INTRODUCTION

After capture, fish is immediately degutted to avoid putrefaction. During gutting and processing of fish, a large amount of waste is produced in the form of scale, skin, fins and body viscera which is buried inland or disposed off in sea, resulting environmental pollution (Nagai; Suzuki, 2000). Effective use of this waste for extraction of some valuable products like collagen, gelatin and enzymes can reduce dispose off cost and generate revenue (TIDWELL; ALLAN, 2001). Another possible alternative method to get rid of this waste material is to convert it into animal feed after proper processing (KROGDAHL et al., 2005).

Protein is a major feed ingredient in aqua feeds and major source of this protein is fishmeal (TONGSIRI et al., 2010). With passage of time demand for fishmeal is increasing and its supply is being restrictive. This difference of demand and supply results in high cost and scarcity of fishmeal, efforts are turning to search alternate of fishmeal (LAPIE; BIGUERAS, 1992; NADEESHA et al., 2006). A successful substituent of fish meal is fish silage which can be prepared by fermentation processing of this waste body viscera (GERON et al., 2007).

Fish silage contains large amount of essential nutrients like protein and fats (CHAKRABARTI; SHARMA, 2005; BOLASINA et al., 2006). Fats of fish silage are usually categorized as saturated, unsaturated, mono unsaturated and poly unsaturated fatty acids. Polyunsaturated fatty acids (PUFA) are important component of fish feed and play a dynamic role in fish growth, and other animals including human beings (NETTLETON, 1991; KLOR et al., 1997; JABEEN; CHAUDHRY, 2011). Therefore, lipid profile analysis is an essential tool to determine and assess nutritional value of any diet. Since fats possess double amount of energy as compared to non-protein energy sources so utilization of fats spare protein for other constructive purposes in fish body (LEE; PUTNAM, 1973). Different researchers have used fish silage successfully in diet of different fish species e.g. Atlantic salmon (Salmo salar) (PARRISH et al., 1991), rainbow trout (Oncorhynchus mykiss) (AKSNES et al., 2006),
tilapia (*Oreochromis niloticus*) (FAGBENRO et al., 1994) and for carp larvae (*Cyprinus carpio*) (CARVALHO et al., 1997). They proved that digestibility of fish silage varies according to raw material used for silage preparation, type of experimental fish and enzymes concentration in fish digestive tract (SEENAPPA; DEVARAJ, 1995; STONE, 2003). These digestive enzymes along other bacteria and micro-organisms help in food digestion so they affect feed digestibility (GARCIA-CARREN; HAARD 1993; FURNE et al., 2008). Digestion of protein mainly depends upon ingredient composition and enzymes concentration in the digestive tract of fish (ALEXIS, 1990). The information obtained by enzyme study can help to design feed management and feed type (FERNANDEZ et al., 2011). In south Asia, and especially in Indian sub-continent, *Labeo rohita*, is most preferred fish species and cultured in ponds (MISRA; SAMANTARAY, 2004). Hence, the same fish species was selected as an experimental fish in the current project.

The aim of present research project was to analyze fatty acid profile of fish fermented silage and to check possible effects of silage incorporated diets on digestive enzyme activity in *Labeo rohita* intestine.

**MATERIAL AND METHODS**

Present experiment was performed in at The University of Lahore, Pakistan. Fish body viscera were collected from local fish market. Collected viscera were washed with distilled water twice to remove mud and other un-necessary material. After washing, collected viscera were minced with electric mincer and poured in plastic tubs having lids. Molasses 5% (as source of sugar), yogurt 5% (as source of starter culture), orange peels 30% (as filler) and already minced body viscera 60% were mixed and thoroughly stirred (FAGBENRO; JAUNCEY 1998). Whole mixture was stored at ambient temperature in an-aerobic conditions. Initially, the mixture was solid but with the passage of time lactic acid was produced by *Lactobacillus* bacteria its pH started to decrease and mixture started to liquefy. At the end of thirty days storage period, dark brown pasty type material with strong fishy smell was produced. At the top of fermented silage there was liquid lipid layer which was removed manually.

**Sample analysis for fatty acid analysis**

For lipid extraction Bligh; Dyer (1959) was followed. Boron rifluoride-methanol complex was used for esterification. The fatty acid methyl esters were used as standards for fatty acid comparisons. Fattori et al (1987) was followed for preparation of methyl esters. Reagent 2, 7-dichlorofluorescein was used for color spots of lipid under ultra violet light; $\lambda$ 366 nm. Fatty acid analysis was performed by gas-liquid chromatography (GLC) at Pakistan Council of Scientific and Industrial Research (PCSIR), Lahore. Fatty acid methyl esters (FAME) were separated identified and quantified using GLC. Helium gas was used as a carrier agent. Methyl esters were identified by their comparison with chromatograms of standards. Peaks were identified by their retention time (KIESSLING et al., 2001).

**Diets Preparation**

Already prepared fermented fish silage was oven dried at 80°C in an electric oven and grinded in fine powder form by grinding mill. Proximate analysis the silage was performed to check nutritional value of silage (AOAC, 1990). Four iso-nitrogenous diets were prepared by incorporation of fish silage at different ratios viz. 100%, 75%, and 50% along with other co-ingredients i.e. rice bran and soybean meal. These diets were designated as $T_1$, $T_2$, and $T_3$ while fourth control diet was prepared by inclusion of fish meal instead of fish silage and was termed as $T_0$ (Table 1).

**Table 1.** Percentages of various feed ingredients and protein levels used in four treatment diets

| Ingredients         | $T_1$      | $T_2$      | $T_3$      | $T_0$      |
|---------------------|------------|------------|------------|------------|
|                     | Weight (g) | Weight (g) | Weight (g) | Weight (g) |
| Fish meal           | -          | 100        | 75         | 50         |
| Fish silage         | 100        | 18         | 0.8        | 100        |
| Soybean meal        | 18         | 7          | 20         | 100        |
| Rice bran           | 32         | 21         | 30         | 14.15      |
| Total               | 100        | 30.18      | 30.2       | 31.33      |

Fattori et al (1987) was followed for preparation of methyl esters. Reagent 2, 7-dichlorofluorescein was used for color spots of lipid under ultra violet light; $\lambda$ 366 nm. Fatty acid analysis was performed by gas-liquid chromatography (GLC) at Pakistan Council of Scientific and Industrial Research (PCSIR), Lahore. Fatty acid methyl esters (FAME) were separated identified and quantified using GLC. Helium gas was used as a carrier agent. Methyl esters were identified by their comparison with chromatograms of standards. Peaks were identified by their retention time (KIESSLING et al., 2001).
Feeding Trials
Labeo rohita fingerlings with an average weight of 13.25± 5.23 g were obtained from Fish Hatchery Bhagtanwala, district Sargodha, Punjab, and brought to Laboratory at The University of Lahore. Fingerlings were acclimatized in glass aquaria dimensions Length × width × height (3’× 2’×2’). One third of aquaria were filled with fresh water and oxygen level was enhanced by electric aerator. Fifteen fingerlings were kept in each aquarium in triplicate. Water quality parameters i.e. dissolved oxygen; temperature and pH were kept within optimum range and recorded on daily basis. Fingerlings were fed twice a day @ 4% of their body weight. Experiment continued for 92 days. Fish weight and length were recorded before initiation of experiment and thereafter these growth parameters including feed conversion ratio (FCR), and survival rate were calculated fortnightly.

Enzyme activity
After completion of feeding experiment, three fish from each aquarium were randomly selected, sacrificed and their gut contents were taken out and stored in 10% formalin solution for further processing and to check their effect on enzyme activity in intestine.

Fish intestines were excised out and washed with cold distilled water. Intestines were homogenized by electrical homogenizer and centrifuged at 20,000 round per minute (RPM) for 10 min at 0°C. The supernatant was removed and used as enzyme assays for further proceeding to check proteolysis enzyme activity.

To check proteolysis enzyme activity prepared supernatant was mixed with phosphate buffer (pH 7.5) and 1% casein. Mixture was incubated at 35°C for 20 min. Thereafter, 5% TCA was added to stop reaction. Optical density of sample was checked by photometer at 280nm. Tyrosine curve was used as standard and activity was expressed as mole of tyrosine released/min/mg protein (KUNITZ, 1947).

For determination of lipase activity, reaction mixture was prepared by mixing phosphate buffer solution (pH 7.5), tissue homogenate, distilled water, and olive oil emulsion. After mixing, mixture was incubated at 24°C for 24 hour in electric oven. Mixture was titrated with 0.05N Sodium hydroxide (NaOH) until appearance of pink color. Amount of Sodium hydroxide (NaOH) required maintaining pH was used to measure amount of fatty acids released. Enzyme source without emulsion and buffer was used as standard (CHERRY; CRANDALL, 1932).

To determine Amylase activity starch solution, prepared in phosphate buffer (pH 7.5) was used as substrate. Already prepared reaction mixture consisting of enzyme supernatant, starch solution and distilled water were mixed well and incubated at 35°C for 30 min in electric oven. After 30 minutes, 5 ml of 5% dinitro salicylic acid (DNS) solution was added in mixture to stop reaction. Mixture was diluted by addition 20mL distilled water. Optical density was recorded by photometer at 540 nm. Standard maltose curve was used for comparison of amylase activity (RICK; STEGBAUR 1974).

Statistical analysis
Statistical analysis software (SAS) version 9.1 was used to analyze data. Analysis of variance (ANOVA) with significance level (p<0.05) was applied to check variations in mean values. The results are presented as means with standard deviation.

RESULTS AND DISCUSSIONS
Anti-nutritional factors present in raw material and palatability of mixtures must be considered (AZEVEDO, 1998) while formulating fish feeds. Other factors viz. price, availability of ingredients used as raw also play significant role in selection of ingredients to prepare fish diets. Soybean meal along with other ingredients such as rice bran, corn flour and milk powder can be successfully used for fish feed formulations (VIDOTTI et al., 2002; SHABBIR et al., 2003). All the factors were considered in preparation of fish feed during current study. Results of proximate analysis of treatment diets are presented in Table 2.

Table 2. Nutrient composition of various prepared treatment diets

| Parameters       | T₁  | T₂  | T₃  | T₀   |
|------------------|-----|-----|-----|------|
| Protein (%)      | 29.98±0.11 a | 30.14±0.07 a | 30.28±0.41 a | 32.14±0.18 a |
| Fats (%)         | 12.58±0.14 a | 11.81±0.41 a | 12.17±1.38 a | 9.41±0.71 b  |
| Ash (%)          | 8.69±0.49 a  | 9.89±1.32 a  | 9.31±0.48 a  | 4.88±0.54 b  |
| Carbohydrates (%)| 5.30±1.82 b  | 4.81±0.79 b  | 7.88±1.23 a  | 5.61±0.76 b  |

Means with same letters in a single row are statistically similar at (P<0.05).
Analysis of results showed that all treatment diets contained reasonable amount of lipid contents. The lipid contents were calculated as 12.58±0.14 %, 11.81±0.41 %, 12.17±1.38 % and 9.41±0.71 % in T$_1$, T$_2$, T$_3$ and T$_0$ diets, respectively. Amount of lipids in prepared silage varied according to type of raw material, life stage of fish and nature of diet to be fed to the fish (DEWAILLY _et al._, 2001; KHODDAMI _et al._, 2009). High concentration of lipid contents in the silage prepared during present project can be defended by NELSON AND COX (2000) who reported that quantity of fat contents in

| Sr. No. | Fatty Acid* | Fermented Fish silage |
|---------|-------------|-----------------------|
|         |             | I         | II        | III       | Average     |
| 1       | C$_{12}:0$  | 1.09     | 1.08  | 1.11        | 1.09±0.01  |
| 2       | C$_{14}:0$  | 13.43    | 14.26 | 15.21        | 14.3±0.89  |
| 3       | C$_{16}:0$  | 21.28    | 23.11 | 22.76        | 22.38±0.97 |
| 4       | C$_{16}:1$  | 12.32    | 12.46 | 13.12        | 12.63±0.42 |
| 5       | C$_{18}:0$  | 5.26     | 5.78  | 6.17         | 5.73±0.45  |
| 6       | C$_{18}:1$  | 28.21    | 27.32 | 26.13        | 27.22±1.04 |
| 7       | C$_{18}:2$  | 15.23    | 16.23 | 15.19        | 15.55±0.58 |
| 8       | C$_{18}:3$  | 1.86     | 1.96  | 1.42         | 1.74±0.28  |
| 9       | C$_{20}:0$  | 0.66     | 0.72  | 0.83         | 0.73±0.08  |
| 10      | C$_{20}:1$  | 0.83     | 0.94  | 0.83         | 0.86±0.06  |
| 11      | TSFA        | 41.72    | 44.95 | 45.97        | 44.2±2.21  |
| 12      | MUFA        | 40.53    | 39.78 | 40.08        | 40.13±0.37 |
| 13      | PUFA        | 17.09    | 18.19 | 16.61        | 17.29±0.81 |
| 14      | TFA         | ND       | ND    | ND           | ND         |

FA→ Fatty acid, TSFA → Total saturated fatty acids, PUFA → Poly unsaturated fatty acids, MUFA → Mono unsaturated fatty acids. Results revealed that saturated fatty acids (SFA), mono un-saturated fatty acids (MUFA), and poly unsaturated fatty acids (PUFA) were present in adequate amount in the silage prepared during present investigation and there amount was calculated as 44.21±2.21 %, 40.13±0.37 % and 17.29±0.81 %, respectively. The results indicated that fatty acids are present in relative pattern of SFA > MUFA > PUFA in pure fermented fish silage.

Our results of Fatty Acid profile are similar to the results as described by Disney _et al._ (1977) and Sales (1995). They analyzed Acid Silage and Fermented Fish Silage and reported 14.84, 12.20 and 14.29 g/100 g Lipid contents, respectively, in different Fish Silage samples. Similar value was also reported by Dapekevicius _et al._ (1998) who studied Acid and Fermented Fish Silages, and obtained values of 14.90 and 10.30 g/100 g respectively. These results are close to our findings. Our results about Fatty Acid profile showed similar pattern of Fatty Acid analysis e.g. SFA > MUFA > PUFA in the profile of Carp fish species from Indus River (JABEEN; CHAUDHRY, 2011). Differences in the pattern of fatty acids profile of fish carcass is influenced by ratio and types of Fats in the fish diet (TOCHER, 2003). Amount of Fatty Acids, especially Unsaturated Fatty Acids, strongly vary in the Wild and Farmed fish species because of variations in their diets (OZOGUL, 2007; KALYXCONCU _et al._, 2010). Eicosapentaenoic Acid (EPA 20:5n-3) Docosahexaenoic Acid (DHA 22:6n-3) and, probably, Arachidonic Acid (AA, 20:4n-6) are considered essential in the diets for normal growth, and survival of marine and fresh water aquatic organisms (SARGENT _et al._, 1999).
According to results, Total weight gain was recorded as 7.71±0.56, 7.27±0.47, 6.59±0.73 and 6.62±0.48 for T₁, T₂, T₃ and T₀, respectively. Statistical analysis showed non-significant difference (P<0.05) among treatments. According to Vincent et al. (2002) nutritional quality and digestibility of feed can be best evaluated by feeding trial. Weight gain, Length gain, FCR, SGR and Survival rate are useful parameters to evaluate feed digestibility and quality (VINCENT et al., 2002). Growth rate is mostly affected by protein concentration (KUMAR et al., 2010) and our all treatment diets were iso nitrogenous so there is non-significant difference in weight gain among all treatment groups. Protein level has severe effect on fish growth. They reported that protein is major nutrient and its concentration significantly affect fish growth rate (TARINDHA et al., 2003; SINGH et al., 2006). FCR was recorded as 2.44±1.45, 3.35±0.67, 3.79±1.58 and 2.59±0.98 for T₁, T₂, T₃ and T₀, respectively. Statistical analysis showed non-significant difference among T₂ and T₃ while these two groups showed significant difference from other two groups T₁ and T₀. Some other nutrients despite Protein affect digestibility and quality of feed which change FCR value (CHAUDHARY AND QAZI ,2007; MUSTAFA et al., 1994) also reported same results while working on effect of nutrition on growth rate of sea bream.

**Enzyme activity**

Enzyme activity of Protease, Amylase and Lipase is presented in Table 5. Protease activity was recorded as 15.63±0.56, 15.32±1.01, 14.02±0.26 and 14.06±0.26 in T₁, T₂, T₃ and T₀, respectively. The results indicated non-significant variation (P<0.05) among different treatments.

**Table 5.** Enzyme activity of Protease, Amylase and Lipase in digestive tract of *Labeo rohita* fingerlings fed on different treatment diets.

| Enzymes | Fermented Fish Silage | Control Fermented |
|---------|-----------------------|-------------------|
|         | 100% | 75% | 50% |                     |
| Protease | 15.63±0.56a | 15.32±1.01ab | 14.02±0.26ac | 14.06±0.26ab |
| Amylase  | 11.55±0.98a | 11.72±0.91a | 12.23±0.55b | 10.09±0.38b |
| Lipase   | 1.29±0.40a | 0.93±0.42b | 0.77±0.11c | 1.17±0.36a |

Means with same letters in a single row are statistically similar at ( P<0.05).

Enzyme activities are expressed as: Protease as Micromol of Tyrosine released/min/g protein; Amylase as Micromol of Maltose released/min/g protein; Lipase as units/mg protein.

Different researchers have shown different results in this regards. Lopez-Lopez et al. (2005) reported that there is no strong Correlation between Protease activity and dietary crude Protein level. With the increase of Carbohydrate and Protein quantity in the fish diet, activity of Amylase and Protease also increases (LE MOULLAC et al., 1994). However, if the concentration of these components increases beyond limits, concentration of Amylase and Protease start to decrease (CECCALDI, 1997). So our results can be explained on this ground that as our all treatment diets were iso-nitrogenous and Protease activity is mainly dependent upon Protein concentration so there is non-significant (P<0.05) variation among results.

Amylase activity was recorded as 11.55±0.98, 11.72±0.91, 12.23±0.55 and 10.09±0.38 in T₁, T₂, T₃ and T₀, respectively. Statistical analysis revealed Non-Significant differences between Amylase activities among all treatment diets. These results can be justified by the
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findings of Sabapathi and Teo (1993) and Hidalgo et al. (1999) who recorded lower amylase activity in Carnivorous fish and higher in Omnivorous fish. Our results about amylase activity are in line with who reported no correlation among amylase and dietary carbohydrates contents in diet in *Homarus americanus* and *Cherax quadricarinatus*, respectively (HOYLE, 1973; LOPEZ-LOPEZ et al., 2005).

Lipase activity was recorded in Fermented Fish Silage as $1.29 \pm 0.40$, $0.93 \pm 0.42$, $0.77 \pm 0.11$ and $1.17 \pm 0.36$ in $T_1$, $T_2$, $T_3$ and $T_0$ respectively. Statistical analysis of Lipase activity showed significant difference ($p<0.05$) among all treatments. $T_1$ treatment showed greater Lipase enzyme activity as compared to other treatments. Different researcher revealed different reasons behind Lipase activity variation. It is proved that that fish feeding habits and type of carbohydrate has strong effect Lipase activity (NRC, 1993; SABAPATHI; TEO, 1993). Temperature of environment and season of fish harvesting can influence the digestive enzyme activity in fish (KUZMINA et al., 1996). So our results may vary to some extent from the previous ones.

This difference may also be due to the variation in Lipid concentration in the treatment diets. Lipase activity varies according to the amount of Lipid contents in diets and this activity is more prominent in the Carnivorous fishes as compared to other fish species (CHAKRABARTI et al., 1995; LUNDSTEDT et al., 2004).

CONCLUSIONS

Fermented fish silage contains high concentrations of mono-unsaturated fatty acids which has positive impact upon *Labeo rohita* growth.

The activity of protease, amylase and lipase do not vary significantly ($P<0.05$) if protein, carbohydrate and lipid concentrations are kept constant in the diet.

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RESUMO: Devido à diferença na procura e na oferta de farinha de peixe há imensa necessidade de qualquer membro suplente da fonte de proteína. Tão presente projeto foi projetada para substituir a dispendiosa farinha de peixe (FM) com barato peixe fermentado de ensilagem (FFS) em alimentos para peixes. FFS foi preparado pelo processo de fermentação usando Lactobacillus bactérias e seu perfil de ácidos graxos e efeito sobre o sistema digestivo de *Labeo rohita* foi investigado. Conteúdo lipídico foram isoladas pelo aparelho de Soxhlet e registadas como $6.23 \pm 1.23$ g/100g de FFS. Perfil de ácidos graxos extraídos de lipídios foi determinada por cromatografia líquida de gás (GLC). Quantidade suficiente de ácidos graxos insaturados foram encontrados com padrão MUFA > SFA > AGPI. Tratamento de três dietas contendo silagem de 100% (T1), 75% silagem (T2) e 50% silagem (T3) foram preparados misturando com farinha de soja (SBM) e farelo de arroz como co ingredientes enquanto FFS foi substituído pela FM na dieta controle (T0). O experimento foi conduzido em aquários de vidro em triplicado. O Crescimento dos peixes foram anotados os parâmetros quínicos enquanto parâmetros físico-químicos de água foram registradas diariamente. Após a conclusão do teste de alimentação, três peixes foram aleatoriamente dissecada a impostos especiais de consumo os seus intestinos e determinar a atividade de protease, enzimas amilase e lipase. As variações não significativas ($P<0.05$) foi registrada em parâmetros de crescimento e atividade enzimática entre as dietas exceto enzima lipase mostrou diferença significativa entre as dietas de tratamento. Aparentemente, é possível concluir que a concentração razoável de FFS tem nutrientes e ácidos graxos insaturados de modo que ela possa substituir com êxito a farinha de peixe na dieta de peixes.

PALAVRAS-CHAVE: Alimentação. TLC. Amilase. Farinha de peixe. Ácidos gordos

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