveness of the green water culture system for the grow-out culture of *P. monodon* in preventing luminous vibriosis maybe attributed to the presence of bacteria, fungi and algae that secrete *V. harveyi* growth inhibitory factors. In addition, the yeasts flora that are significantly abundant in tilapia, is a probable source of beta-glucan that is known to exert an immunostimulating effect on *P. monodon* against *Vibrio* infections. Also, the tilapia mucus that apparently prevents the colonization of *V. harveyi* on the fish skin, may also shed these anti-*V. harveyi* factors into the rearing waters thereby contributing to the biocontrol of this bacterial pathogen. It is likely that the low to undetectable levels of luminous *Vibrio* in the green water
culture system for *P. monodon* is a result of the combined inhibitory effects of factors in the fish mucus and microbial flora of this culture system. Therefore, the microbial flora associated with the "green water culture" system is an excellent source of indigenous microorganisms with probiotic potentials.

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