HYDROPERICARDIUM HEPATITIS SYNDROME IN CHICKENS

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ABSTRACT

Hydropericardium hepatitis syndrome is an acute infectious disease of broiler chickens characterized by high mortality, accumulation of fluids in the pericardial sac and necrosis of hepatocytes, during 1987 the virus was isolated from chickens in Angara near Karachi city in Pakistan and the disease takes it is name of this city. It is reported in China in 2015 in birds of 3-6 weeks in broiler chickens causing sudden death with high mortality reaches up to 75%. Avian adenovirus divided into 3 genrea: Atadodenovirus, Siadenovirus, and Aviadenovirus the last one divided into 5 species A to E subgroup1 strain 4 causing (Hydropericardium hepatitis syndrome). Subgroup 2 causing turkey (hemorrhagic enteritis, marble spleen disease). Subgroup 3 causing (Egg drop syndrome EDS). strain D and E produce inclusion body hepatitis while strain 1 causing (quill bronchitis). outbreaks of the disease producing economic losses for the poultry industry. Both vertical and horizontal transmission have an important role in the spreading of the disease. gross examination of the liver showing friable, enlargement, paleness of the liver. the kidney is swelling and hemorrhage with pale areas of necrosis in the parenchyma, pericardium is filled with clear or jelly fluids but histological changes showing hepatitis, nephritis and presence of basophilic intranuclear inclusion bodies. There are many tests can be used for the diagnosis such as PCR, ELISA, neutralization test. We concluded that the disease has economic importance due to high mortality and high losses of chicken’s meat production, so, biosecurity and vaccination are essential for preventing the infection.

INTRODUCTION

Avian adenovirus is a group of pathogens causing problems for poultry production and economic losses. The first isolation of the virus during 1949 the virus was propagated in Chicken embryo which taken from a case of lumpy skin disease. (Balamurugan and Kataria, 2004). FAV produce inclusion body hepatitis (IBH) in broiler chickens. IBH along with the hydropericardium syndrome. (HPS). Popularly called litchi heart disease in India (Hafez,2011). Isolation of the virus from a quail respiratory infections outbreak during 1950 and affects birds of 3 to 6 weeks old of broiler chicks (Abdul- Aziz and AL-Attar,1991). the disease was recorded in Iraq for the first time in 1991 in Mosul city (Danial et al.,2005).and the serotypes 2, 3 and 5 were isolates and diagnosed in the country in 1995. (Li ,2018). FAV nonenveloped double stranded DNA linear viruses, lie under the genus adenovirus in the family
adenoviridae the disease vertically transmission via the egg. Horizontal transmission through contaminated eggs and tools (Shamin et al., 2009). It may cause in immunosuppression leading to increase the susceptibility to secondary infections (Rahul et al., 2005). The symptoms and lesions vary according to serotypes of the virus as in hydropericardium syndrome, inclusion body hepatitis, egg drop syndrome, gizzard erosion, enteritis and poor growth (Domanska–Blicharz et al., 2011; Asthana et al., 2013; Lim et al., 2012). For detection of viral genome by using (PCR) (Dahiya et al., 2002). An outbreak during last years have been reported in different parts of India (Memon et al., 2006; Gowthaman et al., 2012; Senne et al., 1994)

ETIOLOGY:
The avian adenovirus is nonenveloped double-strand DNA under the genus Aviadenovirus in the family Adenoviridae viral genome have a number of nonstructural and structural proteins. The virion is composed of 252 capsomeres surrounded with a core 60-65nm. The capsomeres are attached to triangular surface and six capsomeres along with every edge, 240 hexons 8–9.5 nm diameter. And 12 vertex capsomeres (penton bases). Vertex carry projections known as fibers. the virus exhibits resistant to lipid solvents such as chloroform, ether, 50% alcohol, phenol 2%. They are resistant to variations in PH between 3 and 9 but inactivated when exposure for 30 minute to heat. (Hafez, 2011; Senne et al., 1994). The family adenoviridae was divided into two family: the genus mastadenovirus isolated from mammalian and aviadenovirus isolated from bird (Benkő et al., 2005). The genus aviadenovirus regarded as group 1 Aviano virus (AAV), Contains 11 of 12 european adenovirus serotypes, they classified into five molecular groups (A to E), the genus siadenovirus regarded as group 2, including hemorrhagic enteritis virus of turkeys (HEV). Adenovirus of avian spleenomegaly in the chickens (AASV). The genus Atadenovirus regarded as group 3 including egg syndrome virus (EDS) (Jensen and Villegas, 2005), group 4 regarded to cause hydropericardium syndrome (HPS). (Balamurugan and Kataria, 2004; Gowda and Satyanarayana, 1994).
Economic Significance:
In the present time it is difficult to detect the economic importance of the virus due to variability in disease so the increase in mortality and poor growth in flocks suffering from the disease results in economic losses (Balamurugan and Kataria, 2004).

Significance of public health:
There is no history of human infection by the virus, therefore the public health dangerous appeared minimal.

TRANSMISSION:
HPS is a contagious disease (Ganesh et al., 2002), transmitted vertically causing spreading of the disease so the virus can be detected in embryonating egg. Transmission of the disease horizontally is another way for spreading via contaminated food with infected feces which are taken orally (Toro et al., 2001).

PATHOGENCITY:
The incubation period 5-18 days, so the virus has the affinity for hepatic and endothelial cells (Akhtar,1992; Todd,2000). The ability of the virus to induce the disease is depend on the variations of different serotypes and the relationship between genotype and virulence not between virulence and serotype. Many factors associated with (HHS) play an important role to induce the disease such as: toxic fat agent mycotoxins, sodium chloride and the infection with the immunosuppressive diseases such as: Gumboro (IBD, and Reo virus infection (CAV) resulting in attenuating of immune response toward the disease (Shane and Jaffery,1997; Naeem et al.,1995; Deepak,1998). The (HPS) virus has affinity to affect the lymphatic tissue that result in immunosuppression (Aliev et al.,1997; Ganesh et al.,2001b). So, these diseases may be predisposing for HPS, when the virus enters the body initial multiplication occur in small and large intestine. Viraemia occur and spreading to different organs like kidney, liver, bursa, bone marrow and respiratory tract the virus can be isolated from these organs (Naeem et al.,2021; Ganesh et al.,2001).

SIGNS AND INCUBATION PERIOD:
The period of incubation of the disease is from 5-18 days with mean of 10 days (Akhtar,1992). Which is depend on the dose, age of the birds, route of infection, (Akhtar,1995). In experimental infection the incubation period is 48-72 hr (Aliev et al.,1997). Another study, the incubation period varied between 2-4 days (Ganesh et al.,2001b). The diseased birds have no appearance of specific symptoms but a sudden death was observed from 2-5 DPI (Anjum,1991; Afzal et al.,1991; Ganesh et al., 2002). The percentage of mortality is up to 80% occur in mature broiler flock of 3-5weeks of age (Jaffery,1988; Asrani et al.,1997; Voss et al.,1996) birds of sporadic cases may become dull and depressed in the terminal stages, huddle in corner, have a ruffled feather they showing different postures the beak and chest are resting on the ground, the eye lids were closed. there are no prominent signs been reported in (HHS) (Asrani et al.,1997).
GROSS LESIONS:

There is accumulation of fluids have a green color or colorless, watery to jelly-like fluid in the pericardium with range between 4-20 ml and PH of 7 in 90% of the affected birds (Ganesh et al., 2002; Gallina et al., 1973). Misshapen of the heart with floating in the apex of pericardial sac (Anjum et al., 1989; Kumar et al., 1997). There is a petechial hemorrhage and yellowish discoloration of cardiac fat. (Asrani et al., 1997). The liver changes include yellow paleness friable, swollen and large areas of mottled focal necrosis and ecchymotic or petechial hemorrhage. the lungs are congested. Urates are deposits in the ureters and tubules of kidney (Anjum et al., 1989; Abdul-Aziz and Hasan, 1995). An experimental study showing thymic and bursal atrophy (Asrani et al., 1997) in the pancreas there are a white pin point of necrosis and ventricular erosions in chickens (Nakamura et al., 2000; Palanivelu et al., 2014).

Picture2: A. enlarged liver with areas of paleness and congestion; B. clear fluids accumulated in the pericardial sac; C. Swelling of the kidneys with pale areas of necrosis and hemorrhage in the parenchyma; Oedema and enlargement in the bursa of Fabricius (Palanivelu et al., 2014).
MICROSCOPIC LESIONS:

The liver is the most important organ which many histological changes are detected such as infiltration of mononuclear inflammatory cells, focal areas of coagulative necrosis, presence of round or irregular basophilic intranuclear inclusion bodies. The hepatocyte is filling in the entire enlarged nucleus. (Anjum et al., 1989). necrosis and degeneration of the hepatocyte. (Abdul-Aziz and Hasan, 1995). The lung lesions showed edema and infiltration of inflammatory cells. Congestion of blood vessels (Asrani et al., 1997). There is hemorrhagic exudates in the alveoli and bronchi (Kumar et al., 1997). Infiltration of macrophage. (Nakamura et al., 2000). In the heart microscopic showed mononuclear cells infiltration, massive edema and sever vascular changes (Kumar et al.,1997), the kidney showed swelling of the tubular epithelium, increase in the glomerular area hemorrhage and necrosis in the parenchyma (Abdul-Aziz and Hasan,1995). Catarrhal infiltration of the mucosa of gastrointestinal tract specially in villi of intestine (Kumar et al., 1997), necrosis and atrophy of follicles and cyst formation in bursa of fabricius, and thymus. necrosis of spleen cells (Ganesh et al., 2002), Pancreatic necrosis of acinar cells (Nakamura et al., 2000).

DIAGNOSIS:

Before birds dying the diagnosis is difficult because there are no specific clinical signs and hardly in spontaneous out breaks (Kumar et al., 1997; Ganesh et al., 2001a). But it can be done in case of HPS-IBH according to gross lesions, histopathology represented by the intranuclear inclusion bodies in the cells of liver (Chandra et al., 1997 ; Kumar et al., 1997). the virus particles can be detected in the hepatic cells by using transmission electron microscope (Kataria et al., 1997; Jadhao et al., 1997). Cell
culture or inoculation in egg embryo can be used for isolation of the virus we can use the specimens from feces cecal tonsil, kidney, liver and heart for isolation (Kataria et al., 1997). For confirmation we can use neutralization test by using serotype- specific sera (Hassan et al., 1993). Other tests can be used for diagnosis such as indirect hemagglutination test, fluorescent antibody technique (Deepak, 1998).and virus modifications of ELISA (Balamurugan et al., 1999; Ganesh et al., 2002). PCR has also been developed for the diagnosis of HPS. Other confirmation of a virus by using electron microscope and immunocytochemistry to detect the infected cells with virus (Dahiya et al., 2002, Ganesh et al., 2002).

![Picture 4: virus particles are found in hepatocyte](image1)

![picture 5: PCR test appears 897 bp](image2)

PREVENTION AND CONTROL

Biosecurity one of the important aspects for prevention and controlling the disease (HPS). One of them by using of disinfection of equipment’s prevent the movement of the visitors inside the farm, proper lighting in the houses and proper ventilation (38,46). Another important aspect of prevention and control is the using of vaccination programs with killed vaccine which prepared from suspension of liver affected cells or FAdV-4 killed oil emulsion vaccine. produce high level of protection. breeders’ vaccination for 1-3 times between 9-12 weeks of age by alive vaccine consist of CAV and FAdV-4 induced high level of protection (Mashkoor et al., 1994; Kumar et al., 1997). While, a single dose of vaccine at 15-18 days of age (Anjum, 1990; Kumar et al., 1997). more effective for prevent the re infection at 35-40 days of age (Ahmad et al., 1990; Odisho et al., 1996).

CONCLUSION

We concluded that adeno virus infection in chickens causing a sever decrease in meat production due to high mortality resulting in poultry industry economic losses. so, biosecurity and vaccination are essential for prevent the infection, and avoided economic losses

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CONFLICT OF INTEREST
None of the authors has a financial or personal relationship with other people or organizations that could inappropriately influence or bias the content of the paper.

متلازمة موه القلب والتهاب الكبد في الدواجن
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الخلاصة:
متلازمة موه القلب والتهاب الكبد هي اصابة حادة لفروج اللحم تتميز بهلاكات عالية وتجمع للسوائل حول القلب مع تناخر لخلايا الكبد. في عام 1987 تم عزل الفيروس في مدينة انكارا القريبة من كراجي في باكستان، اذ تم تسمية المرض باسم هذه المدينة، كما تم تسجيله في الصين عام 2015 في افراخ اللحم بعمر 3-6 أسابيع، إذ يؤدي الى الموت مفاجئ مع نسبة هلاكات عالية تصل الى أكثر من 75% بما تم تسجيل المرض لأول مرة في العراق في العام 1991 في مدينة الموصل. اكتشفت الأنواع المصلية 2و5 في العام 1995 من قبل الدكتور مزاحم العطار وجماعته. صنف الفيروس الغداني الى ثلاثة اجناس وهي: فيروس أتا الغداني و سيا الغداني والطيري الغداني كما صنف الجنس الآخر الى خمسة فصائل من A-H وهي: تحت المجموعة 1 عتة 4 تسبب متلازمة موه القلب والتهاب الكبد تحت المجموعة2 تسبب التهاب الامعاء النزفي والطحال المرمي. تحت المجموعة الثالثة تسبب متلازمة انخفاض البالست كما ان العتة د و ه تسبب التهاب الكبد الاشتمالي بينما العتة D تسبب التهاب القصبات في السمان كما ادت الالدادات الوبائية الى حدوث خسائر اقتصادية في صناعة الدواجن ، إذ يلعب كل من الانتقال العمودي والافقي دورا مهما في انتقال العدوى وانتشار المرض. يبدو المظهر العياني للمرض بهشاشة الكبد مع تضخم وشحوب كما تظهر الكلى تورما مع نزف ومناطق نخريه شاحبة كما يظهر تجمع سوائل شفافة او جلتيني في شغاف القلب. أما الفحص النسجي يظهر وجود التهاب في كل من الكبد والكلى مع وجود اجسام اشتمالية نواتية كبيرة الحجم . يمكن تشخيص المرض باستخدام عدة اختبارات منها تفاعل البوليمرايز المتسلسل المتتر مناعي. اختبار التعامل نتج من ذلك ان للمرض اهمية اقتصادية مهمة لما يسببه من هلاكات عالية تسبب في خسائر في انتاج لحم الدجاج لذلك يعد الامن الحيوي والتلفيق من اهم الوسائط للسيطرة على المرض ومنعه.

الكلمات الدالة: فيروس الادينو,التهاب الكبد, دجاج, اجسام اشتمالية, موه القلب.

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