Dynamic of Stress Response in Victoria Labeo (Labeo victorianus) during transfer from the Hatchery to cages and ponds under Differential Caged Stocking Densities

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Abstract: Variation in fish stocking densities translate to difference in growth performance, yields and economic benefits in fish culture. Transferring fish directly from hatcheries to the cages or ponds may induce stress. We evaluated the stress response of Labeo Victoria (Labeo victorianus) in an integrated cage-cum-pond culture during transfer of fish from the hatchery to the cages and ponds at different cage stocking densities. Cages were stocked at varying densities of 10, 30, 60, 90, 120, 150 and 180 fish/m³ and suspended in a static pond of 200 m². The L. victorianus fingerlings of a mean weight 23.6 ± 1.8 g were stocked in the cages and the pond respectively. 20 fish were sampled during the transfer period from the hatchery to ponds and cages for analysis of primary and secondary parameters of stress response. Primary stress response occurred when fish were directly transferred to cages and ponds at stocking density ≥ 60 fish/m² and 90 fish/m² respectively. Parameters of secondary stress response occurred in fish transferred to the cages at stocking density ≥ 120 fish/m² and in ponds at density ≥ 150 fish/m². Transfer of fish directly from the hatchery to the ponds induce primary and secondary stress.

Keywords: Stress response, Cortisol response, Integrated cage-cum-pond culture, Stocking density, Labeo victorianus

1.Introduction

Due to the increasing demand for fish in the world, intensification of fish culture has increased in many aquaculture units. To utilize space, fish are stocked in cages and in ponds simultaneously (referred to as cage-cum-pond-integrated system) [1-3]. Success of this system depends on the yields obtained from cages and open ponds. A key challenge in this system has been to determine the stocking density to maintain sustainable critical standing crop (CSC). When fish is cultured in low density, the overall yields will be low, yet high stocking density beyond the CSC may negatively affect the fish growth [4]. During stocking, fish are transferred directly from the hatchery or grow-out ponds to the cages and ponds. Since there is no recommended stocking density for all fish, various stocking densities above the CSC may induce stress in fish. Yet very little is known concerning stress response during transfer of fish from hatcheries to the cages and ponds.

The response to stress in fish is characterized by the stimulation of the hypothalamus, which results in the activation of the neuroendocrine system and a subsequent cascade of metabolic and physiological changes [5]. Under conditions of stress, the body of the fish emits immediate responses recognized as primary, secondary and tertiary responses. The primary response is the perception of an altered state by the central nervous system (CNS) and the release of the stress hormones, cortisol and catecholamines (adrenaline and epinephrine) into the bloodstream by the endocrine system [6]. Secondary responses occur as a consequence of the released stress hormones [7], causing changes in the blood and tissue chemistry, e.g. an increase of plasma chloride and sodium and blood ammonia [8]. Therefore, some plasma chemicals may be useful tools to evaluate the health and/or stress condition of the fishes [9]. Yet there is very little work that has been done using stress response parameters to evaluate the stress response when fish is transferred directly from hatchery to the cages and ponds. The aim of the current study was therefore to evaluate the stress response in Labeo Victoria (Labeo victorianus) during transfer from the hatchery to cages and ponds under differential caged stocking densities.

2.Materials and Methods

The study was conducted at Mwea Aquafish Farm in Kenya (0°36.73'S, 37°22.84'E, 1208 m asl). The cages used in this study were 3-m³ floating cages consisting of a wooden frame (2 m × 1.5 m × 1 m) covered with a half inch wire mesh. Each cage had a polyvinyl chloride (pvc) ring of 30 cm diameter feeding tray to prevent direct spillage of the artificial experimental feed (sinking pellets). Twenty one, 200 m² earthen ponds of 1.0 m depth in the shallow end and 1.5 m depth in the deeper ends each with a floating cage were used in this study. Broodstock of the L. victorianus (mean weight = 521.4 ± 34.3 g) were transferred to the hatchery at the Fish Farm. Larvae were obtained through induced breeding and semi-natural spawning. They were cultured for a period of 42 days in a hatchery. During the culture period, the larvae were fed Artemia nauplii. After 42 days, the juveniles were then sorted out and randomly transferred to the experimental cages and open ponds. The stocking weights of the juveniles were 23.7 ± 2.6 g. The
stocking densities in cages were: 10, 30, 60, 90, 120, 150 and 180 fish m\(^{-3}\). All treatments were executed in triplicate. The ponds were stocked with juveniles at a stocking density of 2 fish m\(^2\). The fish were hand fed sinking diet in the morning (0800 h) and in the evening (1700 h). Feed were supplied at 2.5% of the caged biomass.

24 hours after the transfer, a total of 20 fish were sampled from each pond and a similar number from the cages using seine and dip nets respectively. Water samples were taken from four different locations (near inlet and outlet, in the middle of the pond and close to the shore along the pond length) in each pond and within the cages, approximately two hours after morning and evening feeding using 1.12-m long column sampler. Samples from the ponds and those from the cages were pooled separately to provide an integrated sample then a sub sample was drawn from the integrated sample for the analysis of the various water quality parameters. Dissolved oxygen (DO), pH and salinity were measured in situ in the cages and in ponds, using a calibrated JENWAY 3405 electrochemical analyser (Barloword Scientific Ltd, Essex, UK). Water sampled were filtered through Whatman glass filter paper GF/F and analysed for nitrate nitrogen (NO\(_3\)-N) using cadmium reduction method, phosphate phosphorus (PO\(_4\)-P) by the standard ascorbic acid method and total ammonia nitrogen (TAN) by the indophenol blue method following detailed procedures in APHA (1998) [10].

A total of 20 fish from each cage and pond, were randomly captured with dip nets and quickly anaesthetized with benzocaine (5 mg/L) for 2-3 min. Blood was withdrawn from the caudal vein of each sampled fish into 1 ml heparinized insulin syringe. Blood glucose was measured according to King and Garner (1947) [11]. Heparinized blood was centrifuged at 3000 r.p.m. for 10 min and plasma frozen at -196°C in liquid nitrogen for further plasma cortisol (Radioimmunoassay with a Coat-to-Count Kit, Diagnostic Products Corporation, Los Angeles, CA, USA), chloride (SIGMA kit 461, Sigma Diagnostics, USA), blood ammonia (Nessler's method modified by Gentzkow and Masen, 1942) and plasma sodium (flame photometry) analyses. Capillary tubes with blood samples were centrifuged to separate cells from plasma. Plasma osmolality was then determined using Wescor 5520 vapour pressure osmometer.

Normality of the data was determined using Shapiro-Wilk test while homogeneity of variance was ascertained using Levene’s test. Difference in stress response was determined using ANOVA. Multiple comparisons for significantly different means was done using Tukey test. All the statistical analyses were done using the SPSS 17.0. Differences were considered significant at p < 0.05.

3. Results

Changes in water quality parameters during transfer of fish from the hatchery to the cages and ponds at different stocking densities are provided in Fig. 1. DO concentration decreased significantly from 10.6 to 2.83 mg/l in cages and 8.89 to 0.78 mg/l in ponds. Cages had significantly (P < 0.05) higher DO than ponds. Similar patterns of reduction were also observed for pH: in cages, pH reduced from 7.7 to 5.6 while in ponds it reduced from 7.4 to 5.1. Salinity, NO\(_3\)-N, PO\(_4\)-P and T-NH\(_4\)-N all increased significantly with increasing stocking density. Salinity values between 4 to 8 ppm were recorded in cages and ponds when fish was transferred to cages and ponds of stocking density 10, 30, 60 and 90. However above 90 fish/m\(^3\), salinity values increased significantly with increase in ponds being significantly (P < 0.05) higher at all stocking densities. Concentration of NO\(_3\)-N, PO\(_4\)-P and T-NH\(_4\)-N increased with each stocking density however, there were no significant differences in the increase between cages and ponds (P > 0.05).
Stocking fish in cages and ponds directly from the hatcheries resulted in significant (P < 0.05) increase in all physiological parameters (Fig 2). Plasma cortisol increased above the normal values in fish (80-90 ng/mg) at stocking densities ≥ 90 fish/m³ in cages and at stocking density ≥ 120 fish/m³ in ponds. Above density of 60 fish/m³ cortisol levels was significantly higher in cages than ponds. Plasma glucose displayed similar changes being above that for fish under normal condition (70-90 mg/dl) at stocking density ≥ 120 fish/m³ in cages and ≥ 150 fish/m³. Secondary stress parameters i.e. plasma sodium, plasma chloride and blood ammonia, all displayed similar changes when fish was transferred from the hatchery at different caged density; they all increased above that of normal physiological levels in cages at density ≥ 90 fish/m³ and in ponds ≥ 120 fish/m³ respectively. In cages and in the ponds, osmolality of the fish increased above fish at normal physiological condition at caged density of ≥ 120 fish/m³ and ≥ 90 fish/m³ respectively.
4. Discussion

Management practices used daily in aquaculture necessarily generate stressors, to which the fish being grown may respond dissimilarly. Moreover, most freshwater fish have their body fluids hyper-osmotic (osmolality of 260–330 mOsm/kg H2O) with respect to their external medium and may experience osmoregulatory disturbances in freshwater if stressors are introduced. In the current study, directly transferring *L. victorianus* from the hatchery to cages and ponds resulted in a characteristic increase in primary response parameters (cortisol and glucose levels) at ≥ 90 fish/m³ in cages and at stocking density ≥ 120 fish/m³ in ponds. Similar, stress responses have been observed in other species subjected to different forms of abrupt shock or stressful conditions [12-14]. Plasma cortisol and glucose concentrations observed in our study were generally within the range reported for stress elevated values of most teleost fishes (30–300 ng/mL; [5]). Prestress plasma cortisol concentrations in *L. victorianus* (80-90 ng/mg) are somewhat higher relative to that of several other fish species examined. The pre- and poststress concentrations of plasma cortisol of 12 species indicate ranged from 1.0 to 11 ng/mL [5] suggesting that in the environment where the current fish was reared may have been stressful. The increase in cortisol and glucose as primary response factor may have been caused by exhaustion of the endocrine system as a result of prolonged hyperactivity [15].

The secondary response parameters (plasma sodium, plasma chloride and blood glucose) were induced at stocking density ≥ 90 fish/m³ in cages and ≥ 120 fish/m³ in ponds. Significant increase in plasma sodium and plasma chlorides of fish after transfer to ponds and cages at high stocking densities could be due to gains of Na⁺ and Cl⁻. The increase in the plasma sodium and plasma chloride is a possible indicator of impairment in ionic (both Na⁺ and Cl⁻) regulation of fish due to the stressful condition. The high TAN after transfer suggests an increased physiological activity of the fish during osmoregulation. The accumulation of ammonia nitrogen due to increased metabolites may cause serious problems to the fish, such as increased oxygen consumption and heart rate, decreased plasma sodium and alteration of the acid–base balance [16].

This is the first evaluation of stress response during transfer of fish from hatchery to the ponds and cages. Differences in the magnitude of stress responses among fish species is common. Barton's review describes post-stress concentrations that ranged from as low as 3.0 ng/mL in pallid sturgeon to as high as 229 ng/mL for walleye *Sander vitreus* [5]. Several factors can influence the stress response, including prior stress exposure and the intensity and duration of the stressor. In our study the changes in primary and secondary stress responses was closely correlated with water quality changes and is our first suspicion as the main cause of stress response during transfer of fish from grow-out to the hatcheries.

5. Conclusion

Abrupt transfer of fish from hatcheries to the cages and ponds resulted in increased primary and secondary stress response in tandem with water quality changes. The observed levels of stress response parameters are within the range of values observed for other species and provide a baseline for future evaluations of stress physiology and the development of aquaculture conditions and techniques suitable to this species.

6. Future Prospects

Protocol for culture of fish involves larval production and subsequent transfer of fish from the hatchery to the ponds and in the cages for stocking purposes. The present study reveal that such abrupt transfer result in stress response when done at different densities above which some are found to induce both primary and secondary response. Such information is vital for future management of stock recruitment.

References


Author Profile

Miss Gladys M. Kuria has an MSc in Aquaculture from the University of Eldoret and has conducted research in fish culture. She has been actively involved in aquaculture projects and activities that focus on growth, yield, survival and economic impact She is currently enrolled for a PhD in Aquaculture at the University of Eldoret.

Professor Charles C. Ngugi graduated with a Doctorate in Fish Biology from Memorial University, Newfoundland, Canada in 1995 and Master’s in Aquaculture, Rivers State University of Science and Technology, Nigeria. He has wide range of publications in the areas of: Development of Aquaculture Systems (Static and Raceways), Aquaculture Planning and Management, Fish Biology; and Fish Populations Assessment.

Dr. Elijah Oyoo-Okoth graduated with A PhD in Aquatic Ecology and Ecotoxicology from the University of Amsterdam in 2012, and MSc in Fisheries and Aquaculture Sciences from Moi University. He is an international scholar with several publications in the field of Aquaculture. Now he is working on improving fish culture by establishing protocols for culture of Victoria Labeo.