Muscle pigmentation of Nile tilapia (*Oreochromis niloticus*) fed on crude palm oil incorporated fish feed

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Abstract The effects of diets containing crude palm oil on muscle pigmentation and deposition of carotenoids in tilapia (*Oreochromis niloticus*) was studied. A total of 135 advanced fingerlings with an average body weight and standard length of 9.11±4.78 g and 8.1±0.8 cm, respectively were stocked in tanks for 8 weeks and fed with formulated diets containing crude palm oil (CPO) as the oil component. Weight, standard length, and carotenoid levels in fish were measured biweekly. Muscles of fish fed with the control diet and CPO incorporated diets had carotenoid levels of 0.14±0.03 µg/g and 0.28±0.01 µg/g, respectively. Skin of the fish had carotenoid levels of 3.24±0.02 µg/g in the control and 6.06±0.03 µg/g in the treatment. According to the results 3% CPO incorporated diet had a significant effect on enhancing flesh colour in tilapia. The sensory evaluation indicated that the flesh of CPO fed tilapia fillets were more attractive than the fillets of fish fed control diet. CPO oil is recommended as a natural food colourant for improving flesh quality of Nile tilapia fillets.

Keywords: *Oreochromis niloticus*, Carotenoids, β-carotene, pigmentation

INTRODUCTION

Carotenoids are a group of phytochemicals responsible for yellow to orange colour of fish (Rebecca et al. 2014). These pigments are derived from dietary sources because the fish are unable to biosynthesize carotenoids (Foss et al. 1984). The orange-pink color of fish fillets is considered a visual indication of "quality" by consumers. Hence, supplementing feed with astaxanthin or canthaxanthin to obtain red-colored flesh is a common practice in fish farming. Carotenoids are known as pro-vitamin A and immunoregulator which absorb light in the 400-500 nm range of the visible spectrum (Rebecca et al. 2014; Foss et al. 1984; García-Chavarría et al. 2013.). Carotenoids are amenable for development of the skin colour of ornamental fish as well as the colour of flesh of food fish (Jintasataporn and Yuangsoi 2012). Synthetic carotenoids are used as dietary supplements to enhance the pigmentation of fish (Kalinowski et al. 2005). Many fish accumulate carotenoids in their integument, flesh and gonads. The effectiveness of deposition and pigmentation of carotenoids is source and species specific. Certain fish have the ability to convert one form of carotenoids into another (Kang and Ha 1994).

In fish, there is no common mechanism for the metabolism of carotenoids and their subsequent derivatives (Das and Biswas 2016). Scientists believe there is a metabolic pathway for carotenoids in the liver and intestines (Aas et al. 1999). Selvakumar et al. (2011) have classified fish into two groups based on the capacity of metabolization of carotenoids. The first type of fish requires an inclusion of specific oxygenated derivatives in to their diet as they are unable to oxidation of ionone. The second type of fish such as gold fish or the fancy red carp can oxidate 4 and 4’ positions of ionone ring and hence have the potentiality of conversion of zeaxanthin and lutein to astaxanthin (Gouveia et al. 2003).
As the aquaculture feed industry seeks natural, environmentally friendly sources of pigments to improve colour and to enhance consumer acceptability, there is a great potential for using natural carotenoids in the industry. It paves the way for many aquaculture feed industries to promote their products as natural with a clear shift away from synthetic ingredients and colourants (Rebecca et al. 2014). Natural carotenoids derived from animals such as crustaceans are limited in supply as there is a declining trend in catches (Gupta et al. 2007). Plant-based foods such as yellow corn, corn gluten meal, vegetable oils and red pepper are also used as natural sources of carotenoids in aquaculture feed formulations. Crude palm oil (CPO) contains the highest known concentration of carotenoids from vegetable oils (Zeb and Mehmood 2004).

CPO is a lipid, extracted from the fleshy orange-red mesocarp of the fruits of the oil palm tree Elaeis guineensis (Zaliha et al. 2015). It is the world’s richest natural plant source of retinol, which is a kind of carotene (Jain et al. 1990). CPO (known as red palm oil), can be extracted either by wet or dry processes (Mancini et al. 2015). As it has a balanced fatty acid composition, the level of saturated fatty acids is almost equal to the levels of unsaturated fatty acids (Koushki et al. 2015). It contains both healthy and beneficial compounds such as triacylglycerols, vitamin E, carotenoids, phytosterols, as well as impurities such as phospholipids, free fatty acids, gums, and lipid oxidation products (Mancini et al. 2015).

The main carotenes present in CPO are β-carotene (56%) and α-carotene (35%) which are pro-vitamin A. Natural antioxidant, tocopherol and tocotrienol contents are 600 and 1200 ppm respectively (Koushki et al. 2015). The fraction obtained from refining, bleaching and deodorization of CPO is palm olein. The oil obtained after the refining processes is paint yellow, soft and stable (Mancini et al. 2015).

Red Palm Oil (RPO) and palm olein (PO) have good oxidative stability due to the presence of natural antioxidants and the absence of linolenic acid (Koushki et al. 2015).

Tilapia is a widely farmed freshwater fish in the world. The consumer preferences of tilapia could be increased by changing the colour of flesh. Yanar et al. (2007) conducted an experiment using three carotenoid sources; marigold flower, synthetic astaxanthin and red pepper, and the highest carotenoid level was observed in the fish flesh with synthetic astaxanthin followed by red pepper and marigold flower. The effect of different carotenoids present in tilapia fish (Oreochromis niloticus) fed for 80 days with feed containing astaxanthin and the bacterial Rubrivivax gelatinosus have resulted in increasing carotenoid concentration and the redness in the fish fillets (Valente et al. 2016).

The present study investigated value addition of Oreochromis niloticus fillet by carotenoid pigmentation on muscle using β-carotene rich sources of crude palm oil (CPO) in order to increase consumer preference.

**METHODOLOGY**

A total of 180 advanced fingerlings of Nile tilapia (O. niloticus) (~7 cm of standard length) was obtained from Udawalawe tilapia breeding centre, Sri Lanka. Fingerlings were transported in oxygenated polythene bags to aquarium and acclimatized for a period of two weeks. During acclimatization period, fingerlings were kept in 4 fiber-glass tanks (380 L) and fed a commercial fish feed, to the level of satiation.

Circular fiberglass tanks (375 L) were cleaned and filled with de-chlorinated water up to 2/3 of the tank volume (250 L). After acclimation, fingerlings, each with a body weight of 19.11±4.78 g and a standard length of 8.1±0.8 cm, were selected and randomly stocked into six fibre-glass tanks with stocking density of 15 fingerlings per tank.

The diets contained 35% protein and were prepared using fish meal, soybean meal, wheat flour, coconut meal, vitamin and mineral mixture and palm olein (PO) for control diet. PO was replaced by CPO in preparing the test diet.

The required amount of each ingredient was weighed (Table 1) and mixed with hot water (50°C). Feed was prepared according to literature procedures (Craig et al. 2017).

Prepared test diets were crumbled into desirable size according to the size of fish’s mouth gape. Fish were fed three times a day at 09.00, 13.00, 17.00 hours up to satiation. Three tanks fed with diet contain CPO and the other three tanks were fed with diet contain PO as the control diet. Daily feed consumption was measured separately for each tank.

During the period of the experiment, a constant water volume was maintained in all tanks. Each tank
was continuously aerated and cleaned daily before the first feeding by siphoning off accumulated waste materials. Approximately 100 L of water in each tank was replaced with de-chlorinated tap water.

**Table 1** Diet formulation and proximate composition of experimental diets

| Ingredients          | Control | CPO   |
|----------------------|---------|-------|
| Fish meal            | 38      | 38    |
| Soybean meal         | 16      | 16    |
| Coconut meal         | 11      | 11    |
| Wheat flour          | 31      | 31    |
| Vitamin and Mineral mix | 1      | 1     |
| Oil                  | PO-3    | CPO-3 |

**Proximate composition (% dry matter)**

|          | Control     | CPO         |
|----------|-------------|-------------|
| Crude protein | 37.03±0.02  | 37.11±0.04  |
| Crude lipid  | 4.28±0.04   | 4.47±0.11   |
| Ash (%)      | 10.04±0.02  | 10.45±0.04  |
| Moisture (%) | 5.92±0.07   | 6.32±0.07   |

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**Data collection of fish growth performance**

Initial length (cm) and weight (g) of fish stocked in the treatment and control tanks were measured and repeated biweekly. The liver and somatic weights of each dissected fish in each tank were measured biweekly. Percent length gain (LG), percentage weight gain (WG), specific growth rate (SGR) and survival rate were determined as described by Dedeke et al. (2013). Feed conversion ratio (FCR) and feed conversion efficiency (FCE) were calculated by the formulae described by Craig et al. (2017). Protein efficiency ratio (PER) and percentage average daily gain (ADG) were calculated according to formulae described by De Silva et al. (1994) for each fish during the experiment. The feed consumption was calculated using the formula put forward by Güray et al. (2012). The Hepato-somatic index (HSI) of fish was determined as described by Sadekarpawar and Parikh (2013).

**Crude protein**

A 0.5-1.0 g of dried sample (fish and diet) was used to analyse crude protein by AOAC (2003) method by Micro-kjeldhal apparatus.

**Crude lipid**

Crude lipid content was determined by Folch method using dried and ground sample (0.2 g) of flesh of fish Folch et al. (1957).

**Total carotenoid content in flesh and skin**

Total carotenoid content in flesh and skin of fish was determined using the method described by Yanar et al. (2007). Fish were sacrificed without using chemicals. Flesh and skin were separated and ground separately. Ground skin and flesh samples weight of 1g were transferred into dry glass bottles separately. A volume of 10 mL acetone was added and mixed with skin and flesh samples Anhydrous Na₂SO₄ (2 g) was added to the mixture was and it was centrifuged at 5000 rpm for 5 minutes. The sealed glass tubes were then stored at 4°C. After 3 days, the supernatant of the extract was carefully drained off, and the absorbance was measured at 450 nm using UV-Visible spectrophotometer (HACH, DR3900).

**Saponification value**

Saponification value (as mg KOH/1 g of oil) was determined by ASTM D558-95 (2017) method using 1.5-2.0g sample.

**Sensory Evaluation**

Sensory evaluation was performed by displaying the samples of fish fillets obtained from fish fed
with the control and experimental feeds without providing any identification labels. The experts recorded their preference along a four-point hedonic scale that varied from 1 (not attractive) to 4 (very attractive).

**Statistical analysis**

All the data were tested for normality of distribution and homogeneity of variance. Data on pigment levels of fish, standard length and weight which were recorded separately for each test diet were analysed by Randomized block ANOVA from SPSS 16. The significant differences between test diets on WG, LG, SGR, FCR, FCE, PER, feed consumption (FC) and hepato somatic index (HSI) were determined using t-test at minimum significance level of p < 0.05.

**RESULTS AND DISCUSSION**

**Daily Feed Consumption**

The mean daily feed consumption of fish fed experimental diet and the control diet is given in Table 2. The mean daily feed consumption was 4.63±0.58, and 4.23±0.50 for control and CPO diet, respectively, which were significantly different among different treatments (p<0.05). Figure 1 illustrates the mean feed consumption of fish during the study period.

**Growth parameters**

Growth parameters of fish fed control diet and test diet are tabulated in table 2. The mean total body length (SL) of fish ranged from 10.8±0.7 to 11.0±0.1 cm at the end of the experiment. Similarly, the mean total body weight is ranging from 45.14±8.54 to 48.33±12.07 g. Length gains (%LG) of fish fed with control diet and the test diet were 6.1±0.5 % and 6.5±1.25 %, respectively during the experimental period. The mean percentage weight gain has a similar trend which has the values of 63.4 ±11.2% for the CPO diet. There were no significant differences of SL, final weights of fish between the control and experimental diet.

The final mean specific growth (%SGR) rates of fish at the end of the 8-week period fed with control and CPO incorporated diet were 2.02±0.13% and 1.96±0.39%, respectively. There were no significant differences between the 2 types of treatments.

Percentage WG and LG were usually considered as the most important measurements of the productivity of experimental feeds. In the present study, WG and LG were not significantly affected by the different diets. According to Hu et al. (2006), Nile tilapia fed diets supplemented with β-carotene showed gradual increase of weight. Similar results have published showing the effect of feeding
four carotenoids sources; commercially available astaxanthin, Dunaliella salina extract, crayfish meal (Cherax quadricarinatus) and Squilla sp. meal on growth performance, stress resistance, lipid oxidation and carotenoids pigmentation in the flesh of red tilapia (Oreochromis sp.). These studies that were conducted for a period of 14 weeks showed highly significant differences from the control treatment in carotenoids flesh content, survival rate and growth performances. Compared to other groups, Tilapia fed astaxanthin incorporated diet had been exhibited higher resistance to stress test (Arous et al. 2014).

According to the results CPO had the highest crude protein, lipid, and fatty acids as well as the TCL. Compared to the test diet, a higher growth was not shown by the fish fed with CPO diet.

There were no significant differences in growth between the fish fed with control and test diets due to same nutrient content in both diets. According to Valente et al. (2016), no significant differences were observed in final body weight, specific growth rate or voluntary feed intake between Nile tilapia fed with diets treated with lutein rich Ulva spp. and the control.

**Feed Conversion Ratio (FCR)**

The higher FCR value of 1.78±0.12 was observed in the fish fed control diet while FCR value of fish fed test diet was 1.19±0.12 (Table 2). However, they were not significantly different. FCR is an indicator of feed but may also include the performance of the person feeding the fish, the fishes’ health, and cost effectiveness of using a particular feed (Luzzana et al. 2003). The lowest FCR value was shown by fish in CPO treatments.

**Feed Conversion Efficiency (FCE)**

In the present study, the mean FCE values for control and test diets were 40.01±11.18 and 46.74±7.53, respectively (Table 2) and they were not significantly different (p >0.05). This indicates that the fish growth had not been influenced by the amount of carotenoids used in oil samples.

### Table 2 Growth performance of fish in each test diet.

| Parameter                             | Control diet | CPO diet  |
|---------------------------------------|--------------|-----------|
| Initial length (SL-cm)                | 8.0±0.8<sup>a</sup> | 7.8±0.6<sup>a</sup> |
| Final length (SL-cm)                  | 10.9±0.9<sup>a</sup> | 10.8±0.7<sup>a</sup> |
| Initial weight (g)                    | 19.38±0.00<sup>a</sup> | 18.60±0.00<sup>a</sup> |
| Final weight (g)                      | 48.33±0.00<sup>a</sup> | 45.15±0.00<sup>a</sup> |
| percent Length gain (SL)              | 6.1±0.5<sup>a</sup> | 6.5±1.2<sup>a</sup> |
| percent Weight gain (WG)              | 63.41±11.2<sup>a</sup> | 57.76±8.75<sup>a</sup> |
| Daily feed consumption (BW/Day)       | 4.62±1.58<sup>a</sup> | 4.23±1.51<sup>b</sup> |
| Specific growth rate (SGR)            | 2.01±0.13<sup>a</sup> | 1.95±0.39<sup>a</sup> |
| Feed conversion ratio (FCR)           | 1.41±0.07<sup>a</sup> | 1.19±0.12<sup>a</sup> |
| Feed conversion efficiency (FCE)      | 40.00±11.18<sup>a</sup> | 46.74±7.53<sup>a</sup> |
| Average daily gain (%ADG)             | 3.33±0.33<sup>a</sup> | 3.23±0.68 |
| Protein efficiency ratio (PER)        | 1.08±0.30<sup>a</sup> | 1.26±0.20<sup>a</sup> |
| % Survival rate                       | 100.00±0.00<sup>a</sup> | 100.00±0.00<sup>a</sup> |
| Hepato-somatic index (HSI)            | 0.46±0.03<sup>a</sup> | 0.60±0.04<sup>b</sup> |

Means with different superscripts in each row (a, b, c) are significantly different (p<0.05)
Average Daily Gain (%ADG)
The calculated mean %ADG values for control and CPO incorporated diets fed fishes were 3.33±0.33% and 3.31±0.68% consecutively (Table 2) without significant differences.

Protein Efficiency Ratio (PER)
The calculated mean values of PER for the control and CPO diet fed fish were 1.08±0.30 and 1.22±0.8 (Table 2). In the present study, crude protein levels were 37.03±0.02 and 37.11±0.04 in control and CPO diets. PER is also influenced by digestibility and environmental conditions. According to the experiment conducted with Nile tilapia, Hu et al. (2006) reported that PER values were gradually increased with the diets supplemented with β-carotene. The results indicated that the addition of carotene rich sources did not affect none of the above parameters of fish (Bolger et al. 1989).

Survival Rate (%)
The survival rate of fish for test diets of control and CPO were 100%. (Bolger et al. 1989) have conducted a study using sex-reversed red tilapia to determine the effects of synthetic carotenoids of astaxanthin, zeaxanthin and β-carotene over spirulina incorporated diet, and results have shown that neither synthetic carotenoids nor spirulina carotenoid produced an effect on fish growth or survival.

Wang et al. (2006), who investigated whether the dietary carotenoid supplements could make differences in survival, growth, pigmentation and antioxidant capacity of characins Hyphessobrycon callistus, reported that there were no differences in growth and survival rate of the fish after 8 weeks in the experimental period. Also, body astaxanthin and β-carotene content have increased with increasing dietary carotenoid concentration (Wang et al. 2006).

Hepato-Somatic Index (%HSI)
The mean value of HSI of fish fed with two diets of control and CPO showed 0.46±0.03 and 0.60±0.04 respectively (Table 2). Hepato-somatic index (HSI) is associated with the liver energetic reserves and metabolic activity (Ighwela et al. 2014). When the feed is available in large quantity and conditions are favourable, it causes to increase the HSI value. The diets do not effect of HSI or body weight on fish.

Saponification value of oil
The saponification value of CPO and palm olein were 192±0.1 and 194±0.01 in mg KOH/g respectively. The greater the molecular weight, the smaller number of fatty acids is liberated per gram of fat hydrolysed (Yanar et al. 2007).

Carotenoids analysis
Total carotenoid levels were 481.00±0.01, 2.11±0.02 for CPO, and palm olein, respectively before adding to feed. The highest was the CPO which was rich in β-carotenes. The variation of absorbance with the wavelength of each oil sample was shown in Figure 2. The highest total carotenoid level was shown by CPO (Table 3).

Table 3 Total carotenoids level in fish skin and flesh

| Time   | Control | CPO   |
|--------|---------|-------|
|        | Skin    |       |
| Initial| 2.56±0.02a | 2.57±0.02a |
|        | Flesh   | 0.06±0.01a | 0.06±0.01a |
| 3 weeks| Skin    | 2.57±0.04b | 3.96±0.03a |
|        | Flesh   | 0.12±0.05b | 0.19±0.14a |
| 5 weeks| Skin    | 2.88±0.05b | 5.67±0.01a |
|        | Flesh   | 0.13±0.03b | 0.22±0.02a |
| 8 weeks| Skin    | 3.24±0.02b | 6.06±0.03a |
|        | Flesh   | 0.14±0.03b | 0.28±0.01a |

Means with different superscripts in each row (a, b) are significantly different (p<0.05)
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Figure 2 The variation of absorbance of CPO (×40 diluted with acetone) and palm olein ((×10 diluted with acetone) with wavelength.

As mentioned by Scott (2001), CPO shows three absorption maxima similar to pure β-carotene at 425, 450 and 478 nm (Figure 2). Palm olein does not show any absorption peaks within 400-500 nm range which proves that β-carotene level in palm olein is less than detectable limit (Top et al. 2011).

The highest level of carotenoids was accumulated in fish fed with CPO. The skin colour of fish fed with CPO remained largely unchanged although a slight pinkish colour could be observed in their skin. The highest triacylglycerol was obtained in fish fed with CPO (table 3).

Previous studies had been demonstrated that the dominant carotenoids in the flesh of Nile tilapia were astaxanthin and canthaxanthin. Violaxanthin was the second most important pigment identified in Nile tilapia skin followed by zeaxanthin (Seef et al. 2014). Several studies have shown that dietary supply increases the integument concentration of carotenoids in cichlids (Brown et al. 2013; Güroy et al. 2007, 2012; Kaisuyama et al. 1988; Kop et al. 2008).

Most information on carotenoid metabolism in the cichlid integument comes from feeding experiments with Nile tilapia (Kaisuyama et al. 1988). The reconstructed metabolic pathways involve epimerization, reduction or oxidation of dietary canthaxanthin, astaxanthin, zeaxanthin and lutein. Dietary tunaxanthin accumulated in the integument without chemical modifications, whereas dietary β-carotene was neither accumulated nor bio converted in the integument (Kaisuyama et al. 1988). Consistent with the findings in tilapia, dietary β-carotene had smaller effects on skin coloration than astaxanthin. Similar Feeding experiments of with two cichlids Amphilophus citrinellus (Pan and Chien 2009) and Cichlasoma severum (Kop et al. 2008) had proved it.

The carotenoid deposition in flesh was significant over the skin of Nile tilapia. The carotenoids deposition was not evenly distributed in CPO fed fish fillets. As ventral part is thinner, it showed higher deposition than the dorsal part of the flesh. Reddish colour was observed closer to the caudal area than the anterior part due to the excess deposition of carotenoid pigment in that area. Previous studies by No and Storebakken (1991) have shown that caudal part of fish may contain 30-40% more carotenoids than the back and neck part of the fillet. Bjerkeng (1992, 2000) observed a longitudinal variation in carotenoid content and red colour of the fish flesh, which indicated that higher carotenoid deposition occurred in the caudal area than in anterior part of the body of salmonid fishes. This can occur due to the rapid deposition of carotenoids close to the backbone (Weerakkody and Cumaranathunga 2016). Fish fed with control diet containing palm olein also showed a slight carotenoid deposition in flesh. The carotenoids (specially carotene) other than β-carotene present in the palm olein might be the reason for giving a slight reddish colour to fillet in control treatment. It proved that β-carotene was not the only pigment responsible for flesh colouration in the present study. But carotenoids which are absent in palm olein (mainly β-carotene) might give more colouration in fish fed with CPO diet.

The sensory analysis component of the present study also indicates that sensory properties of fillet colour help to increase consumer attraction. A red colour fillet is more attractive and has a higher consumer preference in our sample population. Weerakkody and Cumaranathunga (2016) have conducted a study on carp fed with diets incorporating diets, and have revealed that there was a significantly higher carotenoid accumulation in flesh, as well as in skin.

Red porgy (Pagrus pagrus) fed with an astaxanthin-supplemented diet, β-carotene and lycopene supplemented diets led an increased total carotenoid content in the dorsal skin area (Chatzifotis et al. 2005).

Arous et al. (2014) reported that feed supplemented with β-carotene riched Dunaliella salina extract had given a reddish colour in red tilapia flesh. Similar studies were carried out by
Valente et al. (2016) showed that the total carotenoid concentration was the highest in the skin of Nile tilapia fed with 5% *Ulva* spp. followed by 10% *Ulva* spp. In muscle samples, no carotenoids could be detected.

**CONCLUSIONS**

Total carotenoid levels of CPO is found about 13.627 µg/g. A considerable increase of carotenoids in fish flesh was detected compared to the initial carotenoids content in flesh. Although skin color was not affected by β-carotene, flesh was affected significantly. The tested carotenoid sources of CPO can be used in the aquaculture industry to increase the consumer preference and attraction for fish fillets. CPO can be used as natural plant based carotenoids to improve the colour of fish flesh.

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