

ISOLATION AND IDENTIFICATION OF MYCOTOXIGENIC ORGANISMS IN POULTRY FEED FROM SELECTED LOCATIONS IN ABIA STATE, NIGERIA

ABSTRACT

Feed contamination by fungi can lead to nutrient losses and detrimental effects on animal health and production. This present study was designed to isolate and identify the mycological contamination of poultry feeds in some selected parts of Abia state (farms and feed depots in Umuahia north, Osisioma and its environs). A total of 120 samples were collected and used for the study. The samples were screened and processed using spread plate technique. The isolates were identified using slide culture technique. From the samples collected, the fungi contamination in feed samples from depots in Umuahia was 50%, Osisioma 78% and in farms it was 85%. Five fungi organisms were isolated from the feed sample which includes *Aspergillus*, *Penicillium*, *Fusarium*, *Mucor* and **Yeast** which were seen in almost all the feed samples. *Aspergillus* (87%) recorded the highest percentage occurrence, followed by *Penicillium* (27%), *Fusarium* (24%), **Yeast** (5%) and **Mucor** (2%). The total fungi load was significant at 2.0×10^5 CFU/g=1 for feed samples from Umuahia North Local government Area, 7×10^5 CFU/g=1 from Osisioma feed depot and 1×10^6 CFU/g=1 from poultry farms thereby making the feed samples unsafe for poultry consumption. Therefore, there is need for screening of feeds in these locations in Abia state due to its high fungal load and percentage contamination.

Keyword: Feed, mycological agents, identification, occurrence, location

INTRODUCTION

The presence of microscopic fungi affects the quality of feeds, their organoleptic attributes and nutritional quality [1]. Moulds like other microorganisms will assimilate and utilize the most readily available nutrient in the material they grow upon and spoilage may lead to the loss of some of the nutrients in the feed [2].

29 Among microorganisms, fungi have important effects on the quality of feed. Fungi
30 growth sometimes leads to non-consumption of feed for poultry [3, 4]. Several factors
31 may lead to the spread of fungi infections such as geographical location, storage
32 conditions, processing of various feeds and moisture. Among the mentioned factors,
33 moisture is the most important factor, hence, rendering the moisture in feed constant to
34 lesser percentage will eliminate fungal growth and aflatoxin production will be stopped
35 [5]

36 Mycotoxins are poisonous toxins/substances and secondary metabolites produced by
37 fungi ([6]; [7]; [8]). The filamentous general of fungi produces secondary metabolites
38 which have deleterious effects on human and animal consumers following consumption
39 of contaminated foods and this ultimately affects the economy of the country [9].

40 Most toxic species belong to the genera *Aspergillus*, *Penicillium*, *Fusarium*, *Alternaria*
41 and produce mycotoxins that are of public health importance/concern such as aflatoxin,
42 ochratoxin A, T2-toxin, fursarotoxin, furmonisins, patulin, zearalenone and
43 deoxynivalenol [10];[11]; [12]. Feed contaminated with mycotoxins negatively affect
44 poultry performance and their health [13]. Most mycotoxicosis of poultry is caused by
45 intake of low concentration of contaminants over a long period resulting in the typical
46 chronic symptoms of poor growth, poor feed efficiency and suboptimal production.
47 Ingestion of high concentration however leads to acute clinical symptoms associated
48 with specific vital organs, the immune system and other aspects of avian physiology as
49 well as mortality [14]. Fungi causes a significant loss in the poultry industry being
50 responsible for high morbidity and mortality rate especially in young birds and causes
51 stunted growth and diarrhea and fetal encephalitis [15]. They also cause drop in egg
52 production leading to economic losses [16]. In this work, presence of potentially
53 mycotoxigenic fungi in samples of poultry feed was determined. Abia state is in Nigeria
54 which is a tropical country with a predominant hot humid environment and the
55 environment is much favorable for the propagation of fungi on feed and feed materials.
56 To prevent economic losses in poultry, isolation and identification of birds affected by
57 fungal infection needs to be determined and such studies on commercial broiler feed

58 sample in Abia state is not well reported. This informs the essence of this study in Abia
59 state south east Nigeria.

60

61 **MATERIALS AND METHODS**

62 **STUDY AREA**

63 Samples were collected between April – June from 2 local government areas of Abia
64 state. (Umuahia and Osisioma)

65 Abia state in Nigeria is located in a tropical rainforest between latitude 543N and
66 longitude 752E. The average annual temperature and rainfall are 26.9⁰C and 2193mm
67 respectively [17].

68 **SAMPLE COLLECTION**

69 Poultry feeds were sampled from farms and different feed depots in 2 different local
70 government areas (Umuahia North and Osisioma). The total feed samples collected
71 were one hundred and twenty (120) (which includes Top feeds, Vital, Animal care, and
72 Apex feed) used to isolate and identify the presence of mycotoxigenic fungi. Forty (40)
73 feed samples were collected from each of the locations. Also 40 feed samples were
74 collected randomly from poultry farms within the 2 localities. The representative
75 samples were collected batch by batch using simple random sampling technique. The
76 sampling plan was carried out according to Food and Agriculture Organization [18].
77 Take 10g from each batch and mix them together. Samples were collected two weeks
78 intervals and collection lasted three months (April-June). Fungal contamination and
79 fungal count determination was carried out in each sample to determine the fungal
80 genera and the total fungal population in the Department of Veterinary Microbiology
81 Laboratory of Michael Okpara University of Agriculture, Umudike.

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83 **Fungal Isolation and identification**

84 **Laboratory procedure**

85 Sabouraud dextrose agar medium was used for the isolation of fungi in the feed
86 samples. The medium was prepared aseptically following the manufacturer's
87 description. After autoclaving, a calculated amount of *Penicillium* and streptomycin was

88 mixed with the medium to help inhibit the growth of bacteria. Therefore, the medium
89 was dispensed into sterile Petri dishes in aseptic environment.

90 Serial dilution plate technique [19] was used for fungal isolation and general fungi
91 counts. One gram of each of the representative samples was mixed with 9ml of sterile
92 distilled water on a horizontal position and shake for 30mins to form uniform
93 suspension. For each feed sample, five dilutions 10^{-1} , 10^{-2} , 10^{-3} , 10^{-4} 10^{-5} were made
94 from each dilution, 0.1ml of dilution was aseptically inoculated on Sabouraud dextrose
95 agar supplemented with penicillin and streptomycin [20]. A surface spread plate
96 technique was used to achieve uniform distribution of the spores. Inoculated plates
97 were incubated at 25⁰c for 5-7days for isolation of the fungi and overall quantitative
98 enumeration of fungal colonies per gram of the feed sample; isolates were identified
99 based on colonial and microscopic morphologies ([21]; [22]). Microscopic examination
100 of the isolate was done using wet mount and slide culture technique [23]. The relative
101 occurrence of fungal genera was calculated in percentage using the following

102

103 Percentage occurrence of fungal genus:
$$\frac{\text{Number of isolates} \times 100}{\text{Total Number of Fungi}}$$

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106 Total fungal load CFU/g:
$$\frac{\text{Number of colonies} \times \text{dilution factor}}{\text{Volume used}}$$

108

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110 4.1 RESULTS

111 Plate 1 and Plate 2 show the morphological presentation of the colonies of *Aspergillus*
112 species and *Fusarium* species which appear in form of an emulsion as brownish and
113 whitish coloration. Plate 3 show colonies typical of *Penicillium* species marked with
114 remarkable in-folding while Plate 4 reveals different colorations consistent with colonies
115 of mixed fungi infection.

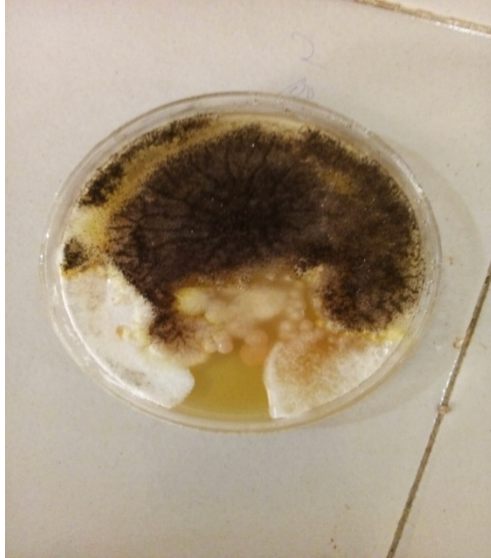
116 Figure1 shows the presence of *Aspergillus* spp at magnification of $\times 100$, the marked
117 evidence of oval dark hypha measuring about 0.5mm can be comparable to similar
118 findings of *Aspergillus* occurrence. Figure 2 presents a remarkable cauliflower with
119 distinctive dark hypha and unique long conidiospore characteristic of *Aspergillus*.
120 Figure 3 shows marked long conidiospore with many branches about (6-10) with long
121 dark hypha and diameter of about 0.5mm. Figure 5 indicates aggregates of fungi hypha
122 called mycelium.

123 From the study, the fungi species isolated and identified down to genus level are
124 *Aspergillus*, *Penicillium*, *Fusarium*, yeast and *Mucor*.

125 Table 1 shows that *Aspergillus penicillium* and *Fusarium* contamination was recorded in
126 the 3 locations, yeast was absent in samples collected from Umuahia while *Mucor* was
127 present only in farms. Table 2 shows the total samples collected from each location and
128 the positive numbers, 85% were positive from farms, 78% were positive from Osisoma
129 and 50% were positive from Umuahia.

130 From the above study, the genus aspergillus had the highest frequency of isolate at
131 85% followed by *Penicillium* (27%), *Fusarium* (25%), yeast (5%) and *Mucor* (2%) as
132 shown in figure 6. Table 3 shows that feed sampled from farms has the highest fungal
133 load followed by samples from Osisioma then samples from Umuahia feed distributors
134 and depot. Table 3 suggests that the feed sampled from poultry farms and Osisioma
135 feed distributors and depots have much fungal load of about 1×10^6 and 7×10^5 compared
136 to that from Umuahia which have fungal load of about 2.0×10^5

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Plate 1. Colonies of *Aspergillus* and *Fusarium*. Morphological view

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Plate 2. Colonies of *Aspergillus* Morphological view.

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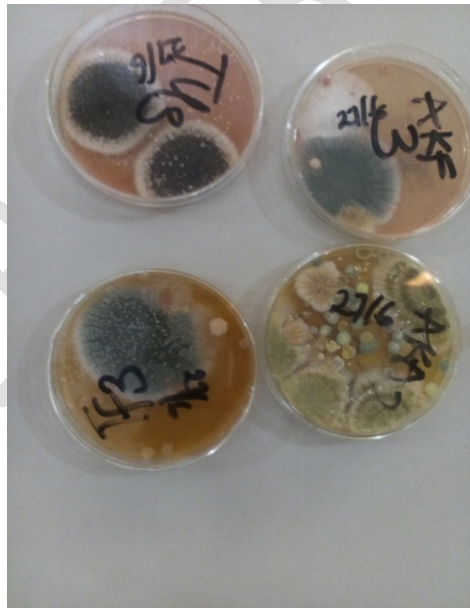


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Plate 3. Colonies of *Penicillium* spp. Morphological view

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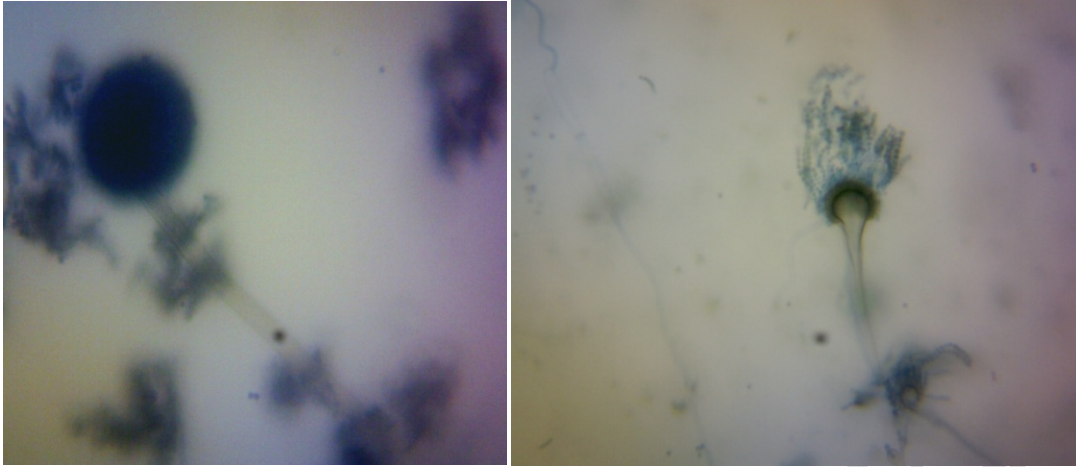


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Plate 4. Colonies of mixed fungi infection (Morphological view).

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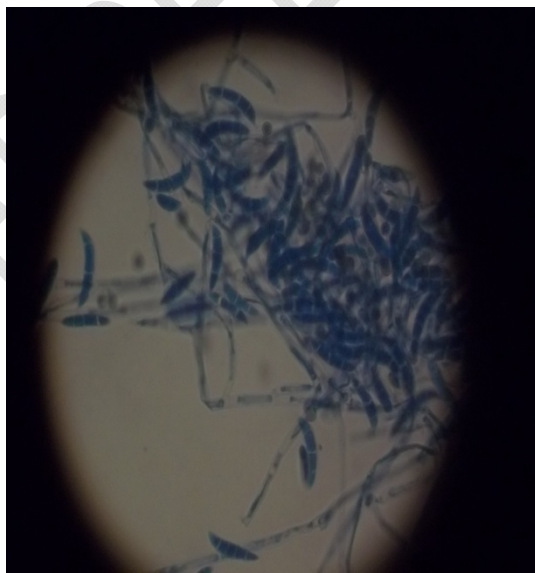
Figure 1. *Aspergillus* x100MG. **Figure 2.** *Aspergillus* view x100MG.

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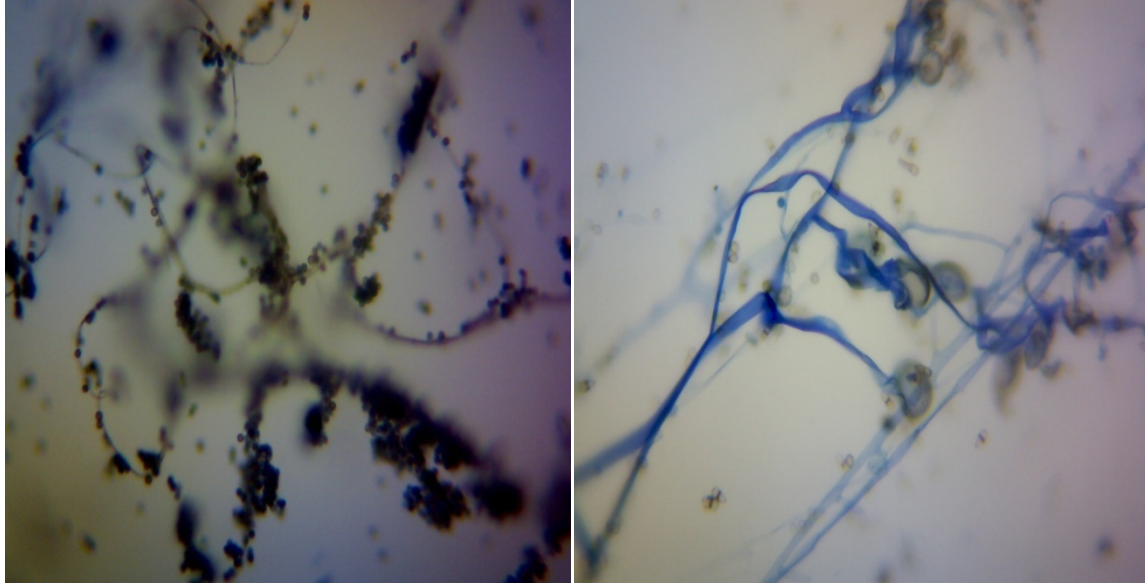


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Figure 3. *Fusarium* spp x100MG.

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Figure 4. *Penicillium* spp x100MG. **Figure 5.** Fungi Mycelium x100MG.

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Table 1. Fungi Genera Isolated From Some Selected Locations in Abia State

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Fungi organisms	Umuahia North	Osisioma	Farms (both)
<i>Aspergillus</i>	+	+	+
<i>Penicillium</i>	+	+	+
<i>Fusarium</i>	+	+	+
Yeast	–	+	+
<i>Mucor</i>	–	–	+

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Keys + (positive) – (Negative)

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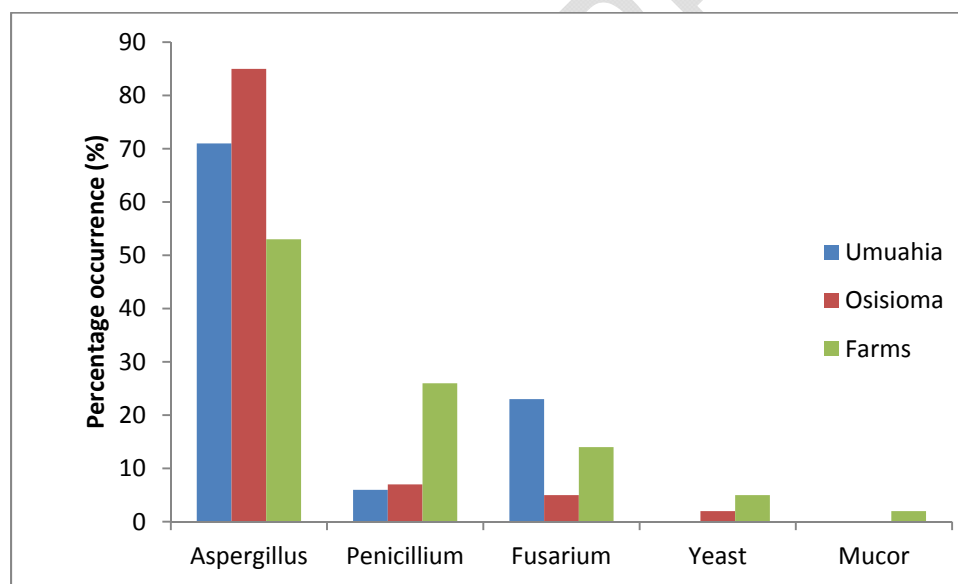
169 **Table 2.** Percentage and frequency of Fungi Contamination of The Feed Sampled
 170 From Various Locations.

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Locations	No of samples	No of samples contamination	Level of % contamination
Farms	40	34	85
Osioma	40	31	78
Umu North	40	20	50

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175 **Figure 6.** Percentage occurrence of Fungi organisms Isolated from 3 different
 176 locations in Abia State.

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Table 3. Total fungal load of feed sampled from each location

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	Locations	Total fungal count CFU/g-1
	Umuahia North	2.0x10 ⁵
	Osisioma	7x10 ⁵
	Farms	1x10 ⁶

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184 **DISCUSSION**

185 The study established that all the poultry feeds sampled harbored one fungi organism or
 186 the other. Most of these organisms found in the poultry feed are those commonly found
 187 in soil and water. The fungi isolated in this study were similar to those microorganisms
 188 reported by (Makun *et al.* [24]; Atehnkeng *et al.* [25]; Kpodo *et al.* [26]). Also from this
 189 result there is indication that feeds from farms has the highest percentage of fungal
 190 contamination of about 85% (Table 2) and this may be due to poor sanitary measures
 191 adopted in the processing and storage or due to poor environmental and personal
 192 hygiene practice in the farm as well as lack of proper biosecurity. *Aspergillus species*
 193 has the highest fungi percentage occurrence affecting most of the poultry feed sampled
 194 and this can be as a result of the organisms ability to thrive in high osmotic pressure
 195 and this is in agreement with Geiser *et al.* [27]. The spores are common component of
 196 aerosols and they drift by air current dispersing themselves both short and long
 197 distances. When these spores come in contact with solid feeds or liquid surfaces they
 198 tend to germinate in the presence of moisture as found by Gioconda and Richard, [28].
 199 From the study, *Aspergillus species* was the predominant organism isolated and this
 200 findings is in agreement with (Rosa *et al.* [29]; Oliveira *et al.* [30]; Figueora *et al.* [31]).

201 This research could not ascertain whether contamination occurred at the manufacturer
 202 level, retailers or farmers, though, several authors (Pitts and Hocking 1997; Monge *et al.*
 203 [32]) established that *Aspergillus* is predominant in cereals and other ingredient used in
 204 producing poultry feeds in the tropics. Contamination of poultry feeds particularly by

205 pathogen may occur prior to processing, distribution and or storage. Other studies have
206 similarly concluded that cereals and other ingredient use in producing poultry feed may
207 be source of product contamination. This does not exclude the fact that environment/
208 moist surface facilitate the growth of fungi. The occurrence of *Aspergillus*, *Penicillium*
209 and *Fusarium* spp could be due to absorption of moisture during storage [33]. The
210 stored poultry feed might have reabsorbed moisture from the environment which then
211 supported the growth of the microorganism in addition to the contamination during
212 processing.

213 The total fungal load in the analyzed finished feed samples in this study were about
214 1.9×10^6 cfu/g-1 which is higher than that reported in Slovakia, in 2003 of 1.9×10^3 cfu/g-
215 1) as reported by Magnoli, [34]. According to mycological quality criterion, good fungal
216 count should be less than 3×10^4 . [35]. The fungal load of poultry in this study was found
217 to be higher than the required load, hence the sampled poultry feeds are not good for
218 poultry consumption because they could lead to aflatoxicosis which results in reduction
219 of both production rate and meat quality. Also, from this result there is indication that
220 feeds from farms has the highest fungal count of about 1×10^6 (Table 4) and this may be
221 due to poor sanitary measures adopted in the processing and storage or due to poor
222 environmental and personal hygiene practice in the farm as well as lack of proper
223 biosecurity, followed by feed samples from Osisioma which have about 7×10^5 which
224 could be as a result of high stocking density. The feed samples from Umuahia has the
225 least fungal load of about 2.0×10^5 which may be due to good sanitary measures and low
226 stocking density adopted by feed distributors and depots in Umuahia.

227 The presence of fungi in the poultry feeds was analyzed using ANOVA of 95%
228 confidence interval and value $p < 0.05$ considered statistically significant. Also the post
229 hoc shows that there was a strong association between the presence of *Aspergillus* and
230 *Fusarium*, *Penicillium*, yeast and *Mucor*.

231 CONCLUSION

232 Since no vaccine exists for any of the fungal diseases of poultry therefore, the timely
233 adoption of good management practices, strict biosecurity, effective disease diagnosis

234 and suitable preventive measures along with necessary treatment like use of probiotics
235 with appropriate chemotherapeutic agents are good measures to have a check and
236 control the fungal disease of poultry apart from the fungal infection. *Aspergillus*,
237 *Fusarium*, *Penecillium* and *Mucor* were the main fungi isolated while yeast is a related
238 fungi organism. Mycotoxins are a major concern as they are the leading cause of
239 immune suppression in birds lowering their resistant level in viral and bacterial disease
240 and increase mortality. Thus a holistic approach is required to combat the adverse
241 effect on high economic returns from the poultry production. There is need for regular
242 surveillance and monitoring of important mycotoxins with the use of conventional as well
243 as modern diagnostic.

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UNDER PEER REVIEW