Effect of different feed ingredients on digestive enzymes activity and on the histology of liver and intestine in *Labeo rohita* Hamilton, 1822

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ABSTRACT

Present study compared the effect of fishmeal and selected plant-origin feed ingredients on digestive enzymes activity as well as on the histology of liver and intestine in *Labeo rohita*. Guar meal (GM), soybean meal (SBM), cotton seed meal (CSM) and canola meal (CM) were used as experimental feed ingredients in the four treatment diets, while fishmeal (FM) incorporated diet served as control. There were three replicates for each of the treatment and control diets. The experiment was conducted in 15 fiber glass tanks, each having 10 fish. Fish in each tank was fed @ 4% of body weight. Protease activity varied significantly (p<0.05) between anterior and posterior part of the intestine while amylase and lipase activities of whole intestine differed significantly (p<0.05) between the treatments. Maximum lipase activity was observed in fish fed with GM while minimum with CM ingredients. Vacuolation of hepatocytes and some variations in the mid intestine were also observed in response to different feed ingredients. Protease activity was greater throughout the intestinal tract when fish were fed with CSM and FM diets. Results of the present study indicated that CSM and FM are better ingredients for feed formulation for *L. rohita*.

Keywords: Amylase, Cotton seed meal, Digestive enzymes, Fishmeal, Histology, Intestine, Lipase, Liver, Protease

Introduction

In developing countries, fish is one of the best sources of animal protein as it provides about 26% of the daily protein requirement (Delgado et al., 2002; Louka et al., 2004). *Labeo rohita* Hamilton, 1822 is one of the prime freshwater fish species, commercially important in Asia in general and in Indian subcontinent in particular (Khan et al., 2004). In aquaculture, nutrition is the basic field where lots of work have been undertaken and there is further scope for future research (Eroldogan et al., 2004; Gomez-Requeni et al., 2004; Eroldogou et al., 2006a, b) because regional differences in the ingredients make diet formulations difficult. Better feed consumption improves fish growth as it implies better enzymatic activities which is greatly influenced by the feeding habits of fishes (Smith, 1980; Lundstedt et al., 2002). For understanding feed digestion mechanism and adaptation to the change in nutritional environment in fish, it is necessary to study the digestive enzymes and their activities in different parts of the intestine (Francis et al., 2001; Sunde et al., 2004; Romarheim et al., 2007; Santigosa et al., 2008). Study of the digestive enzymes is a reliable tool to understand digestive processes and fish nutritional status (Harms et al., 1991; Johnston et al., 2004). Proteolytic and amylase enzymes’ activities can unveil the ability of different fish species to use protein and carbohydrates (Hidalgo et al., 1999). Lipases are also inducible enzymes (Aliyu-Paiko et al., 2010) and can be stimulated by the dietary lipid content (Buchet et al., 2000; Li et al., 2012). A positive correlation between lipase activity and dietary lipid content in juvenile rice field eel (*Monopterus albus*) has been documented (Ma, 2014). In fish it will be easy to understand the nutrient digestibility after understanding enzymatic activity (Kolkovski, 2001) and it will facilitate better feed formulations (El-Sayed et al., 2000; Eusebio and Coloso, 2002). The histological changes in liver and intestine are also associated with nature of feed ingredients.
(Caballero et al., 2004; Uran et al., 2009; Poleksic et al., 2007). In fish, replacement of fishmeal (FM) with plant origin feed proteins causes reduced growth rates and pathological changes, mainly enteritis in the distal part of the intestine (Uran et al., 2009). Present experiment was therefore planned to understand the effect of different plant origin feed ingredients in comparison with fish meal, on digestive enzymes activity as well as on the histological features of liver and intestine in juvenile *L. rohita*.

**Materials and methods**

This study was conducted in the Department of Fisheries and Aquaculture, University of Veterinary and Animal Sciences, Ravi Campus Pattoki, Lahore, Pakistan. Experiments were conducted in fiberglass tanks (12.5×4.5×3.0 feet, length×width×depth) using juvenile *Labeo rohita*.

**Experimental design**

Experiments were designed following completely randomised design (CRD). Fifteen uniform-sized fiber glass tanks were stocked with 10 nos. each of juvenile *L. rohita* having body weight ranging from 80 to 120 g. Four experimental diets incorporated with plant origin feed ingredients viz., soybean meal (SBM), cotton seed meal (CSM), canola meal (CM) and guar meal (GM) were formulated along with a control diet containing fish meal (FM). Fish were regularly fed with the experimental diets in three replicates tanks each, @ 4% of wet body weight, twice a day for three months (90 days).

**Proximate analysis**

Feed ingredients were analysed for fat, moisture, protein, fiber, ash and phosphorus by Buchi NIR Technology (Buchi NIRFlex N-500) (Table I) (Martinez et al., 2003; Iqbal et al., 2015). The ingredients were dried, ground and were placed in sampler cups. The cups were placed in NIR machine for 2 min which then displayed a complete proximate analysis report.

**Mineral analysis**

Well ground sample (0.5 g) was taken in a conical flask, 10 ml of HNO₃ was added, the mixture was then boiled for 15 min at 60°C and then 5 ml perchloric acid was added. It was boiled again for another 15 min at 60°C. The flask was then placed on a hot plate and heated until sample volume reduced to 1 ml. This sample was diluted to 100 ml by addition of distilled water. Sodium (Na) and potassium (K) were measured through flame photometric method while calcium (Ca), iron (Fe), zinc (Zn), copper (Cu) and magnesium (Mg) were determined in an atomic absorption spectrophotometer (AAS) (Iqbal et al., 2014).

**Sample preparation for intestinal enzymes**

Six fish samples each from experiment and control diets (three for whole intestine and three each for anterior and posterior portions of the intestine) were collected. These samples were degutted, removed intestine and homogenised in a homogeniser by adding chilled Tris-HCl. The homogenate was centrifuged at 6000 g for 15 min and the supernatant was collected and stored at -20°C (Ismat et al., 2013) until analyses of intestinal enzymes activity (amylase, protease and lipase).

**Amylase activity**

To estimate amylase activity, the sample homogenate along with 1% starch solution and phosphate buffer (pH 6.9) were added in the sample test tubes in the ratio
of 1:1:1. Starch solution (1 ml) was added in the blank test tube and both sample and blank tubes were incubated at 37°C for 15 min. One milliliter of glucose solution (0.1%) was taken as standard in a test tube. Subsequently, 1 ml DNS (3,5-dinitrosalicylic acid) reagent was added to the sample, blank and standard test tubes and all the test tubes were kept in boiling water bath for 1 min. These test tubes were then cooled at room temperature and then 2 ml distilled water was added in standard and blank test tubes. Absorbance of blank, standard and samples were measured in a spectrophotometer at 540 nm and activity of amylase (units ml⁻¹ min⁻¹) was estimated as per Ismat et al. (2013).

**Protease activity**

Protease enzyme activity was assessed using 1% azocasein in 50 mm Tris–HCl having pH 7.5 (Garcia-Carreno 1992). Ten microliter of homogenate was mixed with 0.5 ml of phosphate buffer (pH 7.5), 0.5 ml of substrate solution and incubated for 10 min at 37°C. The reaction was stopped by adding 0.5 ml of 20% trichloro-acetic acid (TCA) and then centrifuged for 5 min at 14,000 g. Absorbance of the supernatant was measured at 366 nm. Standard curve was prepared using different concentrations (0, 2, 4, 5, 6, 8 and 10 mg) of azocasein (Ismat et al., 2013).

**Lipase activity**

One milliliter of sample was stirred in the presence of 3.5 ml phosphate buffer (pH 7.5) and 0.5 ml olive oil for 30 min at 37°C. Enzyme activity was stopped by adding 1 ml of acetic acid and then 3-4 drops of phenolphthalein indicator was added in the mixture. Then mixture was titrated against NaOH (10 mm) solution till the colour became pink (Ismat et al., 2013).

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\text{Lipase activity (units ml}^{-1}\text{min}^{-1}) = \frac{\text{Vol. of NaOH} \times \text{Normality of NaOH} \times 1000 \times 40 \times 1000}{\text{Vol. of sample homogenate used} \times 1000 \times \text{mol. wt. of oleic acid} \times \text{time (min) of incubation}}
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**Histology**

On termination of the experimental trial, liver and mid intestine samples were collected from three fish per treatment for histological analyses. These samples were immediately fixed in 10% neutral buffered formalin and processed following standard histological procedures i.e., dehydrated in ascending series of ethanol concentrations, and embedded in paraffin. Tissue sections were cut at 4-5 μm, stained with hematoxylin and eosin (Humason, 1979; Markovic et al., 2012).

**Physico-chemical parameters of water**

The water quality parameters viz., dissolved oxygen (DO) was determined using DO meter (YSI 55 Incorporated, Yellow Springs, Ohio, 4387, USA), pH using pH meter (LT-Lutron pH-207 Taiwan) while electrical conductivity, water temperature, salinity and total dissolved solids (TDS) were determined using conductivity meter (Condi 330i WTW 82362 Weilheim, Germany) following APHA (1998).

The data obtained were subjected to statistical analyses using the software package SAS 9.1 and analysis of variance (ANOVA) was applied to compare means.

**Results and discussions**

Water quality parameters (temperature, dissolved oxygen, pH, TDS, salinity and electrical conductivity) remained within the permissible limits and corroborated with previous findings (Ali et al., 2000; Abid and Ahmed, 2009).

During the present study, maximum protease activity was observed in fish fed with CSM while CM fed fish showed maximum amylase activity. Fernandez et al. (2001) stated that the constituents of feed work as driving force for digestive enzymes in fish. Differences (p≤0.05) in protease activity were observed in various parts of the intestine and that of the whole intestine. Amylase activity also showed significant differences in whole intestine of fish fed on different feed ingredients (Table 2). Kumar et al. (2011) documented significant differences in amylase and protease activity in *Labeo rohita* in response to different feed ingredients. Similar effects were observed in *Pagrus pagrus* (Suzer et al., 2007) and in juvenile Senegalese sole (Rodiles et al., 2012) when fishmeal was partially replaced by plant protein sources. Findings of all these investigations are in line with our findings and confirm that variations in dietary quality as well quantity do have lot of bearings on enzymatic activities of digestive tract.

Due to physiological versatility in *L. rohita* fingerlings, higher lipase secretions were observed for lipid digestion (Sethuramalingam and Haniffa, 2002). During the present study, lipase activity in anterior, posterior and whole intestine was found variable (p<0.05) for all the five feed ingredients (Table 3). Lipases are inducible enzymes which could be stimulated by the dietary lipid content (Aliyu-Paiko et al., 2010; Li et al., 2012). Maximum
lipase activity was observed in the whole intestine when fish were fed with GM diets while minimum activity was observed in anterior part of intestine for CM diets. Ma (2014) reported positive correlation between lipase activity and dietary lipid content in fish. The variation in feed composition causes changes in the specific activity of digestive enzymes in *P. pagrus* (Suzer et al., 2007). Ismat et al. (2013) observed highest lipase activity in *Catla catla* fed with SBM diets whereas *Hypophthalmichthys molitrix* showed poor performance. Fernandez et al. (2001) pointed out that digestive adaptation in different species exhibit closer correlation with their diet rather than on their taxonomic category. Similarly, Kuzmina (1996) also stated that changes in digestive enzyme activity could be affected by feeding behaviour and biochemical composition of feed.

In the present study, histological changes were also observed in the liver of fish fed with different feed ingredients. Severe vacuolation of hepatocytes and mild congestion were seen in sinusoids in fish liver fed with CM diet (Fig. 1). Robaina et al. (1995) observed hepatocytes vacuolisation and disorganisation in the liver of the fish fed on 30% SBM diet. Similarly, unstained portions showing fat vacuoles were observed in hepatocytes of fish fed with CSM diet (Fig. 2). Demska-Zakes et al. (2012) reported congestion, vacuolisation in hepatocytes, nuclear chromatin and nucleus disintegration when supplementary feed containing vegetable oil was offered to juvenile tench (*Tinca tinca*). Vacuolation in hepatocytes with pyknotic nuclei were observed in fish fed on FM diet.
Effect of different feed ingredients on digestive enzymes activity in *Labeo rohita* (Fig. 3). GM feed caused severe cytoplasmic vacuolation (Fig. 4) while SBM caused hepatocytes degeneration and mild vacuolation (Fig. 5). Bac *et al.* (1983) and Parpoura and Alexis (2001) observed necrosis and degeneration of hepatocytes cell membranes in European seabass and gilthead seabream (*Sparus aurata* L.) receiving feeds supplemented with soy oil. Similarly, common carp fed with 50% mustard protein showed histological abnormalities in liver tissues (Hossain and Jauncey, 1989).

Several researchers have observed changes in intestine of fish fed with plant origin feed which suppressed fish growth (Van den Ingh *et al.*, 1991; Baeverfjord and Krogdahl, 1996; Refstie *et al.*, 2000; Krogdahl *et al.*, 2003; Knudsen *et al.*, 2008). During the present study mild infiltration of leucocytes in lamina propria and in lumen as well as sloughed tissue debris was seen in mid intestine of *L. rohita* fed with GM diet (Fig. 6). Similarly, SBM diet caused degenerative changes on surface epithelial cells (Fig. 7). In common carp (*Cyprinus carpio*), signs of enteritis were observed when fed with high levels of SBM (Uran *et al.*, 2008) while Bonaldo *et al.* (2006) observed no pathological changes in intestines of *Solea aegyptiaca* when 30% FM was
replaced with SBM. FM diet caused detachment of surface epithelial cells from underlying lamina propria indicating necrotic changes (Fig. 8). Kumar et al. (2010) observed necrosis of enterocytes and denudations of the upper laminar epithelium, accompanied by a massive influx of leucocytes into the lamina propria of the villi when 75% of FM protein was replaced by detoxified Jatropha kernel meal. CSM diet resulted in neutrophillic infiltration in lamina propria with segmented nucleus and there was also goblet cell hyperplasia in mucosa (Fig. 9). Edematous fluid along with leucocytic infiltration in the lamina propria and small blood vessels were observed in intestine when L. rohita was fed with CM diet (Fig. 10). Markovic et al. (2012) observed increased leucocytes infiltration in the epithelium often accompanied by increased mucous production when carp was fed with plant origin feed. Baeverfjord and Krogdahl (1996) and Uran et al. (2008) reported enteritis which involved inflammatory infiltrate in fish intestine fed with soybean meal.

Based on the results of the present study, it can be concluded that every ingredient behaves differently in the intestine due to differences in nature of nutrients. Better protease level was observed in the whole intestine when fish were fed with CSM and FM which ultimately enhanced fish growth.

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K. Javed Iqbal et al.

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