Effects of stocking density on blood chemistry and amino enzymatic activity of juvenile Nile tilapia Oreochromis niloticus (Linnaeus, 1758) in hyperosmotic rearing condition

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Abstract

Nile tilapia Oreochromis niloticus, a euryhaline species, is a good candidate for culture in estuarine areas in the Philippine archipelago. Thus, the effects of stocking density on plasma cortisol, blood glucose, plasma aspartate transaminase (AST), and plasma alanine transaminase (ALT) were determined. The group of juvenile Nile tilapia O. niloticus (14.67 ± 0.12 cm TL and 44.57 ± 0.48 g W) was subjected to three different stocking densities (6, 12 & 24/60 L) for 14 days in 15‰ rearing condition. The experiment was composed of three treatments with triplicated groups arranged in a completely randomized design (CRD).

Highest stocking density affected the plasma cortisol, blood glucose, plasma AST and ALT of Nile tilapia by increasing its concentration level in the blood. These manifestations suggest that overcrowding in Nile tilapia affects the physiological function through hormonal secretion, glucose utilization, and protein catabolism of the species.

Keywords: Cortisol; Aspartate transaminase; Alanine transaminase; Plasma; Glucose.

1. Introduction

Nile tilapia Oreochromis niloticus L. is a euryhaline species that has considerable potential for culture in high saline waters, especially in the Philippines where estuarine environments are common. Adaptation of Nile tilapia in a hyperosmotic condition was previously reported [1]. The tilapia industry is growing rapidly since Nile tilapia O. niloticus became more accepted by the Filipinos [2]. Tilapia is now ranked second to milkfish as the most important cultured fish in the Philippines [3]. Given the limited space for freshwater aquaculture and pressures on providing the food demands of the growing population, tilapia is now being cultured in brackishwater ponds. The brackishwater tilapia farming will further intensify in the years to come to meet the cheap protein requirements of the increasing world human population. With the invasion of tilapia production into the brackishwater, an appropriate stocking density in these farming areas should be put into consideration. And as to optimize their culture condition, rearing density should be considered in fish culture intensification [4].

Rearing density is an important aspect for fish culture, and it has been demonstrated that rearing fish at inappropriate stocking densities may impair growth and reduce immune competence due to factors such as clustering stress and the deterioration of water quality, which can affect both the feed intake and conversion efficiency of the fish [5]. However,
it is not always clear whether the performance of fish is influenced by stress caused by suboptimal water quality parameters associated with high densities (e.g., low oxygen level, elevated ammonia, or carbon dioxide levels) or by aggressive behavior due to crowding [4].

Stress in farmed fish is of considerable significance to both welfare and productivity as it has been linked to a reduction in growth [6]. Particular attention has been paid to stocking density as one of the key factors to influence the perceived level of stress in fish [5]. Physiological alterations might be used as indicators for unsuitable environmental conditions or the presence of stressors encountered in intensive fish culture [4]. Stress in fish affects its overall performance as it undergoes several responses that compromise health and growth. It commences with an initial short-latency increase in plasma levels of catecholamines released from stores in the chromaffin tissue of the kidney, followed by a longer latency but usually more prolonged elevation in plasma cortisol levels following de novo synthesis by interrenal tissue [7]. Another study manifested that stress responses of this species include an increasing level of plasma cortisol [8]. The release of glucose is triggered by the effect of cortisol as fish maintaining its homeostatic state. As stress persisted, the stored protein was converted to glucose through deamination and transamination processes to further combat the stress. Depending on the persistence of stressors until fish become exhausted and eventually die. Amino enzymes catabolism as a coping mechanism in the hyperosmotic medium was previously observed in Nile tilapia [8].

Lowering the stocking density proved to improve the overall growth performance of tilapia species [9,10]. However, extensive farming is no longer practical nowadays as it requires huge production areas. And farming towards the sea in the hope of the future generation to attain food security. Thus, this study was conducted to determine appropriate stocking density in hyperosmotic conditions without compromising the overall health capability of the species. Specifically, the hematological indices such as plasma cortisol, blood glucose, plasma aspartate transaminase, and alanine transaminase of juvenile Nile tilapia reared in hyperosmotic conditions were quantified.

2. Material and methods

2.1. Study site and duration
The study was performed for 14 days (September 12, 2014 to September 26, 2014) at the Multi-Species Hatchery of the Institute of Aquaculture (IA), College of Fisheries and Ocean Sciences (CFOS), University of the Philippines Visayas (UPV), Miagao, Iloilo, Philippines.

2.2. Animal source and acclimation
Tilapia juveniles were obtained from the Freshwater Aquaculture Station (FAS), IA, CFOS, UPV, and immediately transported to the hatchery in oxygenated plastic bags via motorbike tricycle.

Organisms were acclimated in 2 tons capacity fiberglass rectangular tanks at a density of 10 L per individual for 15 days prior to experiment proper. Organisms were fed with floating commercial feeds (Tateh® Feeds) twice a day given in the early and middle afternoon at 1% average body weight (ABW) for body maintenance. Likewise, acclimation tanks were provided with aeration, and 5 to 10% water volume was replaced on a daily basis.

2.3. Test organism and experimental design
Nile tilapia (14.67 ± 0.12 cm TL and 44.57 ± 0.48 g W) stocked at 6, 12, and 24 individuals in 60 L tanks arranged in a completely randomized design (CRD). There were 3 treatments in triplicate groups with a total of 27 experimental units used in the experiment. Nine experimental tanks were taken out in each sampling period until the end.

2.4. Experiment Proper
The experimental tanks were supplied with continuous aeration and about 70% of water volume was changed twice every day. The wastes were siphoned using a small hose every day and tanks were cleaned using sponge every other day. Experimental organisms were fed twice daily at 3% average body weight (ABW) with Tateh® tilapia starter surfer feeds.
2.5. Stocking and monitoring

The experimental tanks having low (6/60 L), moderate (12/60 L), and high (24/60 L) stocking densities were subjected to a rearing exposure of 15% salinity level. Adjustment of salinity was done by diluting seawater with tap freshwater to achieve the desired concentration of salinity. The water salinity was measured by a refractometer (ATAGO HHR-2N). Other water quality parameters such as temperature, dissolved oxygen, pH, and total ammonia were monitored using DO meter (YSI model 55-12 FT), pH meter (HI 99104), and total ammonia test kit (Advance Pharma Co., LTD) in situ.

2.6. Blood sampling in fish

A sampling of the fish blood was done thrice throughout the experiment, which carried out within 24 hrs after stocking, 7 days, and end of the culture period.

Feeding was stopped 24 hours before sampling. Before blood extraction, organisms were anesthetized using 2-phenoxyethanol at 1 ml L⁻¹ concentration as described by Morgan et al., [11]. In every sampling, 1 to 1.5 ml of blood was taken from each fish through caudal vein puncture using a heparinized needle and 1ml syringe. Blood samples were immediately placed in a styrofoam box with ice after extraction. Collected blood was centrifuged at 3000 rpm for 10 minutes to collect blood plasma using a refrigerated centrifuge (HETTICH 4903-02-0). The upper colorless fluid in the microcentrifuge tube was collected and transferred to another tube. Plasma samples were stored at -80 °C in the ultra-low freezer (ILSHIN NKH10579) until analyzed.

2.7. Sample analysis

Quantitative analysis of fish cortisol (ng/ml) was done using an enzyme-linked immunosorbent assay (ELISA) kit (SunLong Biotech Co., LTD). Enzymatic readings were determined using the microtiter plate reader (Ledetect 96 ELISA). Blood glucose (ml/dL) was quantified following the method of Mustafa et al., [12] in which ooze of fresh blood from the severed caudal peduncle was slightly dropped into the tip of the glucose strip that was inserted in a standard glucometer (Apex Biotechnology Corp.) and reading of the result was obtained in situ. Plasma aspartate transaminase (AST) (ng/ml) and alanine transaminase (ALT) (pg/ml) content were analyzed using enzyme-linked immunosorbent assay (ELISA) kit (SunLong Biotech Co., LTD) following the company standard manual.

2.8. Data Analyses

Data on plasma cortisol, plasma aspartate transaminase (AST), plasma alanine transaminase (ALT) were analyzed using Statistical Package for the Social Sciences (SPSS) version 20 in one-way ANOVA (Analysis of Variance). Duncan Multiple Range Test (DMRT) was used as a post hoc analysis to determine the significant difference among treatments. The level of significance was set at α = 0.05 and 0.01 in all data. Data presented in tables are expressed in means ± SEM in triplicate groups.

3. Results and discussion

Within 24 hours of the experiment, results showed no significant difference (p>0.05) in blood glucose. However, analysis of variance (ANOVA) showed a highly significant difference in plasma cortisol (p<0.01), and significantly increased at higher stocking density (Table 1).

Table 1 Blood glucose and plasma cortisol level of Nile tilapia O. niloticus after 24 hrs. of rearing period in 60 L tank.

| Biological Indices | Stocking Density | Cortisol (ng/ml) | Glucose (mg/dL) |
|--------------------|------------------|------------------|-----------------|
|                    | 6                | 67.97 ± 0.86b    | 175.44 ± 21.00a |
|                    | 12               | 64.89 ± 0.40b    | 143.26 ± 21.37a |
|                    | 24               | 76.70 ± 0.69a    | 197.06 ± 38.21a |
| ANOVA (P value)    |                  | 0.001            | 0.196           |

Means ± SE having the same letter are not significant (p>0.05).
Increasing plasma cortisol level was observed in response to high stocking density within 24 hrs of the culture period. Initially, the highest cortisol level was found within a high stocking density group, suggesting that high stocking density resulted in stress manifestation, which rapidly increased blood cortisol levels. The plasma cortisol means of 76.70±0.69 ng/ml found in higher stocking density conditions were higher than the 40 ng/ml characteristic of stress Nile tilapia previously reported [13]. Even though Barreto and Volpato [14] reported that the cortisol baseline for *O. niloticus* ranges from ~5 to 60 ng/ml and thus contradict Barcellos et al., [13], the cortisol level in this study suggests that fish underwent stress during rearing condition. An abrupt increase in plasma cortisol level within 24 hours of exposure might suggest an acute stress response caused by overcrowding in this species. Acute stress causes a rapid increase in plasma cortisol in other teleost species which has been reviewed [15]. And the recovery from stress in terms of the return of plasma cortisol to resting levels may be slowed if stressor still exists in the recovery environment [7].

On the 7th day of rearing, results showed no significant difference in plasma cortisol (*p*>0.05). On the contrary, blood glucose and plasma ALT were highly significant (*p*<0.01). Likewise, plasma AST was also significantly affected by the stocking density (Table 2).

**Table 2** Plasma cortisol, blood glucose, plasma AST and ALT level of Nile tilapia *O. niloticus* after 7 days of rearing in 60L tank.

| Biological Indices | Stocking Density | Cortisol (ng/ml) | Glucose (mg/dL) | AST (ng/ml) | ALT (pg/ml) |
|--------------------|------------------|------------------|-----------------|-------------|-------------|
| 6                  |                  | 90.39±5.24a      | 114.94±2.42b    | 5.01±0.16b  | 1248.83±14.54b |
| 12                 |                  | 82.80±4.99a      | 214.33±10.81a   | 5.34±0.26ab | 1313.90±2.10a |
| 24                 |                  | 84.83±6.71a      | 194.50±10.49a   | 5.65±0.35a  | 1334.89±7.57a |
| ANOVA (*p* value)  |                  | 0.159            | 0.005           | 0.019       | 0.000       |

Means ± SE having the same letter are not significant (*p*>0.05).

In this study, cortisol level did not go down into a resting level but instead even went up after 7 days of the culture period. This means that reared species at different stocking densities underwent physiological stress. The lack of significant differences (*p*>0.05) observed among high, moderate, and low stocking densities might not only be due to overcrowding effect or water quality deterioration but rather it was influenced by behavioral activities including biting and chasing which were more pronounced in a lower density. A similar observation was also reported by Barcellos et al., [13] in Nile tilapia that social stress may result in the occurrence of a chronic stress response which is probably caused by antagonistic encounters among the members of the group. This scenario persisted until day 14 of the rearing period, where high cortisol level was shown by *O. niloticus* raised at a lower density. This suggests that the hierarchy in 6 individuals per 60 L is still pronounces.

A significant increase in blood glucose towards higher stocking densities was found after a week of culture. Normally, the increase of glucose in plasma is not as rapid as for cortisol and it has been reviewed that an increase of glucose occurs hours or even days after the stress has manifested [15]. Glucose elevations are primarily generated by cortisol-mediated gluconeogenesis [7]. According to Barreto and Volpato [14] the blood glucose from 45 to 130 mg/dL is the non-stressed baseline characteristic of the Nile tilapia. A review by Pankhurst [7] elucidated that post-stress responses usually lead to increased plasma glucose. The effects of higher stocking density on the increase in blood glucose have been reported in other species [16,17]. Glucose is responsible for the provision of energy that is used by fish to cope with stress [18,19,20,21,4]. The rise of glucose during this time might be due to the density-dependent factors mainly attributed to social stress brought about by the aggressive encounters and dominant/submissive relationships that are common in this species [21].

By the end of the study, all biological indices were significantly affected by stocking density. However, plasma cortisol level was higher in low stocking density while higher blood glucose concentrations were observed at higher stocking density. Likewise, similar trends were also manifested in plasma AST and ALT (Table 3).
Table 3 Plasma cortisol, blood glucose, plasma AST and ALT level of Nile tilapia *O. niloticus* after 14 days of rearing in 60L tank.

| Stocking Density | Biological Indices                                      |
|------------------|--------------------------------------------------------|
|                  | Cortisol (ng/ml) | Glucose (mg/dL) | AST (ng/ml) | ALT (pg/ml) |
| 6                | 162.72±3.93a    | 134.12±23.24ab  | 4.10±0.06b  | 1292.91±3.64b |
| 12               | 114.50±8.88b    | 114.83±13.41b   | 4.41±0.19a  | 1229.94±3.64bc |
| 24               | 139.50±11.28a   | 200.34±19.27a   | 4.91±0.28a  | 1437.75±20.09a |

Means ± SE having the same letter are not significant (*p* > 0.05).

The plasma aspartate transaminase (AST) was significantly elevated at high stocking density at the end of the study. The degree of response was caused by the influence of stocking density as a stress factor which suggests that increased stocking density caused stress to the experimental organisms. The previous finding stated that aspartate is one of the major glucogenic precursors and important energy substrate for fish thus increasing activities of plasma AST indicated amplified transamination processes [22]. Besides, aspartate is essential to purine nucleotide synthesis in all cell types. Aminotransferases are intracellular enzymes that are normally localized within the cells of the liver, heart, gills, kidney, muscle, and other organs. The levels of these enzymes increase in the plasma when the cells are damaged or their membranes disrupted, allowing the enzymes to leak out of the cells [23]. Higher stocking density caused higher plasma AST as the stress response was also reported previously in Nile tilapia *O. niloticus* and they also concluded that stocking density of tilapia up to 6 fish m$^{-3}$ showed an adverse effect on the liver functions [24].

Similarly, other work reported that 3 fish m$^{-3}$ obtained the lowest value of AST in silver carp, *Hypophthalmichthys molitrix* fingerlings suggesting that fish were essentially normal and within the range consistent with good fish health [17]. Previous study mentioned that stress elevates the internal oxidizing effort towards membrane permeability that increases the fluxes of AST enzymes into the bloodstream [25]. Kamal and Omar [17] reported on silver carp *H. molitrix* that plasma AST increased at higher stocking density. This parameter is known as a sensitive indicator of even minor cellular damage in the liver of fish.

Effect of stocking density as stressor caused higher plasma alanine transaminase (ALT) at the end of the culture period. Aside from aspartate, alanine is also a major glucogenic precursor and important energy substrates for fish. Moreover, alanine is a preferred carrier of nitrogen for inter-organ amino acid metabolism [22], as several amino acids can be converted to alanine, released to the blood, and used as a fuel source in other tissues in fish [26].

Ovie and colleague [23] also suggested that aminotransferases play vital roles in carbohydrate-protein metabolism in fish and other organisms’ tissues. The aminotransferases occupy a central position in the amino acid metabolism as they help in retaining amino groups (to form new amino acid) during the degradation of amino acid and are also involved in the biochemical regulation of intracellular amino acid pools. They help in providing necessary intermediates for gluconeogenesis. The observed alterations in their activities in the exposed fish may, therefore, have an adverse effect on the amino acid metabolism of the tissues and consequently the intermediates required for gluconeogenesis [23]. The levels of these enzymes increase in the plasma when the cells are damaged or their membranes disrupted, allowing the enzymes to leak out of the cells. These enzymes are therefore of major importance in assessing and monitoring liver cytolysis [23]. Thus, the present experiment implies that higher stocking density may cause organ damage. Plasma enzyme levels depend on the rate of release of enzymes from damaged cells, which in turn depends on the rate at which damage is occurring and at the extent of cell damages [23]. This result indicates the consistent stress effect of increasing the stocking density on liver functions of the fish as reported by Kamal and Omar [17] on silver carp *H. molitrix*. This result further indicates fish liver tissue destruction. ALT concentrations would increase because of its role in an initial amino acid compensation and formation that the body needs in physiological changes that require increased energy demands. Further, this also suggests that the increase in plasma ALT was due to infiltration of the enzymes to the bloodstream [25]. This was in line with the work of Melotti et al., [27] in rainbow trout brought by a high density of the
This result corroborates that of Aly et al., [24] who reported in Nile tilapia that ALT levels were significantly increased by increasing stocking density. This was attributed to the stress response that has been associated with increased amino acid levels as a consequence of activated catabolism and inhibited protein synthesis. On the other hand, the elevation in the ALT was attributed to the damage of hepatic cells and other tissues and escaping of this enzyme to the blood [24].

The experiment was done indoor and utilized small containers to single out factors affecting the physiological aspect of Nile tilapia in the brackishwater farm. However, it was revealed that social aggressiveness was also triggered by small confinement in this species. Thus, species in the farm site might show different characteristic behavior compared to the result of the study. With this regard, a future study on the stocking density of Nile tilapia should consider several factors such as; uniformity of size, sex, water flow, and shading.

4. Conclusion

Findings of the present study suggest that stocking density affects the physiological function of Nile tilapia *O. niloticus* by increasing the cortisol level, blood glucose, and levels of amino acid catabolizing enzymes that include ALT and AST. These results add to the present knowledge on the physiological responses and coping mechanisms of Nile tilapia reared in an overcrowding hyperosmotic environment.

Compliance with ethical standards

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Disclosure of conflict of interest

Author declares no possible conflict of interest from any individual, party or entity.

Statement of ethical approval

The experiment was performed based on the ethical approved guidelines set by the UPV on the humane act of handling and dissecting the specimen during sampling.

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