Identification of a novel herpesvirus associated with cutaneous ulcers

in a fisher (*Martes pennanti*)

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Abstract. On December 8th, 2008, a male fisher (Martes pennanti) housed in a quarantine enclosure at the St-Félicien Zoo was found dead with multiple skin ulcers on the muzzle and plantar pads. At necropsy, no major findings were found, and a specific cause of death was not determined microscopically. However, at the borders of ulcerated sites, there were increased numbers of koilocytes, with perinuclear vacuolation and nuclear enlargement. A pan-herpesvirus nested-polymerase chain reaction (PCR) assay was conducted, and an expected PCR product of 230 nucleotides was obtained within tissues collected from around the skin ulcers. Other tissues, including intestines and pool of lung, liver and kidney, tested negative. The obtained PCR amplicon was sequenced and was highly related to the partial viral DNA polymerase (DPOL) gene of a Badger herpesvirus. Virus isolation was negative, and no virion was detected by electron microscopy. The pathogenic potential of this novel herpesvirus and its role in the death of the fisher are unknown.

Key words: fisher, FiHV, herpesvirus, Martes pennanti, skin, ulcer.
The fisher (*Martes pennanti*) is a fur animal of the size of a domestic cat with a long tail and short legs, classified in the *Mustelidae* family. As a generalist predator, the fisher’s diet consists predominantly of North American porcupines, snowshoe hares, squirrels, small mammals, birds, carrion, and, to a lesser extent, fruits and plants. Fishers have been reported in the forest regions of several states of the United States of America (USA) and provinces of Canada.

On October 2nd 2008, two couples of captive born fishers, each aged 2½-years, were bought from a breeding farm in Minnesota, USA, and sent to the St-Félicien Zoo, Quebec, Canada. While in quarantine, a physical exam was performed under anesthesia 6 days after their arrival. Hematology and biochemistry blood tests were normal except for a marginal uremia of 16.9 mmol/L [reference values: 3.57-12.14 mmol/L] detected in one of the males. Fecal and radiographic exams were performed and no significant clinical findings were recognized.

On December 8th 2008, the male fisher that was previously found marginally uremic on October 2008, was found dead in its quarantine enclosure with multiple skin ulcers on the muzzle and plantar pads (Fig. 1). A variety of causes could induced ulcerative skin lesions but with the gross lesions of the muzzle and plantar pads, *Feline calicivirus* (FCV), and *Feline herpesvirus* (FHV), *Canine distemper virus* (CDV) as well as frostbite were considered as possible causes. The three other healthy fishers had shown no clinical signs of illness.

On gross examination, the fisher, which had lost 3.7 pounds in body weight, had no fat in the body cavities or surrounding the organs. There were patchy losses of hair on both sides of the flanks and on the hips. The most remarkable lesions were the sharply demarcated and deep, round to oval ulcerations of approximately 0.5 to 1.0 cm in size located on all paws and footpads, and on the muzzle (Fig. 1). Selected few grams of tissues (brain, lung, heart, liver, kidney, lymph
nodes, spleen, stomach, bladder, intestine, pancreas, adrenals, thyroids, muscles and skin) were taken for light microscopic evaluation examination, toxicology, a bacteriology, b immunohistochemistry, c electron microscopy examination, c virus isolation, and for polymerase chain reaction (PCR) assays.

A specific cause of death was not determined microscopically. Noteworthy, endocrine dermatosis might be considered to explain the marginal alopecia. Thus, sections of the skin with alopecia, adrenal and thyroid glands were examined. Thickness of the epidermis varied from 4-5 cells thick, which might be normal, to 1-2 cells thick, suggestive of epidermal atrophy. In these regions, the hair follicles appeared atrophied. No cortical hyperplasia or hypoplasia, neither nodular hyperplasia were noticed in the adrenal glands. No remarkable changes were noted in the thyroid. No specific tests were done to assess the hormonal status of the animal. Beneath the ulcers noted grossly, there were occasional fibrin thrombi and mild superficial infiltrate of neutrophils, lymphocytes and plasmacytes (data not shown). At the margin of the ulcerated sites, the epidermis is thickened with increased numbers of koilocytes, with perinuclear vacuolation (Fig. 2b) and nuclear hypertrophy. Occasionally, there were scattered discrete cytoplasmic basophilic and, rarely, eosinophilic (not shown) inclusion–like material within the vacuoles (Fig. 2c) and areas with large, pale amphophilic intranuclear inclusions, which displaced the chromatin peripherally (Figs. 2c and 2d). No intralesional bacteria were detected following Gram and Periodic acid-Schiff stains.

A heavy metals screen panel (selenium, copper, arsenic, lead, iron and zinc) performed in the liver was negative. Lung, liver and colon were sent for aerobic culture on blood agar plate (containing 5% (v/v) defibrinated sheep’s blood) with overnight incubation at 37°C. No
pathogenic bacteria were found. Using an avidin-biotin complex immunoperoxidase method, the following antibodies were used to detect viral antigens: 1) rabbit anti-measles virus,\textsuperscript{d} which detects antigen from several viruses classified within the \textit{morbillivirus} species, including CDV; 2) rabbit anti-\textit{adenovirus type 2},\textsuperscript{e} which detects antigen from a number of \textit{mastadenovirus} species, including \textit{Canine, Bovine and Equine adenovirus}; and 3) monoclonal antibodies specific for FHV (clone FHV7-7)\textsuperscript{c} and FCV (clone FVCS-19).\textsuperscript{c} Consistency of staining for each antigen was confirmed using tissue sections from known positive cases stained in parallel with the fisher tissues. Immunostaining for CDV in the skin and brain and for FHV, FCV, and adenovirus in the skin were negative.

Virus isolation was attempted using Madin-Darby canine kidney (MDCK) and Crandell feline kidney (CRFK) cell lines\textsuperscript{f} because they are well known to permit the isolation of several types of viruses.\textsuperscript{2,4,7,8} The fetal mink lung epithelial cell line (ML)\textsuperscript{g} was also used because it is derived from a member of the \textit{Mustelidae} family.\textsuperscript{1} Ten days old embryonated eggs\textsuperscript{h} (inoculated into the chorioallantoic sac and the chorioallantoic membrane) were also used for virus isolation. Unfortunately, all virus isolation attempts were negative. Skin lesions and surrounding tissues were ground in a glass tissue homogenizer and then prepared accordingly (as cell culture supernatants and allantoic fluids) and thereafter negatively stained for electron microscopy visualization. No virus particle could be observed in any of the prepared samples.

A pan-herpesvirus nested-PCR assay was conducted as previously described.\textsuperscript{3,13} Briefly, DNA was extracted from 1g of tissue sample, including skin, intestines and a pool of lung, liver and kidney, with the QIAamp DNA mini kit\textsuperscript{i} according to the manufacturer’s tissue protocol. The PCR was carried out with the QIAGEN Fast Cycling PCR kit,\textsuperscript{j} according to the
manufacturer’s specifications, with a set of three primers (DFA, ILK and KG1) as previously described. Nested-PCR was then performed under the same conditions using 5 µL of the first PCR reaction and two other primers (TGV, IYG), as previously described. An expected PCR product of 230 nucleotides (nt) was obtained only with skin tissues collected from and around the ulcers. The obtained PCR products were sequenced using a standard automated sequencing method and a sequence of 219 nt in length was submitted to GenBank Basic Local Alignment Search Tool (BLAST) for comparison. The BioEdit Sequence Alignment Editor software with the CLUSTAL W alignment method was used for nt and amino acids (aa) comparisons.

Nucleotide comparison showed that the nearest nt homology was with the viral DNA polymerase (DPOL) gene of Badger herpesvirus (BadHV), which is a virus classified in the Herpesviridae family within the subfamily Gammaherpesvirinae. The Fisher herpesvirus (FiHV) partial sequence possesses 85.3% nt identity and 91.7% aa identity with the BadHV DPOL gene published sequence (Fig. 3). Compared to BadHV, FiHV possesses a total of 23 nt silent mutations over a total of 32 nt mutations (Fig. 3). From a total of 73 deduced aa, 67 aa are identical between FiHV and BadHV (Fig. 3). Attempts to increase the length and the yield of the PCR product obtained by the pan-herpesvirus nested-PCR assay, and to amplify other viral genomic regions of the FiHV by designing new PCR primers based on the reported sequences of BadHV were unsuccessful.

Frostbite could cause vascular changes followed by epidermal necrosis but will most often affect the extremities like tips of the ears and tail. The footpads lesions were particularly similar to the cutaneous lesions observed when cats are infected with a systemic virulent strain of FCV but immunohistochemistry results were negative for FCV. CDV is known to infect animals classified within Mustelidae and to induce typical hyperkeratosis skin lesions located at the same...
sites where the fisher's ulcerative lesions were observed. Noteworthy, all CDV diagnostic results were negative and no microscopic lesions associated with CDV infection was found in tissues. Herpesviruses are well known to cause ulcers of the skin and mucosa to several mammals. It is possible that FiHV could be involved in the formation of the skin ulcers, but many animals are known to harbor herpesviruses asymptomatically. Unfortunately, no virus was isolated and observed by EM. If FiHV was the etiological cause of the lesions, it would be expected to observe virions by EM within and surrounding the cutaneous ulcers lesions. Noteworthy, the EM technical approach that was used possesses low sensitivity. In spite of the fact that the macroscopic skin lesions were impressive, the histopathological findings were rather mild and not typical of classic herpesvirus diseases (Fig. 2). It is noteworthy that similar histopathological findings have been previously observed in cases of herpesvirus associated skin lesions reported in cats and green sea turtles. Thus, the detection of FiHV DNA only within skin lesions and not in other tissues as well as the absence of other known pathogens in those lesions suggests that FiHV might be associated with the formation of such ulcers, but these findings alone are certainly not conclusive.

Is it possible that the pan-herpesvirus nested-PCR assay gave a false positive result? It is believed that if it was the case, then there is a good chance that this PCR assay would be positive for other fisher tissues, which was not the case (data not shown). The novel viral DNA sequence of FiHV was most closely related to the badger herpesvirus, a member of the subfamily Gammaherpesvirinae. Like the fisher, the badger is an animal classified in the Mustelidae family. Thus, it is reasonable to assume that FiHV is a new herpesvirus classified in the subfamily Gammaherpesvirinae. Further fisher cases would have to be studied and additional
work is needed to establish the involvement of FiHV in the ulcer lesions and in the death of the animal.

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Sources and manufacturers

a. Direction du laboratoire d’expertises en analyses alimentaires (DLEAA), Ste-Foy, Québec, Canada.
b. Laboratoire d’expertise en pathologie animale du Québec (LEPAQ), Québec, Québec, Canada.
c. Dr D.L. Godson, Prairie Diagnostic Services Inc., Saskatoon, Saskatchewan, Canada.
d. Dr D.B. Ziola, University of Saskatchewan, Saskatoon, Saskatchewan, Canada.
e. Lee Biomolecular Research, San Diego, CA, USA.
f. Dr S. Dea, Institut national de la recherche scientifique – Institut Armand-Frappier, Laval, Quebec, Canada.
g. NBL-7, ATCC CCL 64, American Type Culture Collection, Manassas, VA, USA.

h. Canadian Food Inspection Agency, Nepean, Ontario, Canada.

i. QIAGEN Inc., Mississauga, Ontario, Canada.

j. Sequencing Laboratory, Faculté de médecine vétérinaire, Université de Montréal, St-Hyacinthe, Québec, Canada.

k. BioEdit Sequence Alignment Editor software version 7.0.5.2, Ibis Therapeutics; Carlsbad, CA, USA.

References


**Figure legends**

**Figure 1.** Fisher skin: palmar pads and muzzle ulcer lesions. Enlarged ulcerative lesions of the muzzle can be seen in the left panel and enlarged ulcerative lesions of one foot pad are illustrated in the right panel. Arrows indicate the ulcers. Scale bars equal 500 and 100 μm, for panel A and B, respectively, and equal 30 μm for panel C and D.

**Figure 2.** Fisher skin: representative microscopic findings associated with the ulcer lesions. A) The margin between healthy skin (on the left) and the ulcer lesion (on the right) is indicated by a triangle. B) Koilocytes, with increased perinuclear vacuolation of the cells near an ulcer lesion is indicated by an arrow. C) Enlargement of panel B where koilocyte vacuolation is illustrated. In addition, an arrow indicates a basophilic pseudoinclusion body within the nuclei with a rim of chromatin. D) Localization of large pale amphophilic inclusions filling the nuclei within keratinocytes is indicated by an arrow.

**Figure 3.** Partial nucleotides and deduced amino acids sequences of the *Fisher herpesvirus* (FiHV) viral DNA polymerase (DPOL) gene compared to the *Badger herpesvirus* (BadHV) sequences. A) Nucleotides (nt) comparison of the FiHV (GenBank accession number HM579931) to the previously reported BadHV (GenBank accession number AF376034) nt sequence. B) Deduced amino acids comparison of the nt sequences illustrated in panel A. Numbers at the top of the nt sequences are nt positions based on the BadHV sequence.
HM579931. Underlined nt are silent mutations and bold nt are non silent mutations of the viral DPOL gene of FiHV compared to BadHV.
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