EXPERIMENTAL INFECTION OF ARCHANOBACTERIUM PYOGENES IN ONE DAY OLD CHICKS

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ABSTRACT

The present work aimed to study the pathogenicity of Archanobacterium pyogenes by using one day old chicks and chicken Embryonated eggs. The chicks were given the bacteria (4.2x10^8 CFU/ml) by orally administration and intraperitoneal injection, while the Embryonated eggs inoculated into allantoic sac at 12 days of incubation. The chicks were show dullness, decrease in food intake and nervous sings, most chicks were died after 2-3 days, and the bacteria were re-isolated from internal organs, and yolk sac. The inoculation of Embryonated chicken eggs resulting in embryonic death, reduce hatchery percentage and deformity of embryo. The study showed that A. pyogenes is virulence to both chicken Embryonated eggs and one day old chicks.

INTRODUCTION

Archanobacterium pyogenes is coccobacilli, non-motile, gram positive bacteria belong to genus Archanobacterium. It's usually found in upper respiratory, urogenital, and gastrointestinal tract in many domestic animals as opportunistic pathogen (1, 2). These bacteria were responsible for many pyogenic infections which affecting skin, joint or internal organs in animals and birds (2, 3). Its cause suppurative mastitis and liver abscesses in cattle (1, 4), cellulitis in chickens (5, 6) and osteomyelitis in turkeys (7, 8) septicemia, visceral lesions, cutaneous abscesses and decrease in egg production in chickens (9).

The disease caused by bacteria after an injury or other bacterial or viral infection (3, 10). The ability of bacteria to adhere, colonization, In vivo multiplication and then the damage of host cells result from several virulence factors which A. pyogenes owing. These factors include cholesterol-dependent cytolysin, pyolsin “PLO”, several proteases, several adhesive mechanisms involved collagen binding protein, fibrinogen binding and fibronectin-binding protein, DNase and neuraminidases which play important role in pathogenicity of bacteria against host immunity (1, 10).

Very little studies reported A. pyogenes as a cause of poultry disease, rather that bacteria should be considered as avian pathogen (5, 11). The objective of this study was to determine pathogenicity of A. pyogenes by using chicks and chick Embryonated eggs.
MATERIALS AND METHODS

Bacterial isolates: *A.pyogenes*(isolated from Quails) were obtained from lab. of microbiology/college of veterinary medicine – university of Mosul.

Prepare of inoculating dose:

The isolate of *A.pyogenes* was refreshed on 5% sheep blood agar. Five colonies from *A.pyogenes* were inoculated in Brain heart infusion broth” BHIB”, which incubate aerobically at 37°C for 24 hours. A serial dilution in brain heart infusion broth was done, then 0.1 ml from each dilution tube is inoculated on three nutrient agar plates and incubated aerobically at 37°C for 24 hours(12). Viable counts of bacteria were calculated and we selected the dose 4.2x10⁸ CFU/ml for chicks and egg inoculation(13).

Experimental study:

a- Egg inoculation: 30 fertile eggs was obtained from (Al-Ameen hatchery company) were used for inoculating of *A.pyogenes*. After cleaning and disinfection the shell of eggs they put in egg incubator for 12 days, the eggswereexamined weekly to remove the dead and ungrowing embryonated eggs. *A. pyogenes* (0.1 ml of 4.2x10⁸ CFU /ml) was inoculated in allantoic sac of Embryonated eggs at the 12 days old. The opening in egg shell is sealed by paraffin(12,14). Eggs were examined daily until the day 21 of incubation. The abnormal changes and death in embryo were recorded and bacteriological sample is taken to confirm isolation of *A.pyogenes*.

b- One day chicks inoculation:

Thirty chicks (one day old) were divided equally into 3 groups(G1,G2,G3) G1 contain 10 chicks injected with phosphate buffer saline as control. G2 and G3 contain 10 chicks received 0.1 ml from 4.2x10⁸ CFU/ml of *A.pyogenes*orally (G2) and intraperitoneally (G3)(15). The signs appeared in inoculated chicks were recorded. Dead chicks were taken to bacteriological examination.

c- *A.pyogenes* confirmation:

*A. pyogenes* confirmation was done by culturing of egg samples and internal organs (liver, intestine) and yolk sac of dead chicks on 5% sheep blood agar plates and incubate aerobically at 37°C for 24-48 h. (15,16). Colony which had typical cultural characteristic for *A.pyogenes*, was diagnosed according to the biochemical tests (catalase , oxidase , nitrate , gelatin hydrolysis , urease , and some sugar fermentation ) (16,17).
RESULTS

a- Embryonated Egg inoculation :

Most eggs Embryonated were dead, the hatchery percentage was 3.3% (1:30) most the embryos showed deformities in legs and yolk sac infection which appeared clearly in 5 days after inoculation. Some embryos were showed fibrinous material in yolk sac, and few embryo showed liver hemorrhage (Fig. 1)

![Fig 1: Twenty old day embryo: (A) show fibrinous material in yolk sac (B) leg deformity and yolk sac infection (same age)]](image)

b- One day old chicks inoculation :

The clinical signs after inoculation included dullness, decrease in food intake in groups (G2, G3). More sever clinical sings appeared in some chicks of intrapertoneal group (G3) including incoordination, unable to stand or walk normally and nervous sings developed after 24 hours post inoculation. Most chicks died after 2-3 days of inoculation.

At postmortem in chick group (G2) showed fibrinous membrane and attached in internal organs, the intestine is empty and therewas unabsorbed yolk with congestion in liver and mesenteric vessels in compare with control (fig. 2 :A). chicks in group (G3) showed unabsorbed yolk sac with congested in blood vessels, congestion and petechial hemorrhage in liver, intestine was empty, enlarged gall bladder also appeared in some chicks of this group (fig. 3 :A). Chicksgroup(G3) which suffered from nervous sings showed fibrinous exudates in mesenteric and fragile- pale liver also the intestine is empty and palemesentery( fig. 3 :C ).
Fig. 2: G2 group three Day old chicks : A, C show fibrinous membrane and attached in internal organs, the intestine is empty and there was unabsorbed yolk with congestion in liver and mesenteric vessels, B-Normal

Fig. 3: G3 group: Three Day old chicks “A”: unabsorbed yolk sac with congested in blood vessels, congestion and petechial hemorrhage in liver, intestine was empty, enlarged gall bladder, “B”: fibrinous exudates in mesenteric and fragile pale liver also the intestine is empty and pale mesentery, “C”: Normal
Bacteriological examination for dead chicks and embryonated eggs showed isolation of *A. Pyogenes* bacteria from internal organs of chicks and eggs (Table 1).

<table>
<thead>
<tr>
<th>Sample</th>
<th>Yolk sac</th>
<th>Liver</th>
<th>Internal organs</th>
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<tr>
<td>Embryonated Egg</td>
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<tr>
<td>G2</td>
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**DISCUSSION**

*A. pyogenes* is most widely reported from Archanobacterium species which cause various diseases in different animal species including poultry (18).

The results of our study in embryonated Eggs inoculation were showed high embryos dead, low hatchery percentage (3.3%) and deformities. This may result from infection in yolk sac (19) and septicemia caused by bacteria injected in eggs (9, 20). Also *A. pyogenes* contain extracellular matrix binding protein “collagen binding protein” CbpA which act as virulence factor to collagen rich tissue such as bone and cartilage and CbpA found in all osteomyelitis-*A. pyogenes* isolates (4, 8). This factor may be responsible for colonization of bacteria in cartilage and bone leading to deformity formation appeared in embryo. Clinical sings that appeared in one day chicks were differed from decrease in food intake and dullness (as a result from septicemia) (9), nervous sings like incoordination, unable to stand or walk normally which may be due to bacterial invasion of blood that may lead to form lesions and multiple abscessation in various organs (21), such as brain (22) which may result incerebral and intracranial abscessation with suppuration in central nervous system (23). These nervous signs occur meanly in young animals (24).

Production and severity of gross lesions based on challenge route and affected by immune reaction that differed in different part of lymphoid system (15). In our experiment liver was most affected organ by the two routes (Intrapertoneal & orally) of administration. Some chicks in both G2 and G3 group showed congestion in liver and mesenteric vessels which may result from inflammation occurred in liver after invading by bacteria and subsequent abscess formation (18), the *A. pyogenes* inflammation contributed by up-regulation of TNFexpression from macrophage under action of exotoxin “cholesterol- depending cytolysin, pyolysin” (PLO) (4). Some chicks in G3 were showed signs of fragile and pale liver with petechial hemorrhage due to distraction of cells by action of bacterial enzymes such as protease which may involve in host tissue damage and degraded host protein (4). Enlarged gall bladder appeared in chicks of G3 group was similar to that noted in *A. pyogenes* infection in turkey (7). The damage of red blood cells at liver my result from action of hemolysin effect of PLO exotoxin which able to cause lyses of red blood cells to produce B-hemolysis culture media (4).
Bacteriological re-isolation of *A. Pyogenes* from internal organs of chicks and embryonated eggs indicated that *A. Pyogenes* septicemia was occurred (20). There was high rate of isolation from liver rather than other organs and this may be result from bacterial tropism to hepatocyte, which has more abundant receptors to *A. Pyogenes* (15). Un absorbed yolk sac in chicks of G2, G3 was result from bacterial infection of the yolk and yolk stalk (omphalitis)(25).

This study concluded that *A. Pyogenes* is virulence to both chicken eggs and small chicks and the bacteria is one of the causes of embryonic death and reduce hatchery percentage in poultry industry, also is one bacterial species that cause death and unabsorbed yolk sac infection in chicks.

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