Comparison of the Aflatoxin levels in Fish Feeds from Rainbow Trout Farms Localized at Different Regions (Adana- Ağrı, TURKEY)

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Abstract

Aflatoxins levels in different fish feeds obtained from two different rainbow trout (Oncorhynchus mykiss) farms located in Adana and Ağrı regions were determined between May and July 2017. During the experiment period, “a drought status” was reported according to the Standardized Precipitation Index of Turkey by Turkish State Meteorological Service. According to precipitation index records, while Adana was reported as slightly humid, Ağrı was slightly dry in the period between May and July 2017. The aflatoxin analyses of fish feed samples was carried out in the laboratories of the Ministry of Food, Agriculture and Livestock in Adana, Turkey. The aflatoxin levels of the samples were resolved by liquid-solid extraction, immune-affinity pillar clean-up using a HPLC-FD (High Performance Liquid Chromatography with Fluorescence Detection). The detection limits of aflatoxin were 0.50, 0.23, 0.73, 0.20 ppb for AFB1, AFB2, AFG1, AFG2, respectively. The results showed that, aflatoxin levels of fish feeds were below the detection limits in all samples.

Keywords: Aflatoxin, fish feed, HPLC analysis, rainbow trout farm trout

INTRODUCTION

Aquaculture, husbandry of aquatic food organism, represents the fastest growing animal food-producing sector in many parts of the world. Rainbow trout (Oncorhynchus mykiss) is one of the most important freshwater fish species cultured in Turkey and the total rainbow trout production was 101,166 tons in inland farms and 6,872 tons in marine environment in 2015 (TÜIK, 2015). The increasing market demands to this species have promoted the feed industry to produce specifically formulated, higher quality feeds for rainbow trout.
However, mycotoxin contamination is one of the most important obstacles in feed manufacturing industry. As the fish producers and feed manufacturers realize the mycotoxins significance and their potential effect on production, it becomes an essential topic for aquaculture (Gonçalves et al., 2018).

Mycotoxins are natural secondary toxic metabolites produced by fungi, particularly the moulds. The most detected mycotoxins in manufactured feeds are aflatoxin (AF), deoxynivalenol (DON), zearalenone (ZEN), ochratoxin A (OTA), fumonisins (FN), T-2 and HT-2. Among these, AF are the most studied group of mycotoxins and are mainly produced by two species of the genus Aspergillus which are abundant particularly in areas with hot and humid climates (Marin et al., 2013). The most occurring forms of aflatoxins are AFB1, AFB2, AFG1 and AFG2. Aflatoxins are highly toxic and carcinogenic compounds that cause disease in animals and humans (Huang et al., 2010; Tsakiris et al., 2013). In particular, AFB1, the most hazardous aflatoxin can be produced by Aspergillus flavus, which affects the animals’ growth performance due to the decrease in feed intake. In addition, the histopathological changes in observed in the liver tissue, immunosuppression, hepatotoxicity, mutagenicity were also reported in aquatic animals which cause fish mortalities (Ashley et al., 1965; Black and Baumann, 1991; Tacon, 1992; Richard, 2007; Han et al., 2010; Alinezhad et al., 2011; Rajeev et al., 2011). Baranyi et al. (2015), concluded that the presence of mycotoxins in feed depends on many factors including the season, temperature, humidity, region and the method of storage. Among them, climate is the most important factor for mycotoxin production. Fungi can produce a wide range of secondary metabolites resulting in production of aflatoxin under ecological conditions which are convenient for their growth. Various fungal strains need specific environment to produce mycotoxins and thus, routine monitoring of these toxins particularly in stored fish feeds is crucial in the fish farms where the humidity levels are high during the summer times.

This study was carried out to determine and compare aflatoxin levels of fish feeds samples, taken from rainbow trout farms in two climatologically different regions of Turkey.

MATERIAL AND METHODS
The Origin of Samples and Sampling
The fish feeds were taken from different rainbow trout farms located in Adana and Ağrı regions over a period of three months (May-July 2017). Fish feed samples (500 grams) for four times each months have been moved to black bags in order to avoid exposure to the sun. The samplings were carried homogeneously by mixing original packed bags. Information concerning feed samples was noted as written in original feed bags.
Figure 1 shows 12-month comparative drought conditions in Turkey according to standard precipitation index of the Turkey. It reported that the Ağrı region was slightly dry (Figure 2) while Adana/Kozan region was slightly humid (Figure 3) over the experiment period of three months.

**Figure 2. Drought Analysis in Ağrı (MGM, 2018)**

**Figure 3. Drought Analysis in Adana/Kozan (MGM, 2018)**
Detailed information was also obtained about the environmental conditions in which the feed was kept. Temperatures were ranging from 14 to 35 °C. All the feed samples were homogeneously ground with blender and stored in a plastic container in a refrigerator until analysis. The analysis of feed samples was carried out in laboratories at the Ministry of Food, Agriculture and Livestock in Adana, Turkey.

**Extraction, HPLC-Apparatus and Conditions of Chromatography**

The aflatoxins were resolved by liquid-solid extraction, immune-affinity pillar clean-up using by a high performance liquid chromatography with fluorescence detection (HPLC-FD). The detection limit of the analysis was determined as 0.2 ppb (Vicam, 2007). Fifty grams of fish feed samples were extracted with 100 mL methanol and 25 mL water by using a blender at high speed for 1-2 min. Whatman No. 4 filter paper were used for filtration of the extract. A 10 mL aliquot of the filtrate was diluted with 40 mL ultrapure water, shaken vigorously and then filtered once more by a glass microfiber filter. The final volume of 10 mL filtrate was swiftly passed through column at a rate of 1-2 drop/sec. Afterward 2-3 mL air was passed through column.

AFs were separated from solvent by passing twice 1 ml of methanol through the column at a flow rate of 2-3 mL min⁻¹ and collected in vials. The column temperatures were retained at 35 °C for aflatoxins. The injection volume to HPLC apparatus for both standard and sample was 100 μL. For AFs analysis, the HPLC mobile phase was the mixed with the solution of water-acetonitrile-methanol (6:2:3, v/v/v) and the flow rate was 1 mL min⁻¹. Afterward, 120 mg potassium bromide (KBr) and 350 μL nitric acid (HNO₃; 65%) were added to 1 liter of mixed solution for electrochemical derivatization (Cobra Cell). The fluorescence detector was regulate to an excitation and emission wavelengths of 360 and 430 nm. The total run time for one cycle was configured for 20 min, and under these conditions, the retention times of AFB1, AFB2, AFG1 and AFG2 were 10.4, 8.7, 7.7 and 6.3 min, respectively.

**RESULTS AND DISCUSSION**

The liquid-solid extraction, immune affinity column clean-up in pursuit of HPLC-FD determination was performed to analyse the aflatoxin level in the fish feed samples. Figure 4 shows the level of aflatoxins (AF; sum of aflatoxin B1, B2, G1 and G2) in fish feeds, which were taken from different rainbow trout farms, located in Adana and Ağrı regions over a period of three months (May-July 2017).

The detection limit of this analysis was determined 0.50; 0.23; 0.73; 0.20 ppb for B1, B2, G1, and G2, respectively. The results obtained from this study show that the aflatoxin levels in the feed samples during the research period was found below the detection limits in all fish feed samples.

Mycotoxin contamination of fish feed is a common problem in the countries with a humid tropical climates conditions that allow to mould growth. In addition, improper feed processing methods and storage conditions, which are suboptimal: temperatures about equal or higher than 27°C, and moisture at levels greater than 14% cause mould growth (Santacroce et al., 2008; Russo and Yanong, 2010). In this current study, the Adana region is more humid, the aflatoxin B1 level was found lower in June. Gonçalves et al. (2017) were investigated 25 samples of finished fish feed which were obtained from Asia and Europe for mycotoxins. They found that most of the feed samples were contaminated mainly by Fusarium mycotoxins. However, Hashimoto et al., 2003 were evaluated the aflatoxin and fumonisins (mycotoxin) contamination in 42 feed samples taken from the region of Londrina, Brazil. The aflatoxin levels ranged from non-detectable to 15.60 ppb, where 61.90% showed <4 ppb levels, which are in accordance with the Brazilian guideline (20 ppb). The fumonisins levels ranged from non-detectable to 11.2 ppb and 76.20% samples were less than 4 ppb. Altuğ and Beklevik (2003) were analysed eighty-five samples taken from fish farming processes, feed factories and imported feeds for aflatoxin. As a result of the analyzes, they were found that total aflatoxin levels were in the range of 21.2-42.4 ppb in 20 feed samples, 5.0-20.0 ppb in 22 feed samples and below detection limits in 43 samples. Levels of aflatoxin were higher in samples taken from fish...
Figure 4. HPLC-FD Analysis results of the level of aflatoxins in fish feeds
A. The level of aflatoxins in fish feeds taken from Adana in May
B. The level of aflatoxins in fish feeds taken from Ağrı in May
C. The level of aflatoxins in fish feeds taken from Adana in June
D. The level of aflatoxins in fish feeds taken from Ağrı in June
E. The level of aflatoxins in fish feeds taken from Adana in July
F. The level of aflatoxins in fish feeds taken from Ağrı in July

farming processes than from factory or imported feed samples.

In this study although AFs levels in all fish feed samples was below the recommended limits (50 ppb), it was important to underline the feed intake with toxin at low concentrations along the time since it produced chronic adverse effects in production of animal.

Due to adverse effects of aflatoxin, the limit values are applied in many countries in the framework of safe fish production of the farms in order to protect the consumers. Different countries have established a wide range of aflatoxin standards for fish feeds. The number of countries regulating mycotoxins has significantly increased over the years. Currently, more than 90 countries have adopted mycotoxin regulations for feed to limit mycotoxin exposure. According to aflatoxin standard used in Turkey in 2003, the maximum limits for all feedstuffs were 50 ppb (Grace et al., 2015). However, Food and Drug Administration (FDA) has set the action levels for aflatoxins present in animal feeds as 20 ppb. Establishing legal maximum concentrations of mycotoxins does not indicate that mycotoxins are safe at low concentrations. In small quantities, mycotoxins may also interfere with normal cellular function and even lead to cell death and accumulated pathological changes (Bryden, 2012). It has been reported that aflatoxins at 0.01 ppb levels in fish feed are capable of causing malignant tumours in fish (Cengizler, 2000). El-Sayed and Khalil (2009) described also that a prolonged oral administration of low levels of AFB1 (0.018 mg/kg body weight) to European seabass causes critical health problems and
represents a major risk to consumers through aflatoxin residues in fish muscle.

CONCLUSION
At the present, the feed sector is not keeping pace with developments in disease control, anti-contamination standards or animal genetic improvement. Further information and analysis are required for providing science-based solution to form a policy and the development of standards for fish feed. This study contributed increasing knowledge on presence aflatoxin levels in fish feeds for rainbow trout farm located in Adana and Ağrı, Turkey. Future studies could be conducted about fungi contamination and analyze the other mycotoxins such as fumonisins, zearalenone and ochratoxin A in fish feeds.

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