Introduction

Fish and seafood constitute an important food component for a large section of the world population. They come after meat and poultry as staple animal protein foods where fish forms a cheap source of protein (Wafaa et al., 2011). Fish serve as an important source of human dietary protein worldwide, especially in African countries (Kumolu et al., 2011). Catfish provides food for a lot of the populace in Africa. It allows for improved protein nutrition because of its high biological value in terms of high protein retention in the body. Protein assimilation in catfish is higher as compared to other protein sources and has low cholesterol (Adebayo et al., 2013).

Catfish production is important to the Nigerian economy. It serves as a source of income, reduces the rate of unemployment in the economy and increases the Gross Domestic Profit (GDP). In most countries, it fetches a higher price than tilapia as it can be sold live in the market as they have a market value two to three times that of tilapia (Emokaro, 2010). Catfish production is influenced by availability of fish feed and its quality. Fish feed quality, generally perceived as the responsibility of the feed manufacturer is affected by factors such as handling, storage and use. In the tropics, conducive environmental condition has majorly been attributed to production and growth of microorganisms and toxic substances; particularly fungi which can alter the quality of fish feed, limit the growth of fish and may result to fish mortalities (Adeniji et al., 2014).

These fungi are ubiquitous plant pathogens that are major spoilage agents of food and feedstuff. The infection of fungi can result into reduction in fish feed quality with contamination of poisonous fungal secondary metabolites called mycotoxins (Atanda et al., 2013).

Mycotoxins are toxic secondary metabolites produced by wide range of fungi (molds) toxic to humans, livestock and plants which explains the major concern of food and feed industries in preventing them from entering the food chain (Pierre, 2007). Mycotoxins are mainly produced by Aspergillus, Penicillum and Fusarium genera which invade crops in the field and may grow during storage under favourable conditions of temperature and humidity.
Mold infestation in stored feeds, reduces nutritional value owing to loss of dietary lipids, amino acids and vitamins by enzymatic digestion making the feeds to have poorer flavour, appearance and less palatable (Lim et al., 2008).

Materials and Methods

Experimental Location

This research was conducted in the Zoology Laboratory, Department of Biological Sciences, Benue State University, Makurdi, Nigeria, located on latitude 7° 43’ (7.7285°) North and longitude 8° 33’ (8.5540°) East (Google Earth, 2017).

Collection of Clarias gariepinus Juveniles

Forty Catfish (Clarias gariepinus) juveniles were obtained from a commercial fish farm in Makurdi, Benue State, Nigeria. They were transported to the Zoology Laboratory, in plastic buckets half filled with water.

Determination of Initial Length and Weight

The initial individual weights and lengths were measured. The weights were determined individually using the weighing balance and their lengths were taken using a measuring board. They were then randomly distributed to the two plastic tanks. The volumes of water in the tanks were maintained at 206 litres at a stocking rate of twenty juveniles per plastic tank.

Collection of Feed Samples

A bag of moldy (presence of grey mold) and another bag of non-moldy feeds were sorted and purchased from a fish feed store at High Level market, Makurdi, Benue State, Nigeria. Both feed samples were Top feeds. The nutrient component of the feed was labeled by the manufacturer as shown in Table 1 below.

Table 1: Feed composition

| Nutrient            | % Level |
|---------------------|---------|
| Crude protein       | 45      |
| Crude fat           | 10      |
| Crude fibre         | 3.0     |
| Ash                 | 7.0     |
| Calcium             | 1.5     |
| Total phosphorus    | 1.1     |
| Lysine              | 2.4     |
| Methionine +cystein | 1.4     |

Additives

- Vitamin A: 700 Iu/kg
- Vitamin D: 850 Iu/kg
- Vitamin E: 50 mg/kg
- Vitamin C: 200 mg/kg

Source: Premier Feed Mills Company Limited. A subsidiary of Flour Mills of Nigeria PLC (2018)
Isolation and Identification of Fungal Microorganisms from the Feed

Twenty grams of potato dextrose agar (PDA) was dissolved in five hundred milliliters (500ml) of distilled water and allowed to homogenize. The medium was then heated on the heating mantle to further dissolve particles and later sterilized by autoclaving in an autoclave at 121°C for about thirty minutes. An antibiotics (Chloramphenicol) was added to the medium (to inhibit the growth of any bacterial species that could contaminate the isolate), before pouring the melted medium into sterile Petri dishes and cooled at room temperature to solidify. One gram of moldy feed sample was ground using mortar and pestle to prepare a 10-fold serial dilution. This was then inoculated on the prepared medium plates and incubated at 25°C for 5-10 days. Developed colonies of fungi were sub cultured to obtain pure cultures. The fungi cultures were examined both macroscopically and microscopically for colony, mycelia, hyphae, conidia heads and spore characteristics. The characteristics were compared with those in a standard mycology textbook and chart (Olga, 1986).

Fish Feeding

Feed 1 was moldy free and served as control while feed 2 was moldy in appearance. Fish were fed at 5% body weight twice a day at 9:00am and 6:00pm for a period of 12 weeks. The ration was adjusted every week when the new weights of the juveniles for the various experimental tanks were determined. Feeding was done by hand. After washing hands and dry cleaning with a hand towel, the feed was fetched and spread evenly on the water surface of each tank.

Water Quality Parameters

Temperature

The weekly water temperature was measured using a portable thermometer, model JPB-607. The thermometer was switched on and reset to zero. It was then dipped in the plastic tank (at the center) and readings were taken when stabilized.

Dissolved Oxygen

Weekly dissolved oxygen was measured using a portable dissolved oxygen meter, model JPB-607. The dissolved oxygen meter was switched on and reset to read at zero after which it was dipped in the tank and the readings were read and recorded when they stabilized.

pH

The pH concentration was measured using Hanna water proof pH meter. The pH meter was reset at zero point and dipped in the center of the plastic tank. The readings were taken and recorded after stabilizing.

Data Analysis

The evaluation for growth and survival rate was carried out as follows:

\[
\text{Mean initial weight (MIW)} = \frac{\text{Total initial weight of juveniles}}{\text{Total number of juveniles}}
\]
Mean weight gain (MWG) = mean final weight – mean initial weight \hspace{1cm} (2)

\[ \text{Growth rate} = \frac{\text{Mean final weight} - \text{Mean initial weight}}{\text{Duration of experiment}} \times 100 \hspace{1cm} (3) \]

\[ \text{Percentage survival} = \frac{\text{Initial number of fish stocked} - \text{Number of dead fish}}{\text{Initial number of fish stocked}} \times 100 \hspace{1cm} (4) \]

\[ \text{Mortality rate} = \frac{\text{Number of deaths during period of experiment}}{\text{Total number of juveniles}} \times 100 \hspace{1cm} (5) \]

Results

The results of the fungi isolated are presented in the table 2 below with Aspergillus and Penicillum as the predominant fungi genera.

**Table 2:** characterization of fungal isolates from fish feed sample on PDA

| Macroscopic Characters | Microscopic Characters | Probable Organism |
|------------------------|------------------------|-------------------|
| Colony colour ranged from grayish to dark blue in colour. | The conidia colour was dark green to blue. Conidiophores length was 265-365 μm and it was uncoloured to grayish brown. | *Aspergillus fumigatus* |
| Colony colour was parrot green to deep green. | Conidia were light green in colour. Conidiophores length ranged from 550-678μm | *Aspergillus flavus* |
| The colour of the colony was grayish to dark green with a white dark border. Growth rate moderate to rapid. | Conidiophores and stipes were smooth walled, with stipes terminating in well-defined venticils of 3-5. Conidia were spherical to subspherical with smooth walls borne on fine columns | *Penicillium digitatum* |

Source: Elise, et al., (2016)

**Table 3:** Survival, mortality and growth rate of juveniles during the study

| Growth parameters (grams) |  |
|---------------------------|--|
| Mean initial weight       | 1.61g |
| Mean weight gain          | 15.62g |
| Growth rate               | 130.17g |
| Percentage survival (%)   | 55% |
| Mortality rate (%)        | 45% |
**Fig.1** Moldy feed (Mean length and weight) growth

**Fig.2** Non moldy feed (Mean length and weight) growth

**Fig.3** Mean dissolved oxygen, pH and temperature during the study
Discussion

In this study, the results of fungi isolated from moldy feed showed *Aspergillus* and *Penicillium* as the predominant fungi genera as also stated in the work of Ariyo *et al.*, (2013). Mycological examination was carried out for 150 samples of fish rations by El-Boshy *et al.*, (2008) and a total prevalence of *Aspergillus flavus* was observed to be 66.6%. Further, El-Boshy, isolated fifteen aflatoxin B₁ producers. Again El-Boshy observed that various changes such as sluggish swimming, darkening of skin, loss of reflexes, increase in mucus secretion, loss of scales and ascities were in aflatoxin B₁ intoxicated fish. Internally, the study showed that the liver displayed pale coloration with patches of congestion and hemorrhage, the spleen and the kidneys were darker in colour and appeared to be enlarged and congested. More so, growth rate of catfish juveniles fed moldy feed was seen to be slower than catfish juveniles fed with non-moldy feed. This could be due to less consumption of feeds by juveniles fed with moldy feed. This study is similar to Oluwafemi (2009) who carried out a study to investigate the performance of catfish fed with three different doses of aflatoxins isolated from fish feeds. The study showed that the growth of aflatoxicosed fish was significantly different from the control. More over protein, cholesterol, bilirubin, electrolytes and liver enzymes such as Aspartate transaminase (AST), Alanine transaminase (ALT) and Alkaline phosphate (ALP) were also significantly different from the control and the liver was observed to be severely damaged. Caguan *et al.*, (2004) reported loss of appetite, decreased mean total biomass and low survival percentage in tilapia when fed with aflatoxin contaminated feed. Furthermore, a higher mortality was recorded with the juveniles fed with moldy feed compared to juveniles fed with non-moldy feed. Out of the twenty juveniles stocked and fed with moldy feed, fifteen died and only five survived within a period of twelve weeks while out of twenty juveniles that were stocked and fed with non-moldy feed, three died and seventeen survived within duration of twelve weeks giving a mortality rate of 45% (Table 2). This result is similar to the findings of Adeniji *et al*. (2014) who carried out similar studies on *Heterobranchus longifilis* and obtained similar results.

In addition, lower weight gain and high mortality of juveniles fed with moldy feed as compared to juveniles fed with non-moldy feed was observed in this study (Fig. 1 and 2). This could be due to low feed consumption as presence of fungi reduces the nutritional value of feed and makes it less palatable (Lim *et al.*, 2008). The results showed a significant difference in the growth and survival of catfish fed moldy feed and catfish fed non-moldy feed. The result of this study is similar to Francis *et al.*, (2010) who evaluated the effect of dietary fumonisin B₁ on the fingerlings of *Clarias gariepinus*. The diets used contained approximately 5, 10 and 15 mg FB₁/kg respectively and the weight gain was noted to have reduced significantly.

The water parameters were all within a tolerable range for the growth of the juveniles. This could be attributed to low stocking density of the fish per tank, constant monitoring of the selected physicochemical parameters and addition of fresh water to the tanks. Again, some physico-chemical parameters taken within the period of this experiment showed no significant difference in the tank with juveniles fed with moldy feed and those fed with non-moldy feed. The water parameters were all within a tolerable range for the growth of juveniles (Fig. 3).
was no significant difference in the physico-chemical parameters taken, so the difference in growth, survival and mortality rates of the juveniles was influenced by moldy and non-moldy feed.

**Conclusion**

During the study, *Aspergillus* and *Penicillium* were the predominant fungi genera isolated and identified from the moldy feed. It was observed that juveniles fed with moldy feed had slower growth, lower survival rate and higher mortality as compared to juveniles that fed on non moldy feed. Survival rate of juveniles fed with moldy feed was five and mortality was fifteen. It was observed that the juveniles that were fed with moldy feed showed normal growth from the beginning but eventually died when they could no longer tolerate the feed which resulted to their death. Some water quality parameters such as temperature, dissolved oxygen and pH measured were all within a tolerable range for the growth and survival of the juveniles.

**Recommendation**

Fish farmers should use only non moldy feed to feed their fish. Also, sellers of fish feeds should properly handle fish feeds to avoid infestation and contamination by mold.

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