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VIRAL INCLUSIONS IN RACCOON LIVER CELLS


Abstract: Three young raccoons (Procyon lotor), two from Michigan and one from Arizona, died suddenly from acute infections. Intranuclear inclusion bodies and viral particles typical of herpesvirus were seen in liver cells from all three. Inclusions also were seen in the nuclei of endothelial cells in the lung, liver, glomeruli and reticuloendothelial cells of the spleen. The source of the infection was not determined, but possible transmission from other species could not be ruled out.

INTRODUCTION

Raccoons (Procyon lotor) are naturally or experimentally susceptible to a number of different viral diseases. These include: canine distemper (paramyxovirus), infectious canine hepatitis (adenovirus), Eastern, Venezuelan and Western equine encephalomyelitis (togavirusess), pox virus and pseudorabies (herpesvirus), rabies and vesicular stomatitis (rhabdovirus), feline enteritis (parvovirus), and encephalomyocardial disease (picornavirus). This report describes a possible new viral disease in raccoons.

Case 1

Two immature raccoons were picked up along the highway at different times during the summer 1971 in western Michigan. They were taken to an animal orphanage to be nursed back to health before being released into the wild. At the orphanage they were caged with a tame, healthy kinkajou (Potos flavus). Five weeks after being placed in the cage one raccoon developed anorexia and was inactive. A rectal temperature of 40°C was recorded. The animal died two days later during a convulsion.

Necropsy revealed necrosis of the skin of the feet and slight enlargement of the spleen, iliac and lumbar lymph nodes. Babesia-like organisms were seen in erythrocytes on a stain blood smear. No pathogenic microorganisms were cultured from the internal organs.

Microscopically, the lesions included necrosis of germinal centers in the spleen and lymph nodes and focal necrosis throughout the liver. In addition, large basophilic intranuclear inclusions were seen in hepatic cells (Fig. 1), reticuloendothelial cells of the spleen and lymph nodes, and in capillary endothelial cells in the lungs and liver. The dense inclusions were large and numerous and usually filled the nucleus, although sometimes a halo was apparent. Their density obscured internal detail. Hepatic cells containing the inclusions were somewhat swollen and projected into the sinusoidal spaces.

The second animal died suddenly. Necropsy revealed hemorrhages in the skin, oral and gastric mucous membranes, and the cerebrum and cerebellum. Focal areas of necrosis were visible on the liver surface. The spleen was enlarged, and Babesia-like organisms were...
FIGURE 1. A dense intranuclear inclusion body in a hepatic cell from a young raccoon. H & E stain. ×4100.

seen in erythrocytes on a stained blood smear. No pathogenic microorganisms were cultured from internal organs.

Microscopically, necrosis and hemorrhage were observed throughout the liver, lymph nodes and spleen. Generally, the necrotic areas in the liver were adjacent to the central veins. Large basophilic intranuclear inclusion bodies were seen in many hepatic cells and capillary endothelial cells in the kidney.

Case 2

A weaned female raccoon was purchased from a pet store in Arizona. At 10 weeks of age, she developed clinical signs of anorexia, malaise and diarrhea. Forty-eight hrs. later she was presented at an animal clinic. At this time, the visible mucous membranes were pale and the body temperature was subnormal. A tentative diagnosis of feline distemper was made. Fluids, corticosteroids and antibiotics were administered, but the animal died six hrs. later.

Necropsy examination revealed swollen kidneys, enlarged mesenteric lymph nodes and a small spleen. Microscopically, intranuclear inclusion bodies were seen in endothelial cells of the renal glomerular tufts, in hepatocytes, and in the reticuloendothelial cells of the spleen and lymph nodes. These inclusions were indistinguishable from those seen in the other two animals. Hemorrhage was seen in the lymph nodes, and necrosis of lymphocytes had occurred.

Only formalin-fixed tissue was available for transmission electron microscopic examination (TEM). Nevertheless, TEM examination of inclusion bodies in hepatic cells from all three animals revealed many viral particles typical of herpesvirus (Fig. 2). With rare exception, all the viral particles were within the nucleus.

Tissue sections from five other raccoons submitted to this laboratory for a variety of reasons were examined for inclusion bodies, but none was found.

FIGURE 2. Viral particles typical of herpesvirus in the nucleus of a hepatic cell. Lead citrate, uranyl acetate stain. ×91000.

DISCUSSION

The fact that no pathogenic microorganisms were isolated from the three raccoons strongly suggests that the
herpesvirus observed in intranuclear inclusions was the causative agent of this acute and fatal disease. The lesions, particularly the large intranuclear inclusions, were strikingly similar to lesions seen in raccoons infected with infectious canine hepatitis (ICH), an adenovirus.3 Although no description of such a herpesvirus disease in raccoons was found in the literature, J. King (pers. comm.) stated that he had seen inclusion bodies in a raccoon typical of ICH, but upon TEM examination, they contained herpesvirus. Similar acute hepatic diseases caused by herpesviruses have been reported in chinchillas and a variety of birds.6,11,1,21 It is interesting to note that two of the raccoons were in contact with a kinkajou and the third was purchased from a pet store. Thus, it can be speculated that cross-species transmission occurred. Alternatively, it is possible that the herpesvirus was latent in the raccoons, and the stresses incidental to captivity activated the disease. In any event, herpesvirus “inclusion body hepatitis” may represent a new disease which must be considered when dealing with ICH-like lesions in raccoons. It is unfortunate that the virus was not isolated (no attempt was made) because validation of the disease will require characterization of the virus and pathogenicity studies.

LITERATURE CITED

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