Iridovirus Infection in Chinese Giant Salamanders, China, 2010

To the Editor: The Chinese giant salamander (*Andrias davidianus*) is one of the world’s largest amphibian species and is often referred to as a living fossil. They primarily inhabit drainage basins of the Yangtze River, the Yellow River, and the Pearl River in the People’s Republic of China (1). Because of habitat loss, pollution, and overharvesting, the population of wild Chinese giant salamanders has dropped sharply (2,3). As a result, the Chinese giant salamander is artificially farmed in mesocosms for research and conservation. The mesocosms (ambient temperature <20°C) are maintained primarily in mountainous caves and ditches. During June–October 2010, a high mortality rate was reported in salamanders in ditch mesocosms in Shaanxi, Sichuan, and Henan, reaching an epidemic peak in July. Mortality rate reached 95% in the affected areas. Although bacteria, including *Aeromonas hydrophila* (4), were isolated from sick salamanders, antimicrobial drug treatment did not successfully improve the situation. Further pathologic analysis and viral testing were subsequently performed.

Pathologic changes were similar among the affected salamander populations. Gross anatomical changes included palpebral hyperemia or swelling; mouth pouch erythema; ecchymoses in the oral cavity; petechiae, ulceration; and erythema on the dorsal and ventral body surface; toe necrosis (online Technical Appendix Figure 1, panels A, B, wwwwnc.cdc.gov/EID/pdfs/10-1758-Techapp1.pdf); emaciation; friable and gray-black liver; and mottled, friable lesions of the kidney and spleen (online Technical Appendix Figure 1, panel C). Histologic examination showed hyperplastic lymphoid nodules in the spleen (Figure, panel A). Additionally, nuclear debris, macrophages (Figure, panel A), and intracytoplasmic inclusion bodies (Figure, panel B) were observed in the lymphoid nodules. Liver sinusoids were enlarged and contained large numbers of macrophages. Degenerating hepatocytes were noted (online Technical Appendix Figure 1, panel D). Degenerate renal epithelial cells were shed from the basement membrane and were found in the lumen of the renal tubules (online Technical Appendix Figure 1, panel E). A large number of viral particles were observed in renal epithelial cells (online Technical Appendix Figure 1, panel G). Virus was isolated from the liver, kidney, and spleen. Electron microscopy was performed on random tissue samples from organs positive for an unidentified virus. Icosahedral viral particles ≈150 nm in diameter were observed in the cytoplasm of some cells (Figure, panel B; online Technical Appendix Figure 1, panels F, G).

On the basis of the gross lesions and the appearance of the virus, we suspected that it was a member of the iridovirus family. To test this hypothesis, genomic DNA (gDNA) was extracted from the isolated virus by using a commercial kit (Genray, Shanhai, China). PCR was performed by using 3 sets of primers targeting 681 bp, 568 bp, and 616 bp iridoviral fragments respectively, from the major capsid protein gene (GenBank accession no. U36913; 5′-CCCTCCCCATTCTTCTTCCTCC-3′, 5′-GCGTGGGTGTCAGTCTACCGTA...
AT-3), the ATPase gene (GenBank accession no. M80551; 5′-CCAAGAG GCACATCATACCG-3′, 5′-GCT GGACATCTCCTACGACCC-3′), and the thymidine kinase gene (GenBank accession no. AY837779; 5′-GGGCTATGTATTTGAAGA CGC-3′, 5′-TTGAAAATTTGAGTG GAGGG-3′). Resulting PCR products from 10 salamanders were sequenced and compared with the corresponding sequences of the 5 known iridovirus strains by using a BLAST search (http://blast.ncbi.nlm.nih.gov/Blast.cgi) (frog virus 3, GenBank accession no. AY548484; soft-shelled turtle iridovirus, GenBank accession no. EU627010; tiger frog virus, GenBank accession no. AF389451; epizootic hematopoietic necrosis virus GenBank accession no. FJ433873; and Ambystoma tigrinus stebbensis virus, GenBank accession no. AY150217). The sequences of the 3 PCR products from the virus-infected Chinese giant salamanders (GenBank accession nos. HQ829176, HQ829177, and HQ829178) showed >96% homology with the corresponding sequences of the 5 iridovirus strains. Additionally, neighbor-joining tree analysis showed that the virus was clustered in 1 lineage with frog virus 3, soft-shelled turtle iridovirus, and tiger frog virus (online Technical Appendix Figure 2). These results suggest that the high mortality rates in Chinese giant salamanders were caused by a virus in the iridovirus family.

The iridoviruses are carried in the bodies of vertebrates such as gopher tortoises (Gopherus polyphemus) (5), Chinese forest frogs (Rana dybowskii) (6), and fish (7,8). Iridoviruses are thought to be transmitted horizontally in lower vertebrates, such as bullfrogs (7,9,10). In addition, some iridovirus infections may be chronic or conditional (7). In this study, the virus was isolated from the liver and spleen of 30 sick (n = 7) or dead (n = 23) salamanders that were farmed in ditch mesocosms, where ambient temperatures were unusually high (>25°C) at the time of the epidemic. Although the virus also was isolated from animals living in cooler cave mesocosms (ambient temperature <18°C), these animals showed no apparent signs of illness. Studies have reported that, when infection is detected early in the course of the disease and when exogenous stress is minimized, mildly affected bullfrogs are able to clear the virus (9,10). The high water temperatures in the ditch mesocosms (i.e., >25°C) and the associated stress on the animals may have increased disease in ditch-dwelling Chinese giant salamanders. This seems particularly likely, given the absence of clinical signs of disease in infected salamanders that lived in the cooler cave mesocosms (i.e., <18°C). In addition, absence of exposure of Chinese giant salamanders to other animal carriers of the virus may prevent horizontal transmission of iridovirus.

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Technical Appendix

Figure 1. Gross anatomic and histologic changes in sick Chinese giant salamanders (Andrias davidianus), People’s Republic of China, 2010. Viral particles are present in hepatocytes and renal cells. A) Palpebral hyperemia or edema; mouth pouch inflammation. B) Toe and tail necrosis (arrow). C) A sick Chinese giant salamander with a diseased spleen (arrow), a friable and gray-black liver (asterisk), and
mottled necrotic lesions in the kidney (arrowhead). D) Histologic changes in the liver of a sick Chinese giant salamander. Arrows indicate a large number of macrophages in enlarged liver sinusoids; asterisks indicate degenerating hepatocytes (hematoxylin and eosin stain; scale bar = 80 μm). E) Histologic changes in the kidney of a sick Chinese giant salamander. Asterisks indicate degenerating epithelial cells in renal tubules; arrows indicate epithelial cells in the renal tubule lumen (hematoxylin and eosin stain; scale bar = 80 μm). F) Electron microscopic image of viral particles in hepatocytes (scale bar = 200 nm). G) Electron microscopic image of viral viral particles in renal cells (scale bar = 200 nm). VP: viral particles in cytoplasm; Nu: nucleus; arrowheads, provirus in nuclear membrane; arrows, provirus in nucleus (scale bar = 200 nm).

Figure 2. Neighbor-joining (NJ) tree comparison of gene sequences of viral isolates from Chinese giant salamander (CGSIV) and 5 known viral strains, People’s Republic of China, 2010. Major capsid protein, ATPase, and thymidine kinase gene sequences of 5 known iridovirus strains were obtained from GenBank (frog virus 3 [FV3]: AY548484; soft-shelled turtle iridovirus [STIV]: EU627010; tiger frog virus [TFV]: AF389451; EHNV: FJ433873; Ambystoma tigrinum stebbensi virus [ATV]: AY150217) and corresponding sequences of the 15 CGSIV (GenBank accession nos. HQ829176, HQ829177, and HQ829178). All sequences were edited by using the DNASTAR 5.0 package (DNASTAR, Madison, WI, USA). These sequences were aligned with the ClustalX package (www.clustal.org) and truncated to equal bp, respectively. Insertions/deletions in the aligned sequences were excluded in the analyses. An NJ tree using the Kimura 2-parameter model with 1,000 bootstrap replicates was constructed to identify a possible phylogenetic tree in MEGA2.1 (www.megasoftware.net). A) NJ tree of major capsid protein sequences of CGSIV and the 5 viral strains. B) NJ tree of ATPase sequences of CGSIV and the 5 viral strains. C) NJ tree of thymidine kinase sequences of CGSIV and the 5 viral strains. D) NJ tree of major capsid protein + ATPase + thymidine kinase sequences of CGSIV and the 5 viral strains. The NJ tree showed that CGSIV was clustered in 1 lineage with FV3, STIV, and TFV, indicating that CGSIV belongs to the family Iridoviridae.