Pathology of Wild Strain (DP/As-Km/0019) of Duck Plague Virus Infection Revived in Ducklings

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ABSTRACT

Duck plague is an acute highly contagious disease of duck, geese and swan of all ages caused by Anatid Herpesvirus-1. The disease is characterized by significant decrease in growth, egg production and sudden death along with high morbidity and mortality which results into significant economic losses in duck industry. In present study, the pathology of wild strain (DP/As-Km/0019) of duck plague virus (DPV) was experimentally studied in 2 months old ducklings. The prominent clinical signs observed were depression, loss of appetite, greenish diarrhoea, ruffled feathers, nasal discharge, lacrimation and pasted eyelids with periorbital ring formation. The significant gross lesions were observed in spleen, oesophagus, liver, heart, brain and intestine. Vascular and degenerative changes like congestion, haemorrhages, necrosis and diphtheritic membrane formation were noticed in various parenchymatous organs. Microscopically, focal to diffuse areas of haemorrhages, coagulative necrosis, and fatty changes in liver, focal emphysema in lungs, formation of diphtheritic membrane on the mucosal surface along with presence of numerous infiltrating cells in oesophagus were recorded. Presence of Anatid Herpesvirus-1 viral DNA in liver and spleen tissue samples was detected by PCR.

Keywords: Pathology, Duckling, Assam, Duck Plague

Among the various types of livestock farming, duck farming plays an important role in the socio-economic development of the farmers. Duck farming is a highly profitable industry because of its high prolificacy and adaptability to free range system of rearing. As per 19th Livestock Census (2012), the total duck population of India is 23.54 million and Assam ranks first with 7.31 million ducks. Duck plague was first reported from West Bengal (Mukherji et al., 1963) in India.

Duck Plague is an acute highly contagious and infectious disease caused by Anatid herpes virus-1, a member of the family-Herpesviridae, subfamily-Alphaherpesvirinae. The disease is characterized by severe diarrhoea, sudden death, vascular damage, internal haemorrhages, lymphoid depletion, mucosal eruptions and degenerative lesions in almost all the parenchymatous organs (Dhama et al., 2017). Although mortality and morbidity depends upon the virulence of the virus and immunological status of the...
host (Goldberg et al., 1990), ducks of all age groups are considered susceptible with highest mortality in ducklings. Under natural conditions, ducks are usually infected through close contact to infected ones (Kaleta et al., 2007). The virus replicates principally in the epithelial cells and lymphocytes (Shawky, 2000; Yuan et al., 2005). Clinical signs like partially closed eyelids with photophobia, anorexia, extreme thirst, ataxia, oculo-nasal discharge, and greenish diarrhoea are characteristic to the disease (Dhama et al., 2017; Konch et al., 2009). Post-mortem lesions viz. petechiae over the epicardium, presence of the diphtheritic membrane over the esophageal folds and haemorrhages in intestine, spleen, liver, kidney, lungs, payer’s patches and bursa of Fabricius are consistent findings of the disease (Konch et al., 2009; Doley et al., 2013).

In spite of taking all important measures for preventing the disease, it is still prevalent in the country and threatening the existence of duck population and survivability of the duck farmers in particular. In present study, the pathology of wild strain (DP/As-KM/0019) of duck plague virus (DPV) was experimentally studied in 2 months old ducklings in order to understand the pathology and pathogenesis of the virus.

MATERIALS AND METHODS

This study was carried out in the Department of Pathology and Department of Microbiology, College of Veterinary Science, Assam Agricultural University, Khanapara, Guwahati.

Virus

DPV (DP/AS-KM/0019) positive tissues obtained from the repository of the Department of Microbiology were used for inoculation of ducklings in order to study the pathology of the virus.

Ducklings

A total of 12 ducklings of 2 months age were procured from backyard duck rearing areas of Assam. Ducklings were screened for presence of maternal antibody against DPV using indirect ELISA (I-ELISA). The ducklings were divided into 2 groups with 6 ducklings in each group, one group was used for inoculation and the other was kept as control. The ducklings were maintained using good managerial practice and under keen observation in the Department of Veterinary Pathology, College of Veterinary Science, Assam Agricultural University, Khanapara, Guwahati.

Screening of ducklings for DPV specific antibody by I-ELISA

The ducklings were screened for maternal antibody by I-ELISA as per the method described by Morrissy et al. (2004). A chequer board titration was performed for optimization of working dilution of DPV antigen and antibodies as per standard protocol.

Preparation of inoculum

PCR positive post-mortem samples (liver and spleen) obtained from repository of Department of Veterinary Microbiology was used to prepare the inoculum. The samples were grounded in a mortar-pestle into a 20% suspension (weight/volume) in sterile PBS. The suspension was then centrifuged at 3000 rpm for 10 minutes and the supernatant was collected, this step was repeated thrice. The final supernatant was subjected to cocktail antibiotic treatment for 45-60 minutes and used as inoculum.

Inoculation of DPV in ducklings

The inoculum was inoculated @ 1ml/duckling via oral (0.5 ml) and intramuscular (0.5 ml) route, whereas in control group sterile PBS was inoculated via same route and with same dose. Ducklings were observed for the development of post inoculation signs and were sacrificed on 7th day of post infection. Gross changes present in the different internal organs were recorded and simultaneously tissue samples from liver, spleen, kidney, oesophagus, lungs and heart were collected in 10% neutral buffered formalin for histopathology and on ice for molecular confirmation by PCR.

Molecular studies

Pooled post-mortem tissue samples were homogenized and 20% suspension was prepared by using appropriate volume of PBS (pH 7.4). Extraction of viral DNA from the inoculum, post-mortem tissues, tracheal and cloacal swabs
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preserved in HBSS and whole blood was done by Trizol method as per manufacturer’s recommendations. The primer pairs, F-5’ GAAGGCGGGTATGTAATGTA-3’ and R-5’ CAAGGCTCTATTCCGTAATG-3’ were used for amplification of DPV DNA (UL30 gene) (OIE, 2012).

**Histopathology**

Representative samples from all the organs were further subjected for the histopathological examination. Dehydration and clearing was done by using alcohol and xylene followed by wax impregnation and sectioning at 4μ thickness. Hematoxylin and eosin staining was done as per standard procedure (Luna, 1968).

**RESULTS AND DISCUSSION**

The ducklings used in the study were found free from maternal antibody. Presence of viral nucleic acid, in the inoculum used in the study and pooled tissue samples of experimental ducklings was detected by PCR. The yield of DNA was ranged between 85 to 750 µg/ml. Test spleen and liver samples showed the amplicon size of 446 bp and no amplification was observed in positive control group (Fig. 1).

Pathology of DPV in ducklings was previously reported by Bhowmik and Chakrabarty (1985) and El-Samadony et al. (2013). They reported that duckling was unable to stand with anoraxia, diarrhea and death. The gross lesions are characterized by vascular damage with tissue haemorrhages. Lesions of the lymphoid tissues were more prominent than visceral haemorrhages. Present study revealed good revival of wild strain of DPV in ducklings inoculated with PCR positive tissue samples which indicates well adaptation of virus to primary host. In order to recover DPV isolate from field tissue samples, duckling can be an ideal in vivo host. In context to this, Sarmah (2016) found similar result with revival of the wild strain of DPV in the primary host.

The infected ducklings showed clinical signs between 4 to 7 days of post inoculation (dpi). The clinical signs were noticed as early as 3 dpi onwards. The common clinical signs observed in most of the ailing ducklings were depression, lethargy, huddling, unapparent to the environment, reduced appetite followed by anorexia noticed from 4 dpi, increased thirst, nasal discharge, ruffled feathers, periorbital ring formation with lacrimation, pasting of feathers around the eyes, greenish diarrhea with soiled vent, ataxia and dropping of wings (Fig. 2). Our findings were closely related with the findings of Konch et al. (2009) and Sahariah (2016).

**Fig. 1:** PCR products in ethidium bromide stained 1.7% agarose gel electrophoresis. **L1:** Inoculum used for the experiment; **L2:** Pooled spleen samples of infected ducklings; **L3:** Pooled liver samples of infected ducklings; **L4:** Positive control; **L5:** 1 kb DNA ladder and **L6:** Negative Control.

**Fig. 2:** Clinical signs showing by duck plaque infected ducklings. **A:** Greenish diarrhoea of ailing duck; **B:** Duckling infected with DPV exhibit periorbital ring formation with lacrimation; **C:** Duckling showing paralysis of left limb; **D:** Catarhal nasal discharge showing by infected ducklings.
The most characteristic gross lesion observed was deposition of yellowish diphtheritic membrane along the longitudinal folds of oesophagus. The lesion was focal and limited to certain areas of oesophagus moreover, removal of the plaques did not reveal any haemorrhagic areas. Moderate catarrhal enteritis with absence of gross lesions in annular band was evident. Grossly, almost all affected birds showed - hepatomegaly, focal petechial hemorrhages over the liver parenchyma and gall bladder distended with bile. Heart showed congestion of blood vessels along with a few focal petechial haemorrhages over the epicardium. In kidneys, diffuse petechial haemorrhages with few necrotic foci over the parenchyma were observed. Lung parenchyma revealed emphysema associated with diffuse haemorrhage. The spleen was dark in colour and focal areas of haemorrhages were evident over the parenchyma, however splenomegaly with severe congestion was the regular finding in some of the infected birds. The mucosa of the trachea was congested with mild haemorrhages. Brain revealed mild to moderate congestion of meningeal blood vessels. (Fig. 3). Gross lesions were mainly confined to spleen, oesophagus, liver, heart, lungs, kidney, trachea and brain. These above lesions were in accordance with the findings of Dhama et al. (2017).

Microscopically, sloughing of diphtheritic membrane from the mucosal surface and infiltration of numerous mononuclear cells were evident in the oesophagus. Liver showed haemorrhages, disintegration and coagulative necrosis of hepatic cords along with necrosis of hepatocytes. In heart, haemorrhages between the muscle fibers were observed. Kidneys were characterized by congestion and haemorrhages in interstitial space, shrinkage of glomerular tuft, increased Bowman’s space, coagulative necrosis and desquamation of renal tubular epithelium. In lungs, emphysema associated with diffuse haemorrhages was observed. Depletion of lymphocytes in lymphoid follicles and diffuse areas of haemorrhages were observed in spleen (Fig. 4). Histopathologically, most consistent lesions were observed in liver, spleen, oesophagus, heart, lungs and kidneys in the present study and similar findings were also reported by Shawky (2000) and Sahariah (2016).

CONCLUSION

The present study, concluded that wild strain of DPV present in post-mortem tissue samples could be revived in the primary host. Periorbital ring formation was the constant clinical finding in experimentally infected ducklings. Formation of diphtheritic membrane in the
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-path inhibit in the oesophagus, haemorrhages along with necrosis in liver and depletion of lymphocytes in splenic follicles are the major gross pathological findings. Microscopically, the most prominent changes were characterized by congestion, haemorrhages, degeneration and necrosis of the parenchymatous organs.

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