From the hygienic point of view, some bacteria like pathogenic serotypes of Escherichia coli produce serious disorders in animals and poultry, such as pathogenic serotypes of Escherichia coli which are common in poultry environment and causes some specific diseases such as coli-septicemia, coli-granuloma, air sac disease (chronic respiratory disease), avian cellulitis, peritonitis, salpingitis, swollen head syndrome; Gross (1994). Staphylococcus aureus which are known to be predominant in closed and open houses respectively in spring, Shigella Sp (66.6%) predominated in open houses in autumn. Water samples showed that Pseudomonas aeruginosa (43.48%) and Aspergillus flavus (46.51%) predominated in open houses in spring, Klebsiella oxytoca (50%) and Candida albicans (33.3%) predominated in close houses; E. coli (43.48%) and Aspergillus flavus (46.51%) predominated in open houses in spring, Klebsiella oxytoca (50%) and Candida albicans (33.3%) predominated in closed houses; Pseudomonas aeruginosa (47.06%) and Candida albicans (17.64%) predominated in open houses in spring, Klebsiella oxytoca (36.49%) and Aspergillus flavus (43.48%) predominated in closed samples; E. coli (43.48%) and Aspergillus flavus (46.51%) predominated in open houses in spring, Klebsiella oxytoca (36.49%) and Aspergillus flavus (43.48%) predominated in closed houses; E. coli (35.17%) and Aspergillus niger (35.14%) predominated in open houses in summer, Pseudomonas aeruginosa (31.75%); (42.31%) and Candida albicans (40.32%); (61.4%) predominated in closed and open houses respectively in autumn. Air samples showed that staphylococcus aureus (51.72%-45.45%, 52%-56.17%, 59.52-69.44 and 48.78-75%) was predominating in closed and open houses respectively in winter, spring, summer and autumn respectively, while the fungal growth showed that Aspergillus niger (66.6%); Aspergillus flavus (100%) predominated in closed houses in winter, Aspergillus niger (100%) predominated in open houses in spring, Aspergillus niger (100%) predominated in closed houses in summer, Aspergillus niger (100%) predominated in closed and open houses in autumn.

**INTRODUCTION**

From the hygienic point of view, some bacterial pathogens cause serious disorders in animals and poultry farms, such as pathogenic serotypes of Escherichia coli which are common in poultry environment and causes some specific diseases such as coli-septicemia, coli-granuloma, air sac disease (chronic respiratory disease), avian cellulitis, peritonitis, salpingitis, swollen head syndrome; Gross (1994). Staphylococcus aureus which are known to be
responsible for 30% losses in poultry by causing bacteraemia, endocarditis and yolk sac infection, Calnek et al. (1997). Aspergillosis (Brooder pneumonia) caused by Aspergillus niger causing high mortality in young chicks in the first three weeks of rearing about 10-50% of the affected chicks dies within 24 h. Also, Aspergillosis have been identified as the real cause of embryo mortality, (Gordon and Jordan, 1982) in addition to aflatoxicosis a condition caused by the toxins of Aspergillus flavus which usually occur in young pollets causing mortality which may reach to 50% (Gordon and Jordan, 1982). Penicillium species also causing serious damage among poultry farms due to the production of toxins may contaminate food (Marth, 1990). Candidiasis that caused by yeast like fungus Candida albicans and the disease which may affect birds from 1-2 weeks of age causing high mortality rate up to 75% (Gordon and Jordan, 1982).

In order to reduce human illness, epidemiological measures have been established because poultry and poultry products often serve as the vehicles for human Salmonellosis, (Bean and Griffin, 1990; Persson and Jendteg, 1992), the poultry industry and governmental agencies are focused on eradicating Salmonella in live birds and at the processing plant (Hargis et al., 2001). Salmonella Enterica serovar enteritidis is the world leading cause of salmonellosis and is often implicated in over 60% of human Salmonellosis in Europe, in the United States it remains the second most common serotype of Salmonellae, (Patrick et al., 2004). The current world wide epidemic of Salmonella enteritidis started in the middle of 1980s, (Ward et al., 2000), the (USDA) and Food Safety Inspection Services (FSIS) implemented an “in plant” Hazard Analysis Critical Control Point (HACCP) program to reduce the prevalence of the food borne pathogen contamination in meat and egg. Poultry are recognized as important reservoir of Salmonella; (Henson, 1997; Lynch et al., 2006), the United States Department of Agriculture implemented to reduce the source(s) of infection. Because food animal are the main reservoir of Salmonella enteritidis is mainly poultry often carrying asymptomatic infection, which pass the human pathogen along the food production chain. Especially undercooked or raw eggs and frozen poultry meat represent a high risk for humans.

This study was aiming for conducting an Epidemiological survey to identify the most frequently and persisting pathogenic microorganisms in the broiler’s environment. Culture, isolation and identification of pathogens from the environmental samples collected from broiler’s environment.

MATERIALS AND METHODS

Sampling: A total of 416 environmental samples were collected from two different systems of bird houses (Natural- artificial) each season (winter, spring, summer, and autumn). The samples were litter samples (William et al., 1975) Water samples (Lawrence, 1988), Swabs (Collins et al., 1991) that were collected from walls, floors, feed pads and drinkers, Air samples, (Cruickshank et al., 1975).

Bacterial and fungal culturing: The processing of the samples was carried out by pre-enrichment of all litter, water, swabs and air samples in 9 ml of Buffered Peptone Water (Oxoid, Fisher Int.) and incubated at 37°C for 18-24 h. The samples were then inoculated on non-selective quadrate plates of Blood Agar (Oxoid, Fisher Int.), Mac Conkey agar (Lab-M, Fisher Int.), Cystein Lactose Electrolyte Deficient Agar (CLED) (Lab-M, Fisher Sci.) and Sabaroud Dextrose Agar (SDA) (Oxoid, Fisher Sci.), the plates were streaked for isolation using a sterile cotton swab. The plates were then incubated at 37°C for 18-24 h.

Identification of bacterial isolates: The growing bacterial colonies were first used for gram staining for identification as Gram Negative Bacilli, (Edward and Ewing, 1982), Gram Positive Cocci, (Cherry et al., 1972; Cruickshank et al., 1975).

The colonies were then identified biochemically by inoculation onto TSI (Bialy and Scott, 1978; Edward and Ewing, 1982; Fingegold and Martin, 1982) SIM Citrate; Cruickshank et al. (1980), Indol test (Cruickshank et al., 1980), Urease Utilization test (Cruickshank et al., 1980), MR/VP test (Fingegold and Martin, 1982) Gelatin liquefaction test (Cruickshank et al., 1980), Potassium Cyanide (Cruickshank et al., 1980) for the gram negative bacteria. Meanwhile the gram positive tested against Catalase test (Cruickshank et al., 1980), Coagulate test and Deoxyribonuclease test (Blair et al., 1967).

Confirmation for the bacterial isolates: The colonies suspected to be E. coli were inoculated into 9 ml of Mac-Conkey broth, incubated at 44°C for 18-20 h. The tubes that revealed growth (gas and acid formation) were streaked again on Eosine Methylene Blue Agar (EMBA) and incubated at 37°C for 18-24 h. The positive colonies were these that have metallic green appearance. Biochemical identification was carried to insure the strain.

The colonies suspected to be Staphylococcus aureus: the samples were inoculated into 9 ml Phosphate Buffered Saline (PBS) and incubated for 18-24 h. The samples were then streaked onto Baird Parker Agar and the plates were incubated at 37°C for 20-24 h. The positive colonies showed black colonies on the agar.

The colonies suspected to be Salmonella Sp: Three to five colonies were inoculated into 9 ml of Selenite F broth. The tubes then incubated at 37 degree C for 18-24 h. A loopful was streaked onto S-S agar and the plates were incubated at 37°C for 18-24 h. The positive result for Salmonella appeared as black colonies with hallow zone.
Identification of the fungal and mycotic isolates: The growing fungal colonies were identified using the macroscopical examination of the mould colonies included the rapidity of growth, surface texture of the colony, surface color and reverse color, (Davies, 1976). Further identification for the fungal growth was carried out using the Microscopical examination, Part of the culture was removed with flame sterilized inoculation needle from the agar medium, placed on a slide and mounted into a drop of lacto phenol cotton blue and covered with a cover slip. The coverslip was pressed gently so as to lay the contents on the slide flat. The preparation were viewed under a low power microscope to reveal the desired structure then magnifying it as the conidial stage, head, vesicle, sterigmata, conidiophore and conidia, (Raper and Thom, 1949; Raper and Fennel, 1965; Zycha et al., 1969).

*Candida Albicans* was identified by using the germ tube test (Koneman et al., 1979) which based on the production of filamentous out growth in the form of pseudo-germ or pseudo-mycelium by cells of *Candida albicans* when inoculated into test tube containing 0.5 ml of Rabbit's Serum and incubated at 37°C for at least 3 h. A drop of the incubated yeast serum suspension was placed on a microscope slides, overlaid with a cover slip and examined for the presence of serum tubes known as germ tubes.

RESULTS AND DISCUSSION
Figure 1, the frequency distribution of bacterial isolates in the examined litter samples in different seasons showed that *Staphylococcus aureus* (40%) and *Pseudomonas areuginosa* (24%) predominated in closed houses, *Staph. aureus* (44%) *Klebsiella oxytoca* (12%), *E. coli* (12%), *Pseudomonas areuginosa* (12%), and *Proteus vulgaris* (12%) predominated in open houses in winter. *Klebsiella oxytoca* (33.33%), *E. coli* (26.67%) and *Pseudomonas areuginosa* (23.33%) predominated in closed houses, *E. coli* (50%), *Klebsiella oxytoca* (20%) and *Pseudomonas areuginosa* (15%) predominated in the open houses in spring. *Shigella Sp* (34.5%) and *Pseudomonas areuginosa* (20.69%) predominated in the closed houses, *Shigella Sp* (47.62%) and *Salmonella Sp* (19.02%) predominated in open houses in summer. *Klebsiella oxytoca* (55.56%) and *Pseudomonas areuginosa* (14.81%) predominated in closed houses, *Klebsiella oxytoca* (41.67%) and *E. coli* (29.2%) predominated in open houses in autumn.

Figure 2, the frequency distribution of bacterial isolates in the examined water samples in different seasons revealed that *E. coli* (39.47%), *Pseudomonas areuginosa* (26.32%) and *Proteus vulgaris* (21.05%) predominated in closed houses, *E. coli* (60.97%) and *Pseudomonas areuginosa* (29.27%) predominated in open houses in winter. *E. coli* (67.57%); (89.29%) and *Klebsiella oxytoca* (27.03%) (10.71) predominated in closed and open houses respectively in spring. *Shigella Sp* (42.55%), *E. coli* (21.3%) and *Pseudomonas areuginosa* (17.02%) predominated in closed houses, *Shigella Sp* (36.67%) and *E. coli* (30%) predominated in open houses in summer. *Klebsiella oxytoca* (36.59%), *Pseudomonas areuginosa* (24.39%) and *Proteus vulgaris* (19.44%) predominated in closed houses, *Klebsiella oxytoca* (47.22%) and *Proteus vulgaris* (19.44%) predominated in open houses in autumn.
Figure 3, the frequency distribution of bacterial isolates in the examined swab samples in different seasons revealed that *Pseudomonas aeruginosa* (62.5%), *Staphylococcus aureus* and *E. coli* (15.63%) predominated in closed houses, *Pseudomonas aeruginosa* (47.06%), *E. coli* (29.41%) and *Staphylococcus aureus* (19.61%) predominated in open houses in winter. *Klebsiella oxytoca* (36.49%), *E. coli* (27.03%) and *Pseudomonas aeruginosa* (24.32%) predominated in closed houses, *E. coli* (43.48%), *Klebsiella oxytoca* (28.26%) and *Pseudomonas aeruginosa* (21.74%) predominated in open houses in spring. *Klebsiella oxytoca* (28.98%), *Staphylococcus aureus* (28.99%) and *Pseudomonas aeruginosa* (23.19%) predominated in closed houses, *E. coli* (35.17%), *Staph. aureus* (28.57%) and *Pseudomonas aeruginosa* (17.86%) predominated in open houses in summer. *E. coli* (47.62%), (38.46%) and *Pseudomonas aeruginosa* (31.75%), (42.31%) predominated in closed and open houses respectively in autumn.

Figure 4, the frequency distribution of bacterial isolates in the examined air samples in different seasons revealed that *Staphylococcus aureus* (51.72%), (45.45%), *E. coli* (34.48%), (45.45%) and *Streptococcus Pneumoniae* (13.79%); (9.09%) predominated in closed and open houses respectively in winter, *Staphylococcus aureus* (52%), (56.17%), *E. coli* (40%) (23.33%) and *Streptococcus Pneumoniae* (8%); (20%) respectively in summer, *Staphylococcus aureus* (59.52%) (69.44%), *E. coli* (23.81%), (25%) and *Streptococcus Pneumoniae* (16.67%), (5.55%) respectively in summer, *Staphylococcus aureus* (48.78%), (75%), *E. coli* (36.59%), (13.89%) and *Streptococcus Pneumoniae* (14.63%) (11.11%) respectively in autumn.

Figure 5, the frequency distribution of fungal and mycotic isolates in the examined litter samples at different seasons revealed that Yeast (37.5%), *Pencillium Sp* (31.25%) and *Candida albicans* (18.75%) predominated in closed houses, Yeast, *Pencillium Sp* and *Candida albicans* by (28.57%) predominated in open houses in winter, *Aspergillus nidulans* (22.73%), *Pencillium spp.* (22.73%) and *Aspergillus niger* (18.18%) predominated in closed houses, Yeast, *Pencillium Sp.* and *Mucour* in a percentage of (21.4%) predominated in open houses.
in spring. Yeast (38.46%), Aspergillus niger (26.92%), and Mucour (19.23%) predominated in closed houses, Yeast (60.87%) and Aspergillus niger (8.69%) predominated in open houses in summer. Yeast (40%), Aspergillus niger and Candida albicans (12%) predominated in closed houses, Candida albicans (59.1%) and Yeast (9.09%) predominated in open houses in autumn. Figure 6, the frequency distribution of fungal and mycotic isolates in the examined water samples at different seasons.
Fig. 5: Frequency distribution of the fungal isolates in the examined litter samples at different seasons

Fig. 6: Frequency distribution of the fungal isolates in the examined water samples at different seasons
Fig. 7: Frequency distribution of the fungal isolates in the examined swab samples at different seasons

seasons showed that *Candida albicans* (50%), *Aspergillus flavus* (25%) and *Pencillium Sp* (25%) predominated in the closed houses, *Pencillium Sp* (60%), *Candida albicans* (20%) and *Aspergillus nidulans* (20%) predominated in open houses in winter. *Pencillium Sp* (37.5%) predominated in closed houses, *Pencillium Sp* (53.85%), *Aspergillus nidulans* and *Yeast* (15.38%) predominated in open houses in spring. *Yeast* (50%) (66.6%) and *Pencillium Sp* (37.5%) (22.2%) predominated in closed houses and open houses respectively in summer. *Candida albicans* (33.3%), (47.62%) and *Aspergillus niger* (25%) (19.05%) predominated in closed houses and open houses respectively in autumn.

Figure 7, the frequency distribution of fungal and mycotic isolates in the examined swab samples at different seasons showed that, *Yeast* (29.41%), *Pencillium Sp* (29.41%) and *Candida albicans* (17.64%) predominated in the closed houses, *Candida albicans* (41.67%) and *Yeast* (20.83%) predominated in open houses in winter. *Aspergillus flavus* (43.48) (46.51%) and *Aspergillus niger* (32.16%) (23.26%) predominated in closed houses and open houses respectively in spring. *Pencillium Sp* (34.48%) and *Aspergillus niger* (21.55%) in closed houses, *Aspergillus flavus* (35.14%) and *Pencillium Sp* (27.03%) predominated in open houses in summer. *Candida albicans* (40.32%) and *Aspergillus niger* (32.26%) predominated in closed houses, *Candida albicans* (61.4%) predominated in open houses in autumn.

Figure 8: Frequency distribution of the fungal isolates in the examined air samples at different seasons

*Aspergillus niger* (100%) predominated in closed houses, no fungal growth was detected in open houses. *Aspergillus niger* (100%) predominated in open houses in spring. *Aspergillus niger* (100%) predominated in closed houses, *Aspergillus flavus* (100%) predominated in open houses in summer.
predominated in closed houses, no fungal growth was detected in open houses. *Aspergillus niger* (100%) predominated in closed houses, *Aspergillus niger* (100%) predominated in open houses.

**Conclusion:** The results revealed that the microbial (bacterial and fungal) prevalence was varying form one type of sample to another (litter, water, swabs and air), from one season to another (winter, spring, summer and autumn) and from one housing system to another (naturally ventilated houses or open house system and artificially ventilated houses or closed house system). The bacterial prevalence generally was predominating in litter, swabs, air and water samples respectively in winter, spring and autumn. On the contrary; in summer, the highest prevalence for bacterial isolates was in swabs, air, water and litter. The fungal prevalence generally was predominating in swabs, air, water and litter samples respectively in winter, spring, summer and autumn. Also the results revealed that the highest rate of bacterial isolation in closed houses was carried out during autumn, summer, winter and spring respectively, and in open house system the highest isolation rate was carried out during winter, spring, summer and autumn. On the other hand highest rate of fungal isolation in closed houses was carried out during winter, summer, spring and autumn respectively and in open house system the highest isolation rate was carried out during autumn, spring, winter and summer respectively.

**REFERENCES**


