Analysis of Enzyme Activities of the Gut Bacterial Communities in *Labeo rohita* fed Differentially Treated Animal Fleshing Diets

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Abstract

The present investigation includes isolation, enumeration and comparison of the gut microbial flora of *Labeo rohita* fed differentially treated Animal Flesming (ANFL) as a sole protein source in their diets and related increase in enzyme secretion. ANFL is the proteinaceous tannery solid waste generated during leather processing and inclusion of ANFL in aqua feed will pave way for the novel alternative cheaper protein source in replacement of fish meal. Six experimental diets were formulated with differentially processed ANFL. The isolates from the gut extract were quantitatively screened and quantitatively assayed for amylase, cellulase, lipase and protease activities. The total culturable bacterial count (10 × 10^7 CFU/g) as well as the proteolytic bacterial count (27 × 10^6 CFU/g) was high in the fish gut fed with diet 5 containing the fermented ANFL. Almost all the bacterial isolates possesses protease activity in which strains (FF5, CF3and CF4) isolated from the diet 5 (287U) and 7 (282U) exhibited highest activities. Maximum cellulase (FF2), amylase and lipase (RF6) activity were observed in the strains isolated from the diets 5 and 1 (78U, 186U and 97U). This study proves the existence of diet dependent enzyme producing bacterial community in the gut of *Labeo rohita* through SEM analysis and the information generated from the present study might contribute towards utilization of enzyme producing bacterial isolates as probiotic and in better feed formulations at low cost for rohu incorporating the tannery solid waste ANFL as a protein source.

Keywords: Gut microbial enzymes; Animal Flesming (ANFL); *Labeo rohita*

Abbreviations: ANFL - Animal fleshing, SMA - skim milk agar, CMC - Carboxy methyl cellulose agar, SA - Starch agar, TA - Tween 20 agar

Introduction

The micro-environment of the digestive tract of fish confers a favorable growth for the microorganisms [1-3] by providing ecological niche for these organisms [4]. The microbiota of fish has been shown to be highly dependent on the bacterial colonization during early development, environmental conditions, nutrient composition and dietary changes [5,6,62]. These microorganisms grow upon the food absorbed by the host animal, digestive secrections and fragments scaled off the mucosal epithelium [7] and have a better chance of survival [8].

The indigenous microflora of fish in aquaculture has previously been studied for several purposes [19,21,22] and differs from ovary skin and liver [71]. Previous studies include descriptions of microbial spoilage [9,72], relationship between environment and fish microflora [10], monitoring of changes in fish farms [11], nutritional role of the intestinal flora [12] and the antibiotic resistance profile of the indigenous flora [13]. Preceding research suggests that digestive enzymes present in fish digestive tract can elucidate some aspects of their nutritive physiology [73] and thus be supportive to develop nutritional strategies for fish feeding and diet formulation [14-17].

In general, the bacterial flora of the gastrointestinal tract with diversified enzymatic potential plays a vital role in major part of the metabolism of the host animal [18]. From the studies made so far, it appears that different species of fish and crustaceans have a specific resident gut microbiota [60,62,85] and the isolated species include *Acinetobacter* spp., *Enterobacter* spp. and *Pseudomonas* spp. in trout (*Onchorhyncus* sp.); *Aeromonas* spp., *Flavobacterium* spp. and *Lactobacillus* spp. in Arctic char (*Salvelinus alpinus*) [58], *Enterobrevibrio* spp. from the intestinal tract of turbot (*Scophthalmus maximus*)[59]. *Vibrio* spp., *Bacillus* sp., *Pseudomonas* sp., *Photobacterium* sp. and *Plesionomonas* sp. in white shrimp, *Fenneropenaeus* (Pentaeus) indicus and *P. glacincola* in the hind-gut of Arctic char [6], and Atlantic cod [22]. *Vibrio* species from gut of halibut larvae [69] and Atlantic cod [74,22]. *P. phosphoreum* farmed Norwegian salmon [65], and UK halibut [60]. *Providencia* sp. and *Shewanella* sp.[63]. However, the information regarding the enzyme producing intestinal bacteria related to feed in fish is scarce [16].

The inter-relationships between the microbiota and the host are clearly important as the health and imbalance between these systems appear to drive a wide range of mucosal and systemic immune-mediated disorder [26,27]. Several authors revealed the presence of microflora in fish gut [19-22] and enzyme producing bacteria in the digestive tract of *L. rohita* [23,24]. The use of such beneficial bacteria has a long tradition in the animal husbandry [25]. However, the research based on gut microbial population in fish species and digestive enzymes related to animal fleshing feed source, their significance as a means to determine assimilating capacity is perhaps nil.

Animal protein sources present good essential amino acids balance [76], but costs and heterogeneous composition repeatedly limit their use, and the use of alternative sources is required [77]. Hence, amino acid rich proteinaceous tannery solid waste animal fleshing was used as a protein source in fish feed formulation. India is one of the leading
countries in freshwater aquaculture. In 1998, the total production of freshwater fish was 1.7 million tons of which carp contributed about 87%. Rohu (Labeo rohita) is the most important Indian major carp that enjoys high consumer preference in many states with high protein conversion ratio [75]. Hence, the focal theme of the present work was to evaluate the enzyme producing microbiota and activities of enzymes in the digestive tract of rohu, Labeo rohita fed differentially treated animal fleshing incorporated diets.

Materials and Methods

**Acclimatization of fish to experimental conditions**

Labeo rohita fingerlings (average weight 5g ±0.25g) were obtained from nearest farm in Chennai (India) and acclimatized in 1000 L cylindrical tanks filled with dechlorinated tap water for 15 days with control feed. Three hundred and fifty fingerlings were selected for the study and divided into six experimental groups and one control. Each group of 50 fingerlings was again divided into two equal duplicate subgroups. Fish were fed twice daily (9:00 a.m. and 17:30 p.m.) to apparent satiation. The fish were fed 3% of their body weight twice a day. The limed animal fleshing was collected from a tannery processing raw goat skins/cow hides into leather in Chennai. Briefly, the animal fleshing was subjected to various optimized treatments (data not shown) such as microwave for 60 minutes, ozonation for 60 minutes, soaking in H2O2 for 60 minutes, fermentation with bacterial mixed culture from the fish gut and autoclaving (Table 1). The impact of nutritional effects on gut flora and gut enzyme was studied on 60th day.

**Microbiological Examination of Gut Flora**

Fish from each experimental set was collected, after 36 hrs of starvation. The fish was sacrificed and surface sterilized using 1% sodium hypochlorite. The fish was stored in 10 ml chilled buffered saline solution. Diluted samples (0.1 ml) with sterile chilled buffered saline solution. Diluted samples (0.1 ml) were spread aseptically within a laminar air flow on sterilized Schaedler HiVegTM Agar (Hi media, India) to determine the total culturable gut heterotrophic bacterial population. Spread plate technique was used to isolate and enumerate protease, cellulase, amylase and lipase producing bacterial population, diluted samples (0.1 ml) was poured on skim milk agar (SMA), Carboxy methyl cellulose agar (CMC), Starch agar (SA) and Tween20 agar (TA) plates, respectively. Culture plates were incubated at 37°C for 48 hours and were subsequently examined for the development of bacterial colonies. It was assumed that the micro flora, which had formed colonies on the SA plate, had amylolytic activity, CMC- cellulolytic, SMA -proteolytic and TA plates- lipolytic activities. Colony numbers per unit sample volume of gut homogenate were determined by multiplying the number of colonies formed on each plate by the reciprocal of dilution. The well separated colonies producing various extracelluluar enzymes with apparently different morphology were streaked separately on Tryptone Soy Agar (TSA) (Hi Media, India) plates to obtain pure cultures [74]. Single, isolated colonies from the streaked plates were transferred to TSA slants for further study. Chemicals used for analysis were procured from Merck and Hi Media, India. No conflict of interest existed during the entire study period.

**Qualitative assay for enumeration of enzyme producing bacteria**

The intensity of extracellular enzyme production by the pure cultures was determined on agar plates with selective media. For extracellular amylase production, the samples were inoculated on SA plates and incubated at 37°C for 48 h. The culture plates were then flooded with 1% Lugol’s iodine solution [56]. Formation of transparent zone surrounding the colony indicated amylase activity. Similarly, for extra-cellular protease, the samples were inoculated on SMA plates and incubated at 37°C for 24 h. The appearance of a clear zone around the colony confirmed the proteolytic activity. For determination of cellulase production, the samples were grown on CMC plates at 37°C for 24 h and flooded with 0.1% Congo red dye and destained with 1M sodium chloride [57]. Congo red selectively binds with unhydrolyzed CMC. Appearance of clear halo due to the presence of hydrolyzed CMC surrounding bacterial colony indicated cellulase production in the medium. For assaying lipase activity the samples were inoculated on the TA plates. Formation of calcium laurate white crystals due to the reaction between the fatty acids released and CaCl2 present in the medium around the colonies confirm the lipase activity of the colony.

**Media composition**

Schaedler HiVegTM Agar (g L⁻¹): Amino acid, 1.00; Lactose, 5.00; Skim milk agar (SA), Carboxy methyl cellulose agar (CMC), Starch agar (SA) and Tween20 agar (TA) plates, respectively. Culture plates were incubated at 37°C for 48 hours and were subsequently examined for the development of bacterial colonies. It was assumed that the micro flora, which had formed colonies on the SA plate, had amylolytic activity, CMC- cellulolytic, SMA -proteolytic and TA plates- lipolytic activities. Colony numbers per unit sample volume of gut homogenate were determined by multiplying the number of colonies formed on each plate by the reciprocal of dilution. The well separated colonies producing various extracelluluar enzymes with apparently different morphology were streaked separately on Tryptone Soy Agar (TSA) (Hi Media, India) plates to obtain pure cultures [74]. Single, isolated colonies from the streaked plates were transferred to TSA slants for further study. Chemicals used for analysis were procured from Merck and Hi Media, India. No conflict of interest existed during the entire study period.

**Quantitative enzyme assay of gut and the bacterial isolates**

Fish gut from each feed trial was aseptically removed on 60th day after 36 h starvation for quantitative assay of cellulase, amylase, and protease and lipase activity of gut enzyme. The gut was homogenized in 10 ml chilled buffered saline to make a suspension and then centrifuged at 4500 rpm for 15 min. The clear supernatant was used

<table>
<thead>
<tr>
<th>Experimental diets</th>
<th>Rice Bran (g)</th>
<th>Wheat Flour (g)</th>
<th>Oil Cake (g)</th>
<th>Sunflower Oil (g)</th>
<th>Protein Source (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DIET 1</td>
<td>20</td>
<td>20</td>
<td>10</td>
<td>1</td>
<td>Raw ANFL (rf)</td>
</tr>
<tr>
<td>DIET 2</td>
<td>20</td>
<td>20</td>
<td>10</td>
<td>1</td>
<td>Microwaved treated ANFL (mf)</td>
</tr>
<tr>
<td>DIET 3</td>
<td>20</td>
<td>20</td>
<td>10</td>
<td>1</td>
<td>Ozone Treated ANFL</td>
</tr>
<tr>
<td>DIET 4</td>
<td>20</td>
<td>20</td>
<td>10</td>
<td>1</td>
<td>H2O2 treated ANFL (hf)</td>
</tr>
<tr>
<td>DIET 5</td>
<td>20</td>
<td>20</td>
<td>10</td>
<td>1</td>
<td>Fermented ANFL (ff)</td>
</tr>
<tr>
<td>DIET 6</td>
<td>20</td>
<td>20</td>
<td>10</td>
<td>1</td>
<td>Autoclave ANFL (af)</td>
</tr>
<tr>
<td>CONTROL</td>
<td>20</td>
<td>20</td>
<td>10</td>
<td>1</td>
<td>Fish meal (cf)</td>
</tr>
</tbody>
</table>

*Vitamin premix mg L⁻¹: retinol palmitate 400,000IU, thiamine 5, riboflavin 4, niacin 20, folic acid 2, pyridoxine 6, cyanocobalmine 5, ascorbic acid 12, cholecalciferol 55,000IU, tocoferol 3, biotin 0.3, choline chloride 90, pantothenic acid 10. Mineral premix for 100g calcium phosphate dibasic 7.35; citric acid 0.03; ferrous sulphate 50 mg, magnesium oxide 3, manganese sulphate 0.7; potassium phosphate dibasic 9; sodium chloride

**Table 1:** Aqua feed composition.
for assaying various enzyme activities. Selective broth media were used as production media for a quantitative assay of amylase, cellulase, and protease and lipase production of selected bacterial isolates. A loopful of selected strain was inoculated into Tryptone soya broth and incubated for 24 h at 37°C. The liquid production medium about 25 mL was inoculated with 2% of the inoculums obtained from seed culture. The culture flasks were incubated for 48 h at 37 ± 1°C. After incubation, the contents were centrifuged at 4500 g for 15 min, at 4°C and the cell-free supernatant was used for enzyme assay. Ten fish were harvested from each experimental setup for further analysis.

### Amylase assay

Amylase activity was assayed by the dinitrosalicylic acid (DNSA) method based on the estimation of reducing sugars at 540 nm using maltose as the standard (Bernfeld, 1955). 1% of starch in sodium phosphate buffer (0.1M, pH 7.0) was used as the substrate for the assay. One amylase unit was defined as the amount of enzyme per milliliter of sample that releasing one micromgram (µg) reducing sugar (maltose) per minute.

### Cellulase assay

Cellulase activity was measured according to the method of Denison and Kohen (1977) using 1% CMC in potassium phosphate buffer (0.1M, pH 7.0) as substrate. The reducing sugar production was measured at 540 nm by dinitrosalicylic acid method using glucose as the standard. One cellulase unit was defined as the amount of enzyme per milliliter of sample that released one micromgram (µg) reducing sugar (maltose) per minute.

### Protease assay

Protease activity was determined according to Kunitz caseinase assay method using 1% Casein in sodium phosphate buffer (0.1M, pH 7.0) as substrate. One unit of enzyme activity represents the amount of enzyme required to liberate 1 µg of tyrosine per milliliter sample under standard assay conditions.

### Lipase assay

Lipase assay was carried out based on titrimetric method using olive oil as a substrate and polyvinyl alcohol (PVA) as an emulsifier.

### Scanning Electron Microscopy (SEM)

The gut sections designated for SEM analyses were processed according to Ghosh et al. [29]. dehydrated with a graded ethanol series and subsequently freeze dried. The dried cells were sputtered coated with gold, and finally photographed by a scanning device attached to a JEOL JM – 5600 electron microscope at 20 kV accelerating voltage for an electron beam of wavelength 5–6 nm.

### Statistical Analysis

Statistical analysis of the experimental data was made by analysis of variance (ANOVA) followed by Scheffe’s F-test for multiple comparison [30].

### Results

Analysis of the bacterial flora in the gastrointestinal tract of the experimental fish showed that the total culturable aerobic gut bacterial population on Schaedler HiVeg TM Agar plate were in the order of fermented ANFL (100000x10³ CFU/g gut tissue) > control > microwave treated > autoclaved ANFL > raw ANFL. The total aerobic bacterial population was found to be least in ozone treated (120x10³ CFU/g gut tissue) and H₂O₂ treated ANFL (15x10³ CFU/g gut tissue) (Table 2).

While enumerating specific enzyme producing bacterial flora, it was observed that the amylolytic strains were highest in microwave treated (4200x10³ CFU/g gut tissue) and fermented ANFL (3300x10³ CFU/g gut tissue) incorporated diets, followed by raw autoclaved > ozone > H₂O₂ treated ANFL incorporated diets. The cellulolytic population exhibited maximum activity in fermented (48000x10³ CFU/g gut tissue) and autoclaved ANFL (30000x10³ CFU/g gut tissue) followed by microwave > raw > ozone > H₂O₂ treated ANFL. Lipolytic bacterial flora were detected in the experimental fish and the maximum population density was recorded in fermented (3700x10³ CFU/g gut tissue) and autoclaved ANFL (3400x10³ CFU/g gut tissue) followed by raw > microwave treated > ozone > H₂O₂ treated ANFL. Proteolytic bacterial flora were found in all the experimental fish and the maximum count was observed in the gut of rohu (Labeo rohita) fed with fermented (27000x10³ CFU/g gut tissue) and microwave treated (24000x10³ CFU/g gut tissue) ANFL followed by raw, autoclaved > ozone > H₂O₂ treated ANFL. On comparison with control the protease producing bacterial population in the experimental feeds containing fermented and microwave treated ANFL, were quiet high. Lipase producing bacterial population was similar to control samples in fermented and autoclaved ANFL fed fish guts and low in all other diets (Table 3).

The intensity of extracellular enzyme activity of the intestinal extract of Labeo rohita gut from each experimental feed was assayed quantitatively and represented as shown in Figure 1. All the diets exhibited higher protease activity. Peak specific amylase and cellulase activities were exhibited by gut extracts of microwave and fermented

### Table 2: Aerobic bacterial count in the gut of rohu fed with experimental diet.

<table>
<thead>
<tr>
<th>Experimental feed (Protein Source) Diets</th>
<th>Total Bacterial Count (CFU × 10³/g gut tissue)</th>
<th>Cellulolytic Bacterial Count (CFU × 10³/g gut tissue)</th>
<th>Amylolytic Bacterial Count (CFU × 10³/g gut tissue)</th>
<th>Proteolytic Bacterial Count (CFU × 10³/g gut tissue)</th>
<th>Lipolytic Bacterial Count (CFU × 10³/g gut tissue)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Raw ANFL</td>
<td>2800000</td>
<td>15000</td>
<td>300</td>
<td>20000</td>
<td>230</td>
</tr>
<tr>
<td>2. Microwave ANFL</td>
<td>700000</td>
<td>12000</td>
<td>4200</td>
<td>24000</td>
<td>200</td>
</tr>
<tr>
<td>3. Ozonated ANFL</td>
<td>1200</td>
<td>120</td>
<td>10</td>
<td>10</td>
<td>70</td>
</tr>
<tr>
<td>4. H₂O₂ treated ANFL</td>
<td>15</td>
<td>13</td>
<td>15</td>
<td>15</td>
<td>10</td>
</tr>
<tr>
<td>5. Fermented ANFL</td>
<td>100000</td>
<td>48000</td>
<td>3300</td>
<td>27000</td>
<td>3700</td>
</tr>
<tr>
<td>6. Autoclave ANFL</td>
<td>70000</td>
<td>30000</td>
<td>2000</td>
<td>15000</td>
<td>3400</td>
</tr>
</tbody>
</table>

Initial (0th day) 40000 23000 1000 20000 4000

Results are means of triplicate treatments. Values with different superscript in the same column are significantly different.
ANFL fed fish than other diets. Specific activity of lipase was found to be highest in the fermented ANFL fed fish and maximum protease activity was observed in fermented ANFL and Microwave treated ANFL fed fish. The protease and lipase activity of the fermented ANFL fed rohu gut extract was similar to control.

The intensity of extracellular enzyme production by the bacterial strains isolated from the gut of rohu was assayed qualitatively (Table 3) in which each '+' indicates a zone diameter of 4mm (radius–2mm). Among these isolates, eight amylase, eight cellulase, nine lipase and fourteen protease producers were selected for quantitative enzyme assay. Peak cellulase (Figure 2) and amylase (Figure 3) activities were exhibited by the bacterial strains FF2 and RF6 isolated from experimental diet groups 1 and 5 respectively. Maximum protease (Figure 4) activity was observed in the strain FF5 isolated from experimental diet 5, followed by the strains CF3 and CF4 isolated from the control groups. The strain rf6 from experimental diet 1 showed highest lipase activity followed by ff5 and cf4 (Figure 5). In the present study, proteolytic bacteria were detected in the gut of all the fish examined and maximum density of proteolytic bacterial population was observed in the experimental diets 5 and 2 (2.7 x 10^7 and 2.4 x 10^7 CFU· g–1 intestinal tissue). However, assay of extracellular protease activity of the bacterial isolates showed highest value in FF5 (287 U), a strain isolated from the diet 5 fed fish gut. SEM analysis of Labeo rohita gut (Figure 6), the arrow marks indicates the position of the bacterial cells attached to the gut wall.

<table>
<thead>
<tr>
<th>Experimental diets</th>
<th>Bacterial strains</th>
<th>Amylase activity</th>
<th>Cellulase activity</th>
<th>Protease activity</th>
<th>Lipase activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Raw ANFL</td>
<td>RF3</td>
<td>++</td>
<td>+++</td>
<td>+++</td>
<td>+++</td>
</tr>
<tr>
<td></td>
<td>RF5</td>
<td>+++</td>
<td>+++</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td></td>
<td>RF6</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>2. Microwave treated ANFL</td>
<td>MF1</td>
<td>—</td>
<td>+++</td>
<td>+++</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>MF2</td>
<td>+++</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>MF3</td>
<td>+++</td>
<td>—</td>
<td>+</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>MF6</td>
<td>+++</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>3. Ozone treated ANFL</td>
<td>OF2</td>
<td>+</td>
<td>++</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td></td>
<td>OF5</td>
<td>+</td>
<td>+</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td>4. H2O2 treated ANFL</td>
<td>HF4</td>
<td>++</td>
<td>++</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td></td>
<td>HF5</td>
<td>—</td>
<td>+</td>
<td>+</td>
<td>—</td>
</tr>
<tr>
<td>5. Fermented ANFL</td>
<td>FF1</td>
<td>++</td>
<td>+++</td>
<td>+++</td>
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<td>FF5</td>
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<td>6. Autoclaved ANFL</td>
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<td>AF4</td>
<td>+</td>
<td>+++</td>
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<td>+</td>
</tr>
<tr>
<td></td>
<td>AF5</td>
<td>+</td>
<td>+++</td>
<td>+++</td>
<td>+</td>
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<tr>
<td>7. Control</td>
<td>CF2</td>
<td>++</td>
<td>++</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td></td>
<td>CF3</td>
<td>+</td>
<td>+</td>
<td>+++</td>
<td>+</td>
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<tr>
<td></td>
<td>CF4</td>
<td>+</td>
<td>+</td>
<td>+++</td>
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</tr>
</tbody>
</table>

'+-' sign indicates the intensity of enzyme production (zone diameter of 4mm). ++++, very high; +++, high; ++ moderate; +, low; —, nil.

Table 3: Qualitative extracellular enzyme activity of bacterial strains isolated from Labeo rohita fish gut fed with experimental diets.
Discussion

The factor contributing to the bacterial assemblage is diet [71]. Protein is the dietary macronutrient and energy source whose requirement is prioritized in nutritional studies, either because it represents the highest fish feed cost, or greatly affects specific growth rate [78,80]. Digestive enzyme activities of fish are associated with innate feeding habit and diet composition [81,82]. Hence, the study of digestive enzymes is an essential step towards understanding the mechanism of digestion and how organisms adapt to changes in the nutritional environment [83,84]. The proteinaceous ANFL generated from Leather industry after various treatments physical (autoclave and microwave), chemical (H₂O₂ and O₃) and microbial fermentation were included in diets of Labeo rohita to analyze the ANFL incorporated diet assimilating bacterial population with enzyme producing capacity. The processing techniques of ANFL includes heat treatment to obtain sterile product as defined by Codex Alimentarius (FAO) for the reduction in viable count in ANFL [31]. chemical process through ozonation and hydrogen peroxide treatment which are strong oxidizers that kills micro-organisms by blocking their enzyme control system and deodorizes both gaseous and particulate matter by an oxidative mechanism and removes the contaminants which can destroy the protein resulting in obtaining better level of protein [32-34] and microbial fermentation with mixed culture which is a best approach to obtain the desired product [35].

The gastrointestinal (GI) tract of fish is a complex ecosystem possessing a dense bacterial population levels consisting of aerobic, facultative anaerobic and obligate anaerobic bacteria [64-69] that can be defined as either autochthonous (indigenous) or allochthonous (transient) depending upon its ability to adhere and colonize the mucus layer in the digestive tract [36,37,70].

In the present investigation, the presence of a considerable population of bacterial flora in the gastrointestinal tracts of the fish species is confirmed through SEM analysis and supports the earlier reports that the bacteria entering along with the diet of fish during ingestion may adapt themselves in the gastrointestinal tract to form a symbiotic association and may be considered to have possible contribution in nutritional process of fish via extracellular bacterial enzymes. Moreover, certain strains of bacteria isolated from the gut extract exhibited quantitative amount of amylolytic, cellulolytic, lipolytic and proteolytic activities [15,38-41]. The Fermented ANFL increases protease producing bacterial population on gut and thereby aids in the digestion of proteinaceous diet. On the other hand, the H₂O₂ and Ozone (oxidizing and bactericidal agent) treated ANFL had the least bacterial count and low enzyme activity. The complexity and intact structure of raw ANFL minimizes the gut microbial protease activity of fish fed with diet 1 but cellulase and amylase activity were seen due to the presence of rice bran and wheat flour which are plant based natural
diet for the fish. In diet 2 the microwave heat treatment process would have un tangled the closely knit collagen structure present in ANFL thereby facilitating its easy uptake from the feed. In the diet 3 and 4, the native protein present in raw ANFL reduced assimilation; therefore, both the bacterial population and enzyme activity remained dormant (Table 2 and Figure 1). The gut extract of diet 5 fed fish showed maximum protease activity than control fish meal based feed. Elevated bacterial population with synchronized increase in protease activity of Labeo gut fed with fermented ANFL reveals the adaptability of rohu to new protein source. It is further noteworthy to state that diet 5 showed the maximum number of proteolytic bacterial count when compared to all other diet fed fish groups (Table 2). The probable reason could be the introduction of heavy bacterial population exogenously to the gut from the feed containing fermentation bacteria. The fermentation bacteria help the host in digestion of food and give less chance for the pathogens to colonize the gastrointestinal tract. Diet 6 fed fish groups also exhibits protease activity higher than diet 1, 3 and 4 but lower than the diet 5. Cellulase, amylase and lipase activities of the gut extracts in all the experimental fish were found to be more or less stable due to the similar feed components in the formulated diet.

Since the experimental diets contain carbohydrate sources such as wheat flour and rice bran considerable population of amylolytic bacterial strains were detected. The highest amylase producing strain RF6 (186 U) was isolated from the experimental diet 1. Similar results have been reported by previous researchers with carbohydrate based feed [24,42,43].

Microbial intestinal cellulase activity has been detected earlier in carp [44], grass carp [43], and in [38]. Fingerlings. Quantitative assay of the selected strains showed relative cellulase activity due to the presence of natural complex cellulose containing plant products like wheat flour and rice bran in the diet (Figure 2 and Table 3). Hence, the cellulase activity was low in the gut extract compared to that of the proteases due to the formulation of protein rich diets. However, the strains FF2 and FF4 were observed to have the highest activity among all other isolates for cellulase production. These bacterial strains present in the gut might indigenously posses the cellulolytic activity or may have derived due to ingestion of rice bran (plant source).

Although fish have an endogenous source of protease in their digestive tracts, not much attention has been paid to the microbial source of protease in fish. Ghosh et al.,[15] suggested that Bacillus circulans, B. pumilus, and B. cereus, isolated from the alimentary canal of Labeo rohita fingerlings were good producers of proteolytic enzymes, though they did not quantify the enzyme activity. Since the experimental diets were protein rich the bacterial strains as well as the gut extracts showed higher protease activity than other enzymes and this corroborates with the prior report wherein increased proteolytic enzyme activity in common carp were observed with bovine-trypsin based diet [45]. Kawai [86] reported adaptive changes in the proteolytic enzyme in common carp (Cyprinus carpio) in relation to the type of the diet. The occurrence of proteolytic bacteria in the gut of the isolated fish species seems to support the presence of a diet dependent microbial population [37]. Further these investigations have suggested that microorganisms have a beneficial effect in the digestive processes of fish. A number of studies on the use of proteases to improve the digestibility of feed ingredients in poultry, pigs and cattle have been published. In broiler chickens, the addition of protease to lupin-based [89] or soya bean-based [90, 91] diets resulted in improved digestibility of protein.

The micro flora of the fish gut is generally culturable and occurrence of protease, amylase and cellulase and lipase producing bacterial population is noteworthy in the digestive tract of rohu. Digestive enzyme activities has also been reported earlier using Bacillus sp. containing probiotics in common carp [46]. B. subtilis in four live-bearing ornamental fishes [47]. Lactobacillus spp. [48] in beluga (Huso huso) and Persian sturgeon (Acipenser persicus) [49] and live yeast in European sea bass larva [50]. Cellulolytic and Amylolylotic gut microflora has been reported earlier [4,38,40,92]. Protease producing organisms and their activity was observed in the gut of the six ANFL formulated diets fed groups and is highest in the fermented ANFL fed Labeo rohita which may be due to the presence of fermentation facilitating bacterial strains. Furne et al. [87] observed that it is possible to predict the ability of a species to utilize different nutrients based on its digestive enzyme profile. The bacterial diversity in the intestines of abalone fed with artificial food was higher than in individuals fed with sea algae [87] and differed between puffer fish fed a natural diet and those fed with an artificial diet [71,87]. The information generated from the present study might contribute towards the research of diet dependent change of micro biota in the gastrointestinal tract of Labeo rohita as well as the low cost feed formulation process utilizing the tannery solid waste as protein source. The study also indicates that there is a distinct microbial source of the digestive enzymes – amylase, cellulase, lipase.
and protease, apart from endogenous sources in fish gut through ANFL administered feed which helps the fish to digest the nutrients well and lead to increase in the total enzyme producing microbial population.

The information generated from the present investigation might contribute to the incorporation of these bacteria in commercial aquaculture as supplement in formulated fish feed or in form of bacteria biofilm to achieve colonization in the fish gut at a higher degree. However, further research involving potent bacterial strains should be conducted for evaluating their efficacy under actual farm conditions. The results are also in agreement with the previous research findings that type of food can influence the enzymatic activity [81,92], and the bacteria, initially introduced with food, further become adapted to particular gastro-intestinal environmental conditions, and develop as a distinct or transient community of facultative aerobic which, being metabolically active, play an active role in digestive processes [51–55]. However, the influence of these bacterial populations on the fish productivity is not known and requires further investigation.

In fish, data pertaining to digestive enzyme activity and profiles have helped overcome nutritional problems associated with formulation of artificial diets that best meet an animal’s nutritive capability [86]. The results also present a scope for fish nutritionists to utilize the enzyme-producing bacterial isolates as a probiotic in formulating cost-effective aquafeeds, especially for the larval stages when the enzyme system is not efficient. However, further investigations are required to know about the metabolic pathways used by these microorganisms in the alimentary tracts of fish, to explain the lifelong host-microbe homeostasis.

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