A Study on culture of *Litopenaeus vannamei* (preparation, water quality parameters) compared with different stocking densities using standard operational procedures

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Abstract

In India, the shrimp culture today has industrialized into an improved farming system and is evolved day by day into a well talented for the management. Present work was carried out in the shrimp *Litopenaeus vannamei* culture in Sattanathar aqua farm in the east coast of Nagai District, Tamil Nadu, India from April 2015 to August 2015. Water quality parameters for the entire culture ponds was properly monitored. Pond water pH, temperature and DO readings is measured in early mornings and in the late evenings. For the three culture ponds, the average pH value was found between 7.9 and 8.6 in the early morning, while fluctuation of pH value was between 8.2 and 8.6 in the evening. Dissolved oxygen values varied between 4 mg/L to 8 mg/L. In overall, early morning readings recorded lesser and the cycle proceeded and the standing crop was enlarged. Average pond temperatures were 26 to 31°C, Generally, the temperature tendency in the production cycle started around 28°C and got reduced to 26°C due to cold conditions were found during the 21st and 28th day, and it was found to be increased at 29–31°C. During the culture period, the highest salinity was recorded as 41ppt and reduced salinity was found as 25 ppt in all the culture ponds.

Keywords: *Litopenaeus vannamei*, shrimp culture, fluctuations, Salinity and water quality

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Introduction
In India, the shrimp culture today has industrialized into an improved farming system and is evolved day by day into a well talented for the management. The last half a decade has proved adequately that the possible for its shrimp production is quite good in India with its warm tropical climate, suitable soil, along the main estuaries and lagoons, suitable water availability and potential force of highly industrious farming community (Lightner, 1996; Flegel, 1997). Due to last outbreak of White Spot Syndrome (WSSV) in P. monodon culture indications to devastating of shrimp culture in India. In the meantime, the Coastal Aquaculture Authority of India (CAA) has presented a new species (L. vannamei) in India and the Coastal Aquaculture Authority of India is actual intense in the Biosecurity and endorsement for the cultures of L. vannamei.

There is very limited research work was done on the culture and growth performance of L. vannamei with different high stocking densities in Tamil Nadu. So, hence the present study was investigated to assess the water quality, survival, growth and FCR of L. vannamei cultured in different stocking densities.

Materials and methods
The current work was commenced at a shrimp farm culture in Tranquilbar, Nagai district, Tamil Nadu, India. The study was carried out in four shrimp rearing culture ponds. Three ponds (A1, A2, and A3) were 0.5ha, pond A4 and A5 was 1.0ha in area. Besides that there was a reservoir present in the A4 pond with a size of 0.5ha, a sedimentation pond and a chlorination pond are in the size of 0.5 ha. Water re-circulation method was trailed to prevent cross infection during the culture period. All the investigational cultural ponds were measured to about 1.2–1.5m depth. The type of the soil should be maintained as sandy clayey soil. Ponds were primarily organized by drying, tilting (to remove the pests and predators and oxidize bottom soil) and liming to adjust the pH of the soil (Figures 1-4).
Figure 3: Ploughing or tilting the bottom of pond

Figure 4: Lime applications in the bottom of pond

Figure 5: Bird fencing

Many inorganic fertilizers like urea and triple superphosphate were applied to enrich the natural food organisms present in the water. Bird netting and crab fencing were performed before pumping water to prevent the auto entrants (Figs. 5 and 6). The filter bags (Fig. 9) were periodically checked, and it was fitted in the inlet and outlet pipe followed by the pumping the water to the whole ponds. After satisfying, the water was allowed to stand for 1 day for sedimentation. Then the water was chlorinated (60 ppm/ha) and excess of chlorine was neutralized by de-chlorination process takes nearly 72 hours. After that, the water was supplemented with probiotic to provide the good beneficial bacterial environment. After 7 days, the algal bloom was noticed slowly on the external of the water in the ponds. The Post larval *L. vannamei* seeds (stage 14) were procured, it was acclimatized, maintained to a salinity of 17 ppt and it’s an evidence for the negative symptoms of the WSSV and Taura Syndrome Virus (TSV) confirmed by using molecular tools like (PCR – Polymerase Chain Reaction) were obtained from Sigma, India and the seeds were procured from Rank marine hatchery, Marakaanam, Pondicherry. The seeds were transported carefully in double-layered oxygenated polyethylene bags with wrinkled ice packs placed in between the covers of the bag to maintain an optimal temperature for the less stress shrimps and the whole set up was packed in a container.

The seeds were immediately transported to the farm site and the seed bags were kept in the pond water for one hour to acclimatize the seeds. Then, the pond water was further added gradually into the seed bag to adjust the physical factors like salinity and pH. Next, the shrimp seeds were released gradually and deliberately in to the ponds. The density
of the stocking must be sustained with 80/m², 120/m² and 160/m² for ponds A1, A2 and A3 respectively. RNK feed pellets (Plates 9 and 10) were fed to the stocked post larvae for four times daily at 7am, 10am, 1pm and 4pm respectively.

In the meantime, water should be added from the water reservoir at steady intermissions to maintain the water loss due to evaporation or soil seepage. During harvesting time, the water from entire culture ponds were drained completely by allowing them to form sediment in the pond and eventually reached to reservoir pond. At any occasion, the culture pond water should not be driven out side from the farm due to bio secure reasons. From the 40th day of culture the DOC, forwards cast net sampling method was employed at each week for monitoring the status of growth and health of the shrimps (Fig. 10a-f).
The water level was measured periodically by using a usual scale with centimeter marking. The physiological water quality parameters like salinity by refractometer, pH was measured by using readymade pH pen, temperature by thermometer, dissolved oxygen by DO meter and light transparency were measured respectively. Long arm aeration equipment was fitted and supplied to the entire ponds (Fig. 11). Totally 20 horse power (hp) aerator was fitted for the individual culture pond. The aerators are kept can perform dissolve maximum dissolved oxygen level (DO) into the pond water and makes the culture pond eco-friendly.
Feed conversion ratio (FCR) and Average daily growth (ADG) were calculated as follow:

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\text{FCR} = \frac{\text{Total weight of the harvested shrimps}}{\text{total feed used}}
\]

\[
\text{ADG} = \frac{\text{Total weight gained by the shrimps}}{\text{Total days of culture}}
\]

**Collection of water samples**

Periodically, the water samples were collected from the culture ponds. The samples for dissolved oxygen were collected at the depth of water in order to prevent the direct contact with air. The determination of turbidity, temperature, pH, salinity, calcium, nitrate and ammonia was carried using the similar sample was taken to reduce the duration between the sample collection and analysis of sample. The samples were investigated by using standard methods.

**Analysis of water quality parameters**

The below mentioned parameters are analyzed at each ten days once for all the entire culture ponds. The level of water was measured by standard scale (cm) with marking. Normally temperature measurement is made with good, Celsius thermometers as a minimum the thermometer must have a scale marking at each 1°C with marking edged on the capillary glass, for field operators a thermometer having a metal case to movement breakage (APHA, 1998). Turbidity of water is measured by a standard Sachi disc meter scale, with pin (APHA, 1998). The water salinity was measured by using a hand refractometer. Hydrogen ion concentrations of the culture pond water were estimated with the help of pH meter manufactured by Hanna Instrumental Company, Japan. The water samples were collected and transferred to the beaker. There are many methods for determining the pH of the water samples. These may be broadly divided into electrometric method and calorimetric method. Here the hydrogen ion concentration was determined with wide range pH papers in the field and later the values were collected with pH meter in the laboratory. In the electrometric method the glass electrodes were cleaned and allow drying with filter paper and immersed into the sample. And the pH was recorded calibration of the scale was necessary and so buffer solution was prepared. The instrument was standardized with the help of standards. A buffer tablet (commercial) dissolved in100 ml of distilled water will form a buffer solution of pH. The presence of calcium in the pond water was estimated by using EDTA Titration method. Brucine method was used widely to calculate the nitrate level in the pond water followed by ammonia using Nessler method.
Microbiological analysis
The sediment and pond water samples were collected aseptically stored in sterile polythene bags from various location of the ponds and were mixed to make a one. The similar procedure was repetitive for every culture pond and the final samples were transported to the laboratory immediately and were analyzed for their microbial counts. The samples were transferred to a sterile 150 ml conical flask containing 99mL of sterile dist. water and 1gm of sample thoroughly mixed followed by serial dilution for estimating the total microbial load in different dilutions like $10^{-1}$ to $10^{-5}$ suspension samples. The Zobell marine agar medium was used to enumerate the Total Heterotrophic Bacteria (THB), (Table 5), for Vibrio spp TCBS media was purchased from Hi-media, Mumbai.

Isolation and enumeration
Spread plate methods were employed for enumerating the microbial load. The sterilized medium was decanted in Petri dishes under aseptic conditions, to allow the media to solidify. 0.1µl of the sample was pipetted out and the sample was spread by using sterile L-rod by rotating in clock and anticlockwise direction for 2-3 times. Ensure the sample should be spread the entire area for proper microbial growth. Then the plates were placed in inverted position and incubated at 28±2°C. All the experiments were carried out in triplicates. After 2-3 days, the colonies were counted in all the plates using colony counter and the Colony Forming Units (CFU’s) in one gram of sample was calculated by using formula,

Total microbial load
In the given sample (CFU/g) =Total number of colonies/ Samples of volume plated (0.1) X Dilution

Statistical analysis
Two way analyses - ANOVA was employed to distinguish the statistical significance between growth and stocking densities of the shrimp. Data was expressed in mean ± standard error (SPSS- package).

Present work was carried out in the shrimp L. vannamei culture in Sattanathar aqua farm in the east coast of Nagai District, Tamil Nadu, India from April 2015 to August 2015. Water quality parameters for the entire culture ponds are summarized in Table 1. Pond water pH, temperature and DO readings is measured in early morning and in the late evening. For the three culture ponds, the average pH value was found between 7.9 and 8.6 in the early morning, while fluctuation of pH value was between 8.2 and 8.6 in the evening. DO values varied between 4 mg/L to 8 mg/L. In overall, early morning readings recorded lesser and the cycle proceeded and the standing crop was enlarged.
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**Table 1: Mean water quality parameters of the culture ponds**

| Parameters         | A1            | A2            | A3            |
|--------------------|---------------|---------------|---------------|
| Salinity (ppt)     | 25 – 40       | 25 – 38       | 25 – 41       |
| Temperature °C     | 26 – 31       | 26 – 31       | 26 – 31       |
| pH - AM            | 7.9 - 8.5     | 7.9 - 8.6     | 7.9 - 8.6     |
| pH - PM            | 8.2 - 8.5     | 8.2 - 8.6     | 8.2 - 8.6     |
| Transparency       | 60 – 30       | 65 – 30       | 65 – 25       |
| Dissolved oxygen   | 5 – 8         | 4 – 8         | 4 – 8         |
| Ammonia            | .1 – .3       | .1 – .4       | .1 – .5       |
| Calcium            | 500 – 650     | 400 – 650     | 400 – 700     |

**Table 2: Average population of yellow colony in the A1, A2 & A3 ponds**

| PONDS | Days of culture (DOC) |
|-------|-----------------------|
|       | 10 | 20| 30 | 40 | 50 | 60 | 70 | 80 | 90 | 100 | 110 | 120 |
| A1    | 90 | 100 | 110 | 130 | 160 | 200 | 320 | 270 | 250 | 220 | 300 | 210 |
| A2    | 90 | 90 | 100 | 100 | 90 | 210 | 130 | 110 | 100 | 190 | 180 | 160 |
| A3    | 120 | 150 | 214 | 200 | 320 | 312 | 250 | 520 | 352 | 450 | 380 | 415 |

**Table 3: Average population of the green colony in various ponds**

| PONDS | Days of culture (DOC) |
|-------|-----------------------|
|       | 10 | 20 | 30 | 40 | 50 | 60 | 70 | 80 | 90 | 100 | 110 | 120 |
| A1    | 60 | 10 | 30 | 100 | 80 | 40 | 32 | 100 | 90 | 120 | 100 | 80 |
| A2    | 40 | 50 | 40 | 40 | 60 | 50 | 60 | 40 | 50 | 40 | 50 | 40 |
| A3    | 50 | 60 | 90 | 90 | 130 | 160 | 200 | 180 | 200 | 200 | 180 | 150 |

Average pond temperatures were 26 to 31°C, respectively (Table 1). Generally, the temperature tendency in the production cycle started around 28°C and got reduced to 26°C due to cold conditions was found during the 21st and 28th day, and it was found to be increased at 29–31°C. During the culture period, the highest salinity was recorded as 41ppt and reduced salinity was found as 25 ppt in all the culture ponds. The population of bacterial communities were changed during every sampling. The bacterial population was found to be yellow colony (beneficial bacteria count) in pond A1 maximum 320 and minimum 90 was recorded, in pond A2, A3 the maximum yellow colony were recorded 190, 520 and minimum 120, 90 were recorded. The bacterial population green colony count (harmful bacteria) was recorded in maximum 120, 60 and 200 at pond A1, A2 and A3, respectively. The minimum 10 was recorded in pond A1. Weekly growth of the shrimp is presented in Table 4.
After 130th days of culture, the average growth were recorded as 28.5g, 26.8g and 25.3g for ponds A1, A2 and A3 respectively (Table 4). Survivals were 80, 72 and 66% for ponds A1, A2 and A3 respectively; FCR was 1.4, 1.5 and 1.7 for ponds A1, A2 and A3, respectively. The average production was 18250, 23478 and 26640 kg/ha for ponds A1, A2 and A3, respectively (Table 5).

### Table 4: Average body weight of the A1, A2 & A3 culture ponds

| DOC | A1  | A2  | A3  |
|-----|-----|-----|-----|
| 40  | 6.3 | 6.1 | 5.4 |
| 50  | 8.5 | 8.4 | 7.0 |
| 60  | 10.8| 10  | 9.2 |
| 70  | 13.1| 12.8| 11.4|
| 80  | 15.8| 15.2| 13.6|
| 90  | 18  | 17.1| 15.8|
| 100 | 20.6| 19.5| 18.0|
| 110 | 23.1| 22  | 20.4|
| 120 | 25.8| 24.5| 21.9|
| 130 | 28.5| 25.5| 23.7|
| 135 | 0   | 26.8| 0   |
| 140 | 0   | 0   | 25.3|

### Table 5: Composition of Zobell Marine Agar

| Composition   | Amount (g) |
|---------------|------------|
| Peptone       | 5.0        |
| Yeast extract | 1.0        |
| K2 HPO4       | 0.5        |
| Feso4         | Trace      |
| Agar          | 15         |
| 50% seawater  | 1000mL     |
| pH            | 7.2        |
Discussion

The present investigation is on the culture of *L. vannamei* in the estuarine or brackish water shrimp farms in Tranquebar, Nagai district, Tamil Nadu, India. This work shows that high stocking density with proper water quality and feed management can give good growth of *P. vannamei*. Many researchers reported that the growth and survival of *L. Vannamei* is depends on various salinities and densities (Wyban *et al*., 1988; Samocha *et al*., 1993, 1999; Emberson *et al*., 1999, Gunalan *et al*., 2011). The water quality should be upheld properly and it’s very much important for their optimum survival and growth. The quality of water is very much considered for many factors like DO, temperature, salinity and pH. The surplus of feed, their waste matter and other metabolites will employ incredible effect on the water in shrimp culture pond (Soundarapandian and Gunalan, 2008). The salinity was maintained with an average of 25 – 40ppt in all the culture ponds. *L. vannamei*, is extensively cultured in Central and South America (Wen-Young 1988) and it tolerates the salinity of 2-45 ppt (Parker *et al*., 1974; Samocha *et al*., 1998). Numerous authors have stated survival and good growth of *L. vannamei* in brackish water ranges from 1.7-2.3 ppt (Bray *et al*., 1994; Samochaet *et al*., 1999; Emberson *et al*., 1999; Moya *et al*., 1999). Karthikeyan (1994) and Gunalan *et al*., (2010) recommended a salinity range of 10 –35 ppt is perfect for shrimp culture. Samocha *et al*., (2004) described that development of shrimps is high in lower salinity at 2 ppt than in sea water.

The pH value is ranged from 7.9–8.2 in the early sunshine morning and 8.2-8.6 in the sunset. The pond water pH is prejudiced by many factors, which includes pH, water source, and acidity of bottom soil, inputs of shrimp culture and various biological activities. Wang *et al*., (2004) suggested the optimum pH ranges from 7.5 -8.7 in *L. vannamei*. The levels of dissolved oxygen in the entire ponds were ranged from 5.0-8.0 mg/L during the whole culture period. The water quality parameters values revealed that all are in the appreciable range for survival and growth of *L. vannamei* (Van Wyk *et al*., 1999). In the present work, RNK sunny feed pellets were used for the entire ponds and the same quantity was followed as per standard feed chat. The extreme feeds were used in pond A3 followed by A2 and A3. From this present investigation, the average FCR was 1.4 to 1.7 for the whole ponds. Related reports were previously recorded by Paul Raj (1999), Ramakrishna (2000) and Soundararapandian and Gunalan (2008).

In the present work, the bacterial population yellow colony (beneficial bacteria count) in pond A1 maximum 320 and minimum 90 were found in pond A2, A3 the maximum yellow colony were recorded 190,520 and minimum 120, 90 were recorded. The bacterial population green colony counts (harmful bacteria) were found maximum in 120, 60,200 at pond A1, A2 and A3, respectively. The minimum 10 was found in pond A1.

Ruangpan and Kitao (1991) described...
that the high occurrence of luminescent *Vibrio* is reliable with incidence of disease and poor or zero harvest results. The bacterial pathogen like *Vibrio harveyi* will affect the *P. monodon* and it leads to drastic loss (Baticados *et al.*, 1990). The probiotic bacteria application was used either in higher or a very low profusion or a comprehensive absence of luminous *Vibrio* in pond water results in good harvest. The steady and higher productivity will occur, even in the percentage of luminescent *Vibrio* in the water was found to be higher in the sea water source, and the richness of total green colony in the pond water was higher than in source of water. Moreover, luminescent *Vibrio* was totally absent in all the stages of growth and the presence of the super biotic *Bacillus* species was recorded.

In the present study, first sampling was carried out in all ponds at the 40th DOC of the culture. During harvest in A1 pond the shrimps are ranged at a size of 28.5 g, in A2 pond 26.8 g and in A3 pond 25.3 g. Higher survival (80%) was noted in pond A1 and lower survival (66%) was noted in pond A3. Bray *et al.* (1994) was observed similar finding in their research. The survival of shrimp was quite well were considering the area and the size of pond and the hygienic risks of outdoor-reared shrimp (Green *et al.*, 1997; Martinez- Cordova *et al.*, 1998). The considerable FCR revealed better rearing practices shared with an appropriate environment and a good shrimp biological responsiveness. In the present work, higher growth was reported in pond A1 followed by pond A2. This accepts the basic concept of aquaculture that better water quality management, proper aeration and good seed feed management leads to better survival and growth. From the present study it was determined that *L. vannamei* culture is fruitful in brackish water environment and the growth is directly proportional to proper water quality, aeration and feed management.

Reference

APHA Method 4500-F, 1998. Standard methods for the examination of water and waste water, 19th edn. American Public Health Association and Water pollution Control Federation, Washington DC, USA. pp. 59–64.

Atwood, H.L., Young, S.P., Tomasso, J.R. and Browdy, C.L., 2003. Survival and growth of pacific white shrimp *Litopenaeus vannamei* post larvae in low-salinity and mixed-salt environments. *Journal of the World Aquaculture Society*, 34, 518–523.

Baticados, M.C.L., Lavilla-Pitogo, C.R., Cruz-Lacierda, E.R., de la Pena, L.D. and Sunaz, N.A., 1990. Studies on the chemical control of luminous bacteria *Vibrio harveyi* and *V. splendidus* isolated from diseased *Penaeus monodon* larvae and rearing water. *Diseases of Aquatic Organisms*, 9, 133–139.

Bray, W.A., A.L. Lawrence and J.R. Leung-Trujillo 1994. The effect of salinity on growth and survival of *Litopenaeus vannamei*, with observation on the interaction of
IHHN virus and salinity. *Aquacult.*, 122: 133–146.

Emerson, C.R., Samocha, T.M. and Wood, G.F., 1999. Use of ground saline water for commercial production of *Litopenaeus vannamei* in the Sonora desert, Arizona, USA. In: Book of Abstracts. *World Aquacult.*, Soc. Ann. Conf., Sydney, Australia. 668P.

Flegel, T.W. 1997. Special topic review: major viral diseases of the black tiger prawn (*Penaeus monodon*) in Thailand. *World J. Microbiol. Biotechnol.*, 13: 433–442.

Green, B.W., D.T. Teichert-coddington, C.E. Boyd, J. Wigglesworth, H. Corrales, R. Zelaya, D. Martinez and E. Ramirez 1997. Effect of diet protein on semi-intensive production of *Penaeus vannamei* during the rainy season. In: Fifteenth Annual Technical Report. pp. 77–83.

Gunalan, B., P. Soundarapandian and G.K. Dinakaran 2010. Effect of Different Stocking Densities on the MBV Infected Seeds of Black Tiger Shrimp, *Penaeus monodon* (Fabricius) *Asian J. Agricult. Sci.*, 2 (1): 5–8.

Gunalan, B., P. Soundarapandian, R. Kumaran, T. Anand and A. S. Kotiya 2011. First report on White Spot Syndrome Virus (WSSV) infection in white leg shrimp *Litopenaeus vannamei* (Crustacea, Penaeidae) under semi intensive culture condition in India. *AAACL Biofl.*, 4 (3): 301–305.

Karthikean, J., 1994. Aquaculture (Shrimp farming) its influence on environment. Technical Paper submitted to the Seminar Our Environment-Its challenges to development projects. American Society of Civil Engineers, Cucutta, India.

Lightner, D.V. 1996. *A Handbook of Shrimp Pathology and Diagnostic Procedures for Diseases of Cultured Penaeid Shrimp*. World Aquaculture Society, Baton Rouge, Lousiana, USA, pp. 304.

Martinez-Cordova, L.R., Villareal-Colmenares, H. and Porchas-Cornejo, M.A., 1998. Response of biota to aeration rate in low water exchange ponds farming white shrimp, *Penaeus vannamei* Boone. *Aquaculture Research*, 29, 587–593.

Moya, M., A.L. Lawrence, C.A. Collins and T.M. Samocha 1999. Acclimation of *Litopenaeus vannamei* postlarvae to 2 ppt ground saline water in Sonora Desert, Arizona. p. 424. In: Book of Abstracts. World Aquacult., Soc. Ann. Conf., Sydney, Australia.

Parker, J.C., F.S. Conte, W.S. Macgrath and B.W. Miller 1974. An intensive culture system for Penaeid shrimp. *Proc. World Maricult. Soc.*, 5: 65–79.

Paul Raj, B.B. 1999. Ecofriendly feed and management system for sustainable shrimp culture. *Fisheries World*. pp. 13–17.

Ramakrishna, R. 2000. Culture of the tiger shrimp *Penaeus monodon*
Van Wyk, P., M. Davis-Hodgkins, C.R. Laramore, K.L. Main, J. Mountain and J. Scarpa (1999). Farming marine shrimp in recirculating freshwater systems. FDACS contract M520. Florida Department of Agriculture and Consumer Services, Tallahassee, Florida, USA. 120P.

Wang, W., W. Gu, Z. Ding, Y. Ren, J. Chen and Y. Hou (2004). A novel Spiroplasma pathogen causing systemic infection in the crayfish Procambarus clarkii (Crustacea: Decapod), in China. FEMS Microbiol. Lett. 249: 131–137.

Wen-Young, T (1988). Shrimp Mariculture: A Practical Manual (2nd ed.). W.S. Aquaculture, Cannan International Pvt. Ltd., Brisbane, Australia. 282P.

Wyban, J.A., J.N. Sweeney and R.A. Kanna (1988). Shrimp yields and economic potential of intensive round pond systems. J. World Aquacult. Soc., 19: 210–217.