Avian Polyomavirus Infection in Non-Budgerigar Psittacine
Birds in Thailand - A case report

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Abstract

One sick 4-week-old Eclectus (Eclectus roratus) and two carcasses of 3-week-old Macaws (Ara macao) from a
bird farm near Bangkok were submitted in July 2009 for investigation. Necropsy revealed multifocal hemorrhages at
subcutaneous tissue, heart, liver and bursa of Fabricius. The most prominent findings were hepatomegaly and
splenomegaly. Histopathologic examination showed severe splenic necrosis with presence of homogeneous pale
basophilic intranuclear inclusion body in splenic periarteriolar sheath cells. The liver showed severe necrosis with
occasionally intranuclear inclusion body in sinusoid-lining cells. Intranuclear inclusion bodies were detected in
mesangial cells, epithelium of collecting tubules and papillary ducts of kidney and mesenchymal cells of lung and
heart. TEM examination revealed tremendous 45 nm icosahedral non-enveloped capsids consistent with virions of
polyomavirus in intranuclear inclusion body containing cells in kidney and lung. The diagnosis of avian
polyomavirus (APV) infection was made based on histopathologic and electron microscopic results. Avian
polyomavirus cause severe disease in a wide range of psittacine and non-psittacine birds. It occurs most frequently in
budgerigars, macaws, conures, Eclectus parrots, Ringneck parrots and lovebirds. The disease is more severe in
nestlings as also observed in this study. In this report, we document the macroscopic and microscopic findings
including TEM feature of a recent case of APV infection in non-budgerigar psittacine birds from a bird farm near
Bangkok, Thailand. This report may remind avian veterinarians and breeding aviary farmers about the occurrence
of APV infection in this country and make them be aware of the possible outbreak of APV in the future.

Keywords: avian, electron microscopy, histopathology, polyomavirus, psittacine

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Introduction

Avian polyomavirus (APV) was first known as pathogen in young budgerigars (Melopsittacus undulatus) in the early 1980s (Davis et al., 1981) and was recognized as the causative agent of the budgerigar fledgling disease (BFD) at that time. The virus causes severe disease in a wide range of psittacine and non-psittacine birds. APV has been recognized in a variety of avian species like non-budgerian psittaciformes, falconiformes, domestic geese as well as finches and gallinaceous birds. The infection is mostly severe and fatal in young birds (Bernier et al., 1981; Pass et al., 1987; Rahaus and Wolff, 2005). In fledgling budgerigars, mortality rate may reach 100% (Krautwald et al., 1989). Clinical signs observed in acute infections are polyuria, subcutaneous hemorrhages, dyspnoea and depression (Graham and Calnek, 1987; Stoll et al., 1993). APV has been placed into the distinct subgenus Avipolyomavirus. The non-enveloped capsids of APV virions are icosahedral shape and about 45-50 nm in size (Bozeman et al., 1981; Pass et al., 1987). The genome is a circular and double-stranded DNA of 4981 bp in length, which is fully sequenced and molecular characterization of isolated APV based on the whole genome sequence analysis were documented (Katoh et al., 2009; Rott et al., 1988). An outbreak of APV infection in Thailand was reported in 2005 and the diagnosis was confirmed by PCR (Banlunara et al., 2005). Predominant macroscopic lesions were hepatomegaly with focal hemorrhage, splenomegaly, hydropericardium, hydroperitoneum and patchy hemorrhage in the subcutaneous tissue. Histopathology showed hepatic necrosis with karyomegalic fine granular basophilic intranuclear inclusion bodies in hepatocytes, Kupffer’s cells, cells in perivascular sheath of spleen, in lamina propria of intestines, glomerular mesangial cells, renal tubular cells and feather follicular cells (Banlunara et al., 2005).

The aim of this study is to present gross, microscopic and ultrastructural features of a recent case of APV infection in non-budgerigar psittacine birds in Thailand.
Materials and Methods

Case history: One sick 4-week-old Eclectus (Eclectus roratus) and two carcasses of 3-week-old Macaws (Ara macao) from a bird farm near Bangkok were submitted in July 2009 to Mahanakorn Veterinary Diagnostic Center (MVDC) for investigation. The animals showed history of severe and acute onset of illness and were found dead. Necropsy was performed on the two carcasses and tissue samples were fixed in 10% buffered formalin. The tissue samples from three birds were sent to histopathology laboratory for slide preparation using conventional method. Briefly, the tissues were processed and embedded in paraffin, sectioned at 4 microns and stained with hematoxylin and eosin (H&E). All slides were examined under light microscope (Axiolab, Zeiss, Germany). Some pieces of organ including kidney, liver and lung were re-collected from achieve formalin fixed material for further transmission electron microscopic (TEM) study at Pathology Section, the National Institute of Animal Health (NIAH). Briefly, the tissue samples were post-fixed in 1% osmium tetroxide and embedded in epoxy resin (Epon). Semithin sections were cut at 0.5 micron and stained with 2% toluidine blue. The sections were examined under light microscope to locate specific areas for ultrathin sections. The ultrathin sections were cut and stained with 5% uranyl acetate and lead citrate. The samples were examined under transmission electron microscope (JEM-1200EX, JEOL, Japan).

Results and Discussion

Necropsy result: The carcass examination revealed multifocal patchy hemorrhages at subcutaneous tissue. The most prominent gross pathological findings were hepatomegaly (Figure 1) and splenomegaly with multiple white foci. Multifocal ecchymotic hemorrhages were detected at epicardium of heart (Figure 2), liver and bursa of Fabricius. The kidneys were pale and the lungs were congested and edematous. Other organs showed non remarkable macroscopic finding.

Histopathological result: The most pronounce microscopic findings were found in spleen and liver. Splenic lesions composed of red pulp congestion and severe periarteriolar sheath necrosis. Karyomegaly, margination of nuclear chromatin and homogeneous pale basophilic intranuclear inclusion body were observed in many splenic periarteriolar sheath cells (Figures 3 & 4). Homogeneous pale basophilic intranuclear inclusion bodies were commonly detected in glomerular mesangial cells (Figure 5), epithelium of collecting tubules and papillary ducts of the kidney and mesenchymal cells of lung and heart (Figures 6a & 6b). The liver showed severe congestion with multifocal randomly hepatic necrosis and hemorrhages. Occasionally, homogeneous pale basophilic intranuclear inclusion body was found in sinusoid-lining cells (Figure 7). Bursa of Fabricius showed markedly medullary lymphoid necrosis without detectable cytoplasmic and nuclear inclusion bodies. Other organs showed non-remarkable microscopic lesion.
Figure 3. Spleen showing necrosis at periarteriolar sheath with presence of basophilic intranuclear inclusion bodies (black arrows)

Figure 4. Higher magnification of spleen from figure 3, pale homogenous basophilic intranuclear inclusion bodies found in karyomegalic nuclei (black arrows) of periarteriolar sheath cells of spleen

Figure 5. Basophilic intranuclear inclusion bodies seen in karyomegalic nuclei of mesangial cells (black arrows) within renal glomeruli, T: renal tubule

Figure 6a-6b. Intranuclear inclusion bodies in karyomegalic nuclei (black arrows) in lung and heart

Figure 7. Focal hemorrhage in liver with presence of intranuclear inclusion body in nucleus (black arrow) of sinusoidal lining cell

Transmission electron microscopic (TEM) result: On semithin sections of kidney, liver and lung, intranuclear inclusion bodies could be observed as previously mentioned in histopathological findings. Within the affected nuclei, karyomegaly with margination of nuclear chromatin were found. Tremendous 45 nm icosahedral, non-enveloped capsids consistent with virions of polyomavirus were detected in intranuclear inclusion body containing cells in kidney and lung (Figure 8). In the kidney, some virions of polyomavirus were also recognized in cytoplasm of collecting duct epithelium. There was no virion of polyomavirus observed in inclusion body containing cells in liver in this study. Based on shape
and size of virions and host range of the virus, it was classified to avian polyomavirus (APV) which belong to family Papovaviridae and genus Polyomavirus.

Morphological diagnosis: Avian polyomavirus (APV) infection was diagnosed based on histopathology and electron microscopy.

Discussion: Avian polyomavirus infection occurs most frequently in budgerigars, macaws, conures, Eclectus parrots, Ringneck parrots and lovebirds. The disease is more severe in nestlings. In this study, the affected birds were macaws and Eclectus parrot and all of them were nestlings. All three non-budgerigar psittacine birds died rapidly and the most prominent external findings were subcutaneous hemorrhages and edema as reported by many authors (Banlunara et al., 2005; Bernier et al., 1981; Davies, 2000; Garcia et al., 1993; Pass et al., 1987). Petechial and ecchymotic hemorrhages are also the hallmark of many cases of acute death from APV as also observed in this study. Presumptive diagnosis of APV infection can be based on history, clinical findings, and characteristic gross and histopathologic lesions (Raidal, 1995). The presence of karyomegaly with margination of nuclear chromatin and homogenous pale basophilic intranuclear inclusion bodies are pathognomonic lesions of APV infection (Bernier et al., 1981; Pass et al., 1987). In this study, intranuclear inclusion bodies were most frequently found at periarteriolar sheath cells of spleen and glomerular mesangial cells of kidney as also found in budgerigars in Thailand previously reported (Banlunara et al., 2005). In the literatures, inclusion bodies were observed in both epithelial and mesenchymal cells (Pass et al., 1987; Garcia et al., 1993). TEM is used to confirm the presence of polyomavirus virions within the nuclear inclusion body. Polyomavirus virions were detected both in the nucleus and cytoplasm of collecting duct epithelial cells in kidney and in the nucleus of mesenchymal cells of lung. The presence of virions of polyomavirus in cytoplasm of collecting duct epithelial cells is compatible with the study of Pass and colleagues (1987). The exact origin of some inclusion body containing cells was difficult to identify both histopathologically and electron microscopically as also mentioned by same authors. This may due to small amount of cytoplasm of affected cells and alteration of cell morphology from the viral infection. However, the cell position within the affected tissue could give some hints of suggestion about cell origin. Diagnoses of chronic or subclinical cases require molecular technique. Although DNA in situ hybridisation (Garcia et al., 1994; Latimer et al., 1994; Ramis et al., 1994) and polymerase chain reaction (PCR) technique (Banlunara et al., 2005; Hsu et al., 2006; Roy et al., 2004; Tomasek et al., 2008) had recently become the most common and standard methods to confirm APV infection, a conventional tool such as TEM examination is still be useful for diagnostic confirmation especially in laboratories where TEM is available and routinely performed.

In this study, we document the macroscopic and microscopic findings including TEM feature of a recent case of avian polyomavirus infection in non-budgerigar psittacine birds from a bird farm near Bangkok, Thailand. This report may remind avian veterinarians and breeding aviary farmers about the occurrence of APV infection in this country and make them be aware of the possible outbreak of APV in the future.

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