Evaluation of the Efficacy of Fermented By-product of Mushroom, *Pleurotus ostreatus*, as a Fish Meal Replacer in Juvenile Amur Catfish, *Silurus asotus*: Effects on Growth, Serological Characteristics and Immune Responses

Kumar Katya, Yong-hyun Yun, Gunhyun Park, Jeong-Yeol Lee1, Gwangyeol Yoo2, and Sungchul C. Bai*

Department of Marine Bio-materials and Aquaculture / Feeds and Foods Nutritional Research Center, Pukyong National University, Busan 608-737, Korea

ABSTRACT: The present experiment was conducted to evaluate the efficacy of dietary fermented by-product of mushroom, *Pleurotus ostreatus* (FBPM) as a fish meal (FM) replacer in juvenile Amur catfish, *Silurus asotus*. A total number of 225 fish averaging 5.7±0.1 g (mean±standard deviation) were fed one of the five experimental diets formulated to replace FM with FBPM at 0%, 5%, 10%, 20%, and 30% (FBPM0, FBPM5, FBPM10, FBPM20, and FBPM30, respectively). At the end of eight weeks of the experiment, average weight gain (WG) of fish fed FBPM0 or FBPM5 were significantly higher than those of fish fed FBPM10 or FBPM30 diets (p<0.05). However, there was no significant differences in WG among the fish fed FBPM0, FBPM5 or FBPM10, and between fish fed FBPM10 or FBPM30, and also between those fed FBPM10 or FBPM30 diets. Lysozyme activity of fish fed FBPM0 or FBPM5 were significantly higher than those of fish fed FBPM10, FBPM15 or FBPM30 diets (p<0.05). The chemiluminescent response of fish fed FBPM5 was significantly higher than those of fish fed FBPM0, FBPM10 or FBPM30 diets (p<0.05). Broken line regression analysis of WG suggested that the maximal dietary inclusion level for FBPM as a FM replacer could be 6.3% without any adverse effects on whole body composition and on serological characteristics. Therefore, these results may indicate that the maximal dietary inclusion level of FBPM as a FM replacer could be 6.3% in juvenile Amur catfish. (Key Words: Fermented Mushroom, Fish Meal Replacer, Juvenile Amur Catfish, Growth, Immune Response)

INTRODUCTION

Aquaculture is a rapidly growing food-producing industry; however, there is still considerable potential for increased efficiency and efficacy of aquaculture through development of nutritious and cost-effective alternatives to traditional marine protein feedstuffs such as fish meal (FM) (Suarez et al., 2013). The FM cost 500 US dollars per metric ton in the middle of 90s but nowadays its cost has quadrupled (World Bank Commodity, 2010). Lee and Bai (1997a) noted that the world supply of FM increased only about 27% in 20 years and FM output by the major FM producing countries actually declined. Because of the limiting supply of FM around the world, the cost of producing fish is expected to continue to increase. Consequently, many studies have been conducted to replace FM or to reduce its inclusion in fish diets by various less expensive alternative protein sources. Some of the readily available ingredients studied with the view of using them to partially replace FM in fish feed include plant protein sources such as soybean meal, corn gluten meal and soy protein concentrate (Carter and Hauler, 2000; Refstie et al., 2001; Choi et al., 2004; Lim et al., 2004; Kim et al., 2008) and animal protein sources such as meat and bone meal, blood meal, feather meal, poultry by-product meal (PBM) and lysine by-product (Bai et al., 1997,1998; Lee and Bai, 1997a,b).
*Pleurotus ostreatus* is one of the most important mushroom species in Korea and the world. Worldwide production of this species has greatly increased during the previous few decades (Chang, 1999; Royse, 2002). The increase in popularity of this species is attributed to its ease of cultivation, high yield potential, high nutritional value, medicinal properties and other beneficial effects (Banik and Nandi, 2004). *Pleurotus ostreatus*, along with other species of mushroom, has been confirmed to have medicinal value. The biological functionality of these mushrooms ranges from antioxidative and immuno-stimulating to antiviral, anti-carcinogenic, anti-hypercholesterolaemic, and the ability to regulate blood lipid and glucose levels (Gordon et al., 1998; Wasser and Weis, 1999; Lakhanpal and Rana, 2005). The bioactive compounds in these species have been identified as oligosaccharides, polysaccharides, dietary fibers, glycoproteins, proteins, peptides, amino acids, triterpenoids, alkaloids, alcohols, phenols, polyphenols, vitamins, and/or minerals such as zinc, copper, iodine, selenium and iron. *Pleurotus ostreatus* contains high levels of glucans, which are polymers of glucose found in the cell walls of plants, fungi and bacteria (Sonck et al., 2010; Kim et al., 2011). It has been reported that β-glucans have the capacity to activate innate immunity, thereby enhancing defense barriers in animals including fish (Kim et al., 2006; Yoo et al., 2007; Sonck et al., 2010).

Amur catfish, *Silurus asotus*, is one of the important freshwater aquaculture species in Korea. The aquaculture production of Amur catfish in Korea in 2010 was 4,194 metric tons and this was valued at over ten million US dollars (FAO, 2012). However, using FM in fish feed increases the feed cost and is not a sustainable long-term feeding strategy (Naylor et al., 2009; FAO, 2010) and more work is needed to identify alternative protein-rich ingredients suitable for cultured fish (Kader et al., 2010). However, there are few nutritional studies on Amur catfish. In line with the dearth of knowledge regarding Amur catfish, the effects of dietary fermented by-product of mushroom (FBPM) as a FM replacer has not been investigated. Therefore, the present study was carried out to evaluate the optimum inclusion level of FBPM as a FM replacer in Amur catfish, *Silurus asotus*.

### MATERIALS AND METHODS

**Experimental design and diets**

Formulation and proximate composition of the experimental diets is shown in Table 1. Five experimental diets were formulated by replacing fish meal protein with fermented by-product of mushroom (FBPM) at different levels (10, 20, 30 and 40% dry weight basis) of dry matter. Five experimental diets were formulated to assess the effects of dietary FM replacement by FBPM on growth performance, feed utilization, hematological parameters and plasma biochemistry of juvenile Amur catfish. All the ingredients were purchased from local suppliers. The ingredients were weighed and mixed in a suitable proportion to prepare the experimental diets. The experimental diets were formulated by replacing fish meal protein with FBPM at different levels (10, 20, 30 and 40% dry weight basis) of dry matter. The proximate composition and chemical analysis of the experimental diets are presented in Table 1.

Table 1. Formulation and proximate composition of the experimental diets fed to juvenile Amur catfish (% of dry matter basis)

| Items                      | FBPM<sub>0</sub> | FBPM<sub>10</sub> | FBPM<sub>20</sub> | FBPM<sub>30</sub> | FBPM<sub>40</sub> |
|----------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| **Ingredients**             |                 |                 |                 |                 |                 |
| Fish meal<sup>2</sup>       | 30              | 28.5            | 27              | 24              | 21              |
| Fermented mushroom by-product<sup>3</sup> | 0               | 2.2             | 4.4             | 8.7             | 13.1            |
| Soybean meal<sup>4</sup>    | 14.3            | 14.3            | 14.3            | 14.7            | 15.5            |
| Corn gluten meal<sup>4</sup> | 12.1            | 12.1            | 12.1            | 12.1            | 12.1            |
| Squid liver powder<sup>4</sup> | 10.0            | 10.0            | 10.0            | 10.0            | 10.0            |
| Meat meal<sup>4</sup>       | 6.5             | 6.5             | 6.5             | 6.5             | 6.5             |
| Wheat flour<sup>4</sup>     | 19.6            | 19.6            | 19.6            | 18.4            | 16.0            |
| Fish oil<sup>5</sup>        | 1.5             | 1.7             | 1.8             | 2.1             | 2.4             |
| Soybean oil<sup>4</sup>     | 1.5             | 1.5             | 1.5             | 1.5             | 1.5             |
| Vitamin premix<sup>5</sup>  | 1.0             | 1.0             | 1.0             | 1.0             | 1.0             |
| Mineral premix<sup>6</sup>  | 1.0             | 1.0             | 1.0             | 1.0             | 1.0             |
| Cellulose<sup>4</sup>       | 2.5             | 1.6             | 0.8             | 0               | 0               |
| **Proximate composition**   |                 |                 |                 |                 |                 |
| Moisture                    | 10.9            | 9.5             | 9.6             | 10.0            | 10.7            |
| Crude protein               | 47.5            | 47.4            | 47.6            | 47.5            | 47.1            |
| Crude lipid                 | 8.9             | 8.9             | 8.9             | 8.9             | 8.9             |
| Crude ash                   | 10.9            | 10.3            | 10.7            | 10.2            | 10.3            |

FBPM, fermented mushroom by-product.

<sup>1</sup>FBPM<sub>0</sub>: 100% fish meal-based diet; FBPM<sub>10</sub>, FBPM<sub>20</sub>, FBPM<sub>30</sub>, FBPM<sub>40</sub>: 5%, 10%, 20%, and 30% fish meal protein replaced by FBPM, respectively.

<sup>2</sup>Suhuy Co., Busan, Republic of Korea.

<sup>3</sup>Mushroom Research Institute, Gyeonggi Province, Republic of Korea.

<sup>4</sup>Contains (as mg/kg in diets): ascorbic acid, 300; dl-calcium pantothenate, 150; choline bitartrate, 3,000; inositol, 150; menadione, 6; niacin, 150; pyridoxine-HCl, 15; riboflavin, 30; thiamine mononitrate, 15; dl-α-tocopherol acetate, 201; retinyl acetate, 6; biotin, 1.5; folic acid, 5.4; B<sub>12</sub>, 0.06.

<sup>5</sup>Contains (as mg/kg in diets): NaCl, 437.4; MgSO<sub>4</sub>·7H<sub>2</sub>O, 1,379.8; NaH<sub>2</sub>P<sub>2</sub>·H<sub>2</sub>O, 877.8; Ca(H<sub>2</sub>PO<sub>4</sub>)<sub>2</sub>·H<sub>2</sub>O, 1,366.7; KH<sub>2</sub>PO<sub>4</sub>, 2,414; ZnSO<sub>4</sub>·7H<sub>2</sub>O, 226.4; Fe·Citrate, 299; Ca-lactate, 3,004; MnSO<sub>4</sub>, 0.016; FeSO<sub>4</sub>, 0.0378; CuSO<sub>4</sub>, 0.00033; Calcium iodate, 0.0006; MgO, 0.00135; NaSeO<sub>3</sub>, 0.00025.
diets were formulated to contain different levels of FBPM to replace 0%, 5%, 10%, 20%, and 30% of FM (FBPM0, FBPM5, FBPM10, FBPM20, and FBPM30, respectively). The five diets were formulated to contain 47% crude protein and 16.0 kJ available energy/g. Fishmeal, dehulled soybean meal, corn gluten meal, squid liver powder and meat meal served as the major protein sources in the experimental diets; fish oil and soybean oil were used as lipid sources while wheat flour was the carbohydrate source. Cellulose was included in the diets to adjust the crude protein and energy levels. Mushroom by-product was obtained from Mushroom Research Institute, Gyeonggi Province, Republic of Korea. The mushroom by-product was fermented using a fermentation machine for 24 h at 30°C with lactobacillus and yeast at 3.93×10⁷ and 6.56×10⁸ CFU/g, respectively. It contained 42% crude protein, 0.3% crude lipid and 8.2% crude ash on a dry-matter basis. Procedures for diet preparation and storage were as previously described by Bai and Kim (1997). After thoroughly mixing the dry ingredients and fish oil with 30% filtered tap water, experimental diets were pelleted with a laboratory pelleting machine without heating using a 2-mm diameter module (Baokyong Commercial Co., Busan, Korea). After processing, all diets were kept at −20°C in a freezer until use.

**Experimental fish and feeding trial**

Juvenile Amur catfish, *Silurus asotus*, were obtained from Jeong-eup, Republic of Korea. Fish were transported to the experimental station (Pukyong National University, Busan, Republic of Korea) and acclimated to the experimental conditions for two weeks before the feeding trial began. During the acclimation period, fish were fed FBPM0 diet twice daily (1000 and 1800 h) at approximately 4% of wet body weight/day. A total number of 225 fish averaging 5.7±0.1 g (mean±standard deviation [SD]) were randomly distributed in fifteen individual fish groups to each of fifteen aquaria. Each aquarium was then randomly assigned to one of three replicates of five experimental diets. Triplicate groups of fish were fed the experimental diets twice daily (1000 and 1800 h) at approximately 4% of wet body weight/d at the beginning and 3% of wet body weight/d at the end of the feeding trial for eight weeks. Total fish weight in each aquarium was determined every 2 weeks, and the feeding rate was adjusted accordingly.

The feeding trial was conducted in an indoor semi-recirculation system with fifteen 40 L aquaria receiving filtered freshwater from the center tank. Supplemental aeration was provided to maintain dissolved oxygen levels near 6.5±0.5 ppm. Water temperature was 24±1°C (mean±SD); pH was 7.5±0.3 (mean±SD) and the photoperiod was 12:12 (light:dark) during the whole experimental period.

**Sample collection and analysis**

At the end of the feeding trial, all of the fish were weighed and counted for calculation of weight gain (WG), specific growth rate (SGR), feed efficiency (FE), protein efficiency ratio (PER), and survival. Three fish per aquarium were randomly selected and dissected to obtain the liver and white muscle samples, and hepatosomatic index (HSI) was calculated. After the final weighing, three fish were randomly collected from each aquarium and frozen at −20°C for analysis. A proximate composition analysis of the experimental diets was performed by the standard methods of AOAC (1995). Samples were dried to a constant weight at 105°C to determine moisture content. Ash was determined by incineration at 550°C, crude lipid by soxhlet extraction using a Soxtec system 1046 (Tecator AB, Hoganas, Sweden), and crude protein by Kjeldahl method (N×6.25) after acid digestion. The same fish as subjected to HSI calculation, were also used to analyze serological characteristics. Blood samples were obtained from the caudal vessels of the fish with 1 mL syringes. Hematocrit (HCT) was determined on three fish randomly selected per aquarium by the microhematocrit method (Brown, 1980), and hemoglobin (Hb) was measured in the same three fish by the cyan-methemoglobin procedure using Drabkins reagent. An Hb standard prepared from human blood (SIGMA Chemical, St. Louis, Missouri, USA) was used. Serum samples were prepared from blood on clotting by centrifugation at 1,600× g for 10 min and then the samples were stored at −20°C until analyzed. Serum was analyzed using commercial clinical investigation kits (NeoDin Co., Ltd., Seoul, Korea) and the specific method employed as follows: the biuret method for serum total protein and the enzymatic method for Glucose.

**Non-specific immune response assay**

**Chemiluminescent response**: Head kidneys were aseptically removed from three randomly selected fish from each tank, and pushed through a mesh with cold Hank’s balanced salt solution (HBSS) at 4°C. The suspension of leukocytes was immediately placed on a 34/51% Percoll (Sigma) density gradient and centrifuged at 2,000 rpm for 30 min at 4°C. The interphase was collected and the cells were washed twice at 2,000 rpm for 5 min in Dulbecco’s modified Eagle’s medium containing heparin (10 U/mL, Sigma) supplemented with streptomycin (100 U/mL, Sigma), penicillin G (100 μg/mL, Sigma). The cell viability was examined with trypan blue exclusion and it was evaluated to be greater than 98%. The leukocytes including neutrophils and monocytes were adjusted to 0.5× 10⁵ cells/mL HBSS. Samples were pre-incubated in 96 well plate each of 200 μL for 2 h at 25°C. Zymosan (10 mg/mL, Sigma) was mixed with the serum of adult rockfish, which were not used in this feeding experiment, and incubated at
25°C for 30 min. The opsonized zymosan was separated by centrifugation, washed three times and suspended in HBSS. The reactive oxygen intermediates (ROIs) produced by the reactive oxygen intermediates (ROIs) produced by the experimental fish were determined by measuring the amount of enzyme producing a decrease in absorbance of a suspension of *Micrococcus luteus* (Scott and Klesius, 1981). Measurements were made for 100 min until a linear reaction occurred. Measurements were made for 100 min until a linear reaction occurred. Measurements were made for 100 min until a linear reaction occurred.

**Lysozyme activity** The test serum (0.1 mL) was added to 2 mL of a suspension of *Micrococcus luteus* in 0.05 M sodium phosphate buffer (pH 6.2). The reactions were carried out at 20°C and absorbance at 530 nm was measured between 0.5 min and 4.5 min on a spectrophotometer. A lysozyme activity unit was defined as the amount of enzyme producing a decrease in absorbance of 0.001/min (Won et al., 2004).

### Statistical analysis

All data were analyzed by one-way (Statistix 3.1, Analytical Software, St. Paul, MN, USA) to test for the effects of the dietary treatments. When a significant treatment effect was observed, a least significant difference test was used to compare means. Treatment effects were considered at p<0.05 level of significance. Broken line model (Robinson Norton and Baker, 1979) was used to evaluate the maximal dietary level of fermented by-products of mushroom as a FM replacer.

### Results

**Growth performance**

Growth performance of Amur catfish, *Silurus asotus*, fed different levels of dietary fermented mushroom by-product, *Pleurotus ostreatus*, as a FM replacer is shown in Table 2. The average WG of fish fed FBPM₀ or FBPM₁ were significantly higher than those of fish fed FBPM₂₀ or FBPM₃₀ (p<0.05). However, there was no significant difference in WG among the fish fed FBPM₀, FBPM₁, and between fish fed FBPM₂₀ or FBPM₃₀, and also between those fed FBPM₂₀ or FBPM₃₀. The SGR of fish fed FBPM₀, FBPM₁, or FBPM₂₀ or FBPM₃₀ were significantly higher than that of fish fed FBPM₃₀ or p<0.05). The FE of fish fed FBPM₀ or FBPM₁ were significantly higher than that of fish fed FBPM₃₀ (p<0.05). The HSI of fish fed FBPM₀ was significantly higher than that of fish fed FBPM₂₀ or FBPM₃₀ (p<0.05). However, there were no significant differences in SGR, FE, PER, or HSI among fish fed FBPM₀, FBPM₁, or FBPM₂₀ or FBPM₃₀ diets (p>0.05). Also, no significant differences were found in survival rate of fish fed any of the experimental diets (p>0.05). Broken line analysis of WG indicated that FBPM could replace 6.3% of FM without any adverse effects on growth performance of Amur catfish, *Silurus asotus* (Figure 1).

**Table 2. Growth performance of juvenile Amur catfish, Silurus asotus, fed dietary fermented by-product of mushroom, Pleurotus ostreatus, as a fish meal replacer**

| Items                  | FBPM₀ | FBPM₁ | FBPM₂₀ | FBPM₃₀ | Pooled SEM² |
|------------------------|-------|-------|--------|--------|-------------|
| WG (%)                 | 278.3a| 273.7a| 226.9b | 201.8b | 140.4b      |
| SGR(%)                 | 2.37a | 2.35a | 2.11a  | 1.97a  | 1.53b       |
| FE(%)                  | 88.9a | 88.7a | 81.9ab | 80.1ab | 62.1b       |
| PER                    | 1.85a | 1.87a | 1.72ab | 1.69ab | 1.32b       |
| HSI (%)                | 1.77ab| 1.83a | 1.68ab | 1.72ab | 1.54b       |
| Survival (%)           | 77.8  | 77.8  | 77.8   | 77.8   | 68.9        |

FBPM, fermented mushroom by-product; Pooled SEM, pooled standard error of means; WG, weight gain; SGR, specific growth rate; FE, feed efficiency; PER, protein efficiency ratio; HSI, hepatosomatic index; SD, standard deviation.

¹ FBPM₀ 100% fish meal-based diet; FBPM₁, FBPM₂₀, FBPM₃₀; FBPM₅₀; 5%, 10%, 20%, and 30% fish meal protein replaced by FBPM, respectively.

² SD/√n. ³ WG (%) = [(final wt. – initial wt.)/initial wt.]×100. ⁴ SGR (%/d) = [(log final wt. – log initial wt.)/days]×100.

⁵ FE (%) = (wet weight gain/dry feed intake)×100. ⁶ PER (%) = wet wt. gain/protein intake. ⁷ HSI (%) = (wet liver wt./wet body wt.)×100.

Means of triplicate groups of fish where values in the same row with different superscripts are significantly different (p<0.05).

![Figure 1. Broken line analysis on weight gain (WG, %) of Amur catfish fed fermented by-product of mushroom (FBPM) as a fish meal (FM) replacer. Values of the X-axis are the FM:PM levels in experimental diets. Values are means±SD of 3 replication. SD, standard deviation.](image-url)
Whole body proximate composition

Whole-body proximate composition of fish fed dietary FBPM, *Pleurotus ostreatus*, as a FM replacer is shown in Table 3. The crude protein content was significantly lower for the fish fed FBPM than those of fish fed any of the other experimental diets (p<0.05). However, there was no significant difference in crude protein content among the fish fed FBPM, FBPM10, or FBPM20. Data for the moisture content showed, significantly higher moisture content for the fish fed FBPM30 than those of fish fed the other experimental diets (p<0.05). However, there was no significant difference in moisture content among the fish fed FBPM15, FBPM10, or FBPM20. While, crude lipid content was found to be significantly lower for the fish fed FBPM30 than those of fish fed the other diets (p<0.05). However, there was no significant difference in crude lipid content among the fish fed FBPM10, FBPM15, FBPM10, or FBPM20 diets (p<0.05). The data for the whole body ash content showed significantly lower value for the fish fed FBPM30, FBPM15, or FBPM10 than that of fish fed FBPM30 diet (p<0.05). However, there was no significant difference in ash content among the fish fed FBPM10, FBPM15, or FBPM10 and between fish fed FBPM20 and FBPM30 diets.

Serological characteristics

Table 4 shows the serological characteristics of Amur catfish, *Silurus asotus*, fed dietary FBPM, *Pleurotus ostreatus*, as a FM replacer. Hematocrit of fish fed FBPM20 was significantly higher than that of fish fed FBPM30 (p<0.05). There were no significant differences in this parameter among the fish fed FBPM0, FBPM15, FBPM10, or FBPM20 and also among those fed FBPM0, FBPM15, FBPM10, or FBPM30 diets (p<0.05). There were no significant differences in Hb or total protein of fish fed any of the experimental diets except for the Hb of fish fed FBPM0, which was significantly higher than those of fish fed FBPM15, FBPM10, or FBPM20 and the total protein of fish fed FBPM0, which was significantly higher than those of fish fed FBPM15, FBPM10, or FBPM30 diets. Serum glucose of fish fed FBPM0 was significantly higher than those of fish fed FBPM15, FBPM10, FBPM20, or FBPM30 diets (p<0.05). Also, serum glucose of fish fed FBPM0, FBPM15, or FBPM20 were significantly higher than that of fish fed FBPM30 diet (p<0.05). However, there was no significant difference in this parameter among fish fed FBPM0, FBPM15, or FBPM30 and also between those fed FBPM15, or FBPM30 diets (p<0.05).

Non-specific immunological responses

Non-specific immune response parameters of fish fed all the experimental diets are shown in Table 5. Lysozyme activity of fish fed FBPM0 or FBPM30 were significantly higher than those of fish fed FBPM10, FBPM20, or FBPM30 diets (p<0.05). However, there was no significant differences in lysozyme activity between fish fed FBPM0 or FBPM30, and also among those fed FBPM10, FBPM20, or

Table 3. Whole-body proximate composition of juvenile Amur catfish, *Silurus asotus*, fed dietary fermented by-product of mushroom, *Pleurotus ostreatus*, as a fish meal replacer

| Items     | FBPM0 | FBPM5 | FBPM10 | FBPM20 | FBPM30 | Pooled SEM² |
|-----------|-------|-------|--------|--------|--------|-------------|
| Moisture  | 76.7ᵇ | 76.5ᵇ | 77.6ᵇ  | 76.7ᵇ  | 79.6ᵇ  | 0.35        |
| Crude protein | 65.0ᶜ | 67.7ᵇ | 67.1ᵇ  | 67.2ᵇ  | 70.6ᵇ  | 0.51        |
| Crude lipid | 21.8ᵃ | 20.5ᵃ | 19.7ᵃ  | 20.2ᵃ  | 14.6ᵇ  | 0.71        |
| Ash       | 11.9ᵇ | 11.5ᵇ | 12.1ᵇ  | 12.8ᵇ  | 14.1ᵇ  | 0.31        |

FBPM, fermented mushroom by-product; Pooled SEM, pooled standard error of means.
1 FBPM0, 100% fish meal-based diet; FBPM5, FBPM10, FBPM20, FBPM30: 15%, 20%, 25%, and 30% fish meal protein replaced by FBPM, respectively.
² SD/√n. Means of triplicate groups of fish where values in the same row with different superscripts are significantly different (p<0.05).

Table 4. Serological characteristics of juvenile Amur catfish, *Silurus asotus*, fed dietary fermented by-product of mushroom, *Pleurotus ostreatus*, as a fish meal replacer

| Items       | FBPM0 | FBPM5 | FBPM10 | FBPM20 | FBPM30 | Pooled SEM² |
|-------------|-------|-------|--------|--------|--------|-------------|
| PCVᵃ       | 24.1ᵇ | 24.9ᵇ | 25.7ᵇ  | 26.9ᵇ  | 22.4ᵇ  | 0.61        |
| Hb (g/100 mL) | 14.7ᵇ | 18.4ᵃ | 15.2ᵇ  | 14.6ᵇ  | 16.1ᵇ  | 0.53        |
| Total protein (g/dL) | 3.1ᵃ | 2.7ᵇ | 2.7ᵇ  | 2.8ᵇ  | 2.6ᵇ  | 0.05        |
| Glucose (mg/dL)    | 76.3ᵇ | 73.7ᵇ | 69.0ᵇ  | 69.7ᵇ  | 64.3ᵇ  | 1.5         |

FBPM, fermented mushroom by-product; Pooled SEM, pooled standard error of means; PCV, packed cell volume; Hb, hemoglobin.
1 FBPM0, 100% fish meal-based diet; FBPM5, FBPM10, FBPM20, FBPM30: 15%, 20%, 25%, and 30% fish meal protein replaced by FBPM, respectively.
² SD/√n. ³ Hematocrit (%).
Means of triplicate groups of fish where values in the same row with different superscripts are significantly different (p<0.05).
Table 5. Nonspecific immune responses of juvenile Amur catfish, Silurus asotus, fed dietary fermented by-product of mushroom, Pleurotus ostreatus, as a fish meal replacer

|                     | FBPM₀ | FBPM₁ | FBPM₁₀ | FBPM₂₀ | FBPM₃₀ | Pooled SEM² |
|---------------------|-------|-------|--------|--------|--------|-------------|
| Lysozyme activity (U/mL) | 72.1a | 69.2a | 60.5b | 56.4b  | 54.6b  | 4.13        |
| CL (RLU/s)          | 42.32a | 78.435b | 71.839bc | 63.304bc | 39.868c | 1.43        |

FBPM, fermented mushroom by-product; Pooled SEM, pooled standard error of means; CL, chemiluminescent responses.

₁ FBPM₁₀: 100% fish meal-based diet; FBPM₁, FBPM₁₀, FBPM₂₀, FBPM₃₀: 5%, 10%, 20%, and 30% fish meal protein replaced by FBPM, respectively.

₂ SD/√n.

Means of triplicate groups of fish where values in the same row with different superscripts are significantly different (p<0.05).

FBPM₂₀ diets (p<0.05). The chemiluminescent (CL) response of fish fed FBPM₁ was significantly higher than those of fish fed FBPM₀, FBPM₂₀, or FBPM₃₀ diets (p<0.05). Furthermore, CL of fish fed FBPM₁, FBPM₁₀, or FBPM₂₀ were significantly higher than those of fish fed FBPM₀, or FBPM₃₀ diets (p<0.05). There were no significant differences in CL between fish fed FBPM₁ or FBPM₃₀, and between fish fed FBPM₁₀ or FBPM₂₀ and also between fish fed FBPM₁₀ or FBPM₂₀ diets (p<0.05).

**DISCUSSION**

Growing ecological and economical concern associated with the use of FM has exerted pressure on nutritionists to evaluate viable alternative protein sources to FM in aquafeed. Consequently, numerous studies have been done on replacement of FM in fish feed with plant protein sources such as soybean meal and other soy protein products (McGoogan and Gatlin, 1997; Day and Gonzalez, 2000; Choi et al., 2004; Lim et al., 2004; Sun et al., 2007; Kim et al., 2008). Animal protein sources such as feather meal, Hb powder, and meat and bone meal have equally been researched on as replacers for FM in fish feed (Bishop et al., 1995; Hasan et al., 1997; Lee and Bai, 1997a,b; Bureau et al., 2000). Although by-products of mushroom, Pleurotus ostreatus, have been used in ruminant feed (Adamovic et al., 1998; Silvana et al., 2006; Song et al., 2007; Oh et al., 2010; Kim et al., 2011), information on the use of these ingredients in fish feed is scarce. The present investigation showed that fermented mushroom, Pleurotus ostreatus, by-product can be used as a dietary FM replacer in juvenile Amur catfish, Silurus asotus.

Although WG, SGR, FE and PER decreased with FBPM supplementation, no significant differences were recorded in these parameters between fish fed the FBPM free diets and those fed diets supplemented with 5% and 10% FBPM (p<0.05). Since little information is available on FM replacement in Amur catfish and on the use of FBPM as a FM replacer in fish, it is difficult to conclusively state the cause of the reduction in growth performance of fish in this trial. However, FM replacement studies in other species using various plant and animal protein sources suggest that FM could be successfully replaced by these alternative protein sources up to certain levels without any adverse effects on growth performance/nutrient utilization but beyond such levels, growth/nutrient utilization could be impaired (Alexis et al., 1985; Brown et al., 1985; Shimizu et al., 1990; Hughes, 1993; McGoogan and Gatlin, 1997; Abdel-Warith et al., 2001; Lim et al., 2004). Abdel-Warith et al. (2001) recorded good growth of Clarias gariepinus up to 40% replacement of FM with PBM but noted depression in growth and abnormality in histological structure of fish with increasing levels of PBM in diets. Similarly, Sun et al. (2007) recorded an optimum replacement level of 30% of FM with fermented fisheries by-products and soybean curd residues mixture. Some of the suggested causes of a reduction in performance above a certain level of replacement in these studies include variations in protein content, amino acid profiles and digestibility, as well as the presence of some antinutritional factors and palatability depressants (Lim et al., 2004; Fasakin et al., 2005; Sun et al., 2007).

Studies in ruminants indicate that certain levels of various mushroom by-products can be included in the animal feed to replace other forage sources. Oh et al. (2010) reported that Pleurotus eryngii and Pleurotus ostreatus could be used to replace 40% of rice straw without any severe negative metabolic effects in Hanwoo steers. Xu et al. (2010) suggested that timothy hay can be replaced by 6.5% of spent mushroom substrate in a silage-based total mixed ration for wethers. Similar to the practice in ruminant nutrition, our results suggest that FBPM can be effectively incorporated in fish feed, replacing 6.3% of FM, the major ingredient in aquafeed.

Whole-body proximate composition of fish was not seriously affected by FBPM inclusion as a FM replacer in the diets (p<0.05). No significant differences were recorded by Lim et al. (2004) in whole-body proximate composition of growing rockfish, Sebastes schlegeli, fed dehulled soybean meal as a FM replacer. Furthermore, although significant differences were found in whole-body proximate composition of juvenile rockfish by the same authors, no clear trends were observed. Also, no clear trends were found in whole-body proximate composition of juvenile
olive flounder, Paralichthys olivaceus, fed fermented fisheries by-products and soybean curd residues mixture as a FM replacer (Sun et al., 2007).

Although significant differences were recorded in serological parameters of Amur catfish fed FBPM as a FM replacer, no clear trends were found. Only serum total protein and glucose of fish fed FBPM0 were significantly higher than those of fish fed other diets while rest of the parameters fluctuated among the different treatments (p<0.05). Serological characteristics can be used as an index of health status of fish (Blaxhall, 1972). Serological changes have been detected following different types of stress conditions like exposure to pollutants, diseases, hypoxia, etc. (Duthie and Tort, 1985). Hence, it could be suggested that any unhealthy condition caused by poor nutrition could affect the serological characteristics of fish. However, results in the present experiment do not indicate any observable adverse effects of FM replacement with FBPM, suggesting that FM replacement by FBPM does not adversely affect the serological characteristics of Amur catfish. Even the significantly high serum protein and glucose levels in fish fed FBPM0 do not conform to the trend in growth performance of fish in the present experiment. This may suggest serological parameters may not be greatly affected by FBPM inclusion. Similar to our findings, although growth performance was significantly affected by β-1,3 glucan and feed stimulants inclusion in olive flounder feed, out of the six serological parameters tested only HCT and serum glutamic oxaloacetic transaminase were significantly different among treatments (Yoo et al., 2007).

Lysozyme activity and CL response have been frequently used as indicators of nonspecific immune functions, which are of primary importance in combating infections in fish (Vazzana et al., 2003; Saurabh and Sahoo, 2008). During phagocytosis, phagocytic cells increase their oxygen consumption and ROIs are produced in a process called the respiratory burst (Scombbes, 1994). The CL response measures the photon emission resulting from the ROIs formation. Lysozyme, a mucolytic enzyme of leucocytic origin, is an important defense molecule of the innate immune system, which is important in mediating protection against microbial invasion (Saurabh and Sahoo, 2008). Increased lysozyme activity and CL response are indications of improved immunity responses in fish (Panigrahi et al., 2004; Kim and Austin, 2006; Taoka et al., 2006). Lysozyme activity of fish decreased with FBPM levels in diets. However, no significant differences were found between fish fed FBPM0 and those fed FBPM5, while CL response actually increased and then decreased after the FBPM inclusion level of 5% (p<0.05). These results show that immune responses of fish were not adversely affected due to FM replacement by FBPM up to 5%. However, as these two parameters had their peaks at two different inclusion levels, it is difficult to determine the inclusion level of FBPM which actually improved the nonspecific immune responses of Amur catfish in this trial.

However, the present experiment opens a new avenue to evaluate the optimum level of dietary FBPM as a FM replacer with supplemental phytase and amino acids viz. methionine and lysine. In conclusion, based upon the broken line regression analysis of WG the maximal dietary inclusion level for FBPM as a FM replacer could be 6.3% without any adverse effects on whole body composition and on serological characteristics in Amur catfish.

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