

Acknowledgments

We thank Catherine Graham for the elk CWD prions and Richard Rubenstein for the 3F4 monoclonal antibodies.

This work was supported by the Alberta Prion Research Institute, Natural Sciences and Engineering Research Council, the Alberta Livestock and Meat Agency, and the Genome Canada Large Scale Applied Research Program.

Dr. Herbst is a research associate and Dr. Duque Velásquez is a postdoctoral fellow at the University of Alberta. Their primary research interest is the mechanism(s) of pathogenicity underlying neurodegeneration, as exemplified by prion diseases in animals and humans.

References


Address for correspondence: Debbie McKenzie, University of Alberta, 120 Brain and Aging Research Building, Edmonton, AB T6G 2R3, Canada; email: debbie.mckenzie@ualberta.ca


Shuilian Chen,1 Heng Zhang,1 Meiling Luo, Jingfang Chen, Dong Yao, Faming Chen,2 Ruchun Liu,2 Tianmu Chen2

Author affiliation: Changsha Center for Disease Control and Prevention, Changsha, Hunan, China

DOI: https://doi.org/10.3201/eid2309.161704

We report rabies virus transmission among solid organ transplantation recipients in Changsha, China, in 2016. Two recipients were confirmed to have rabies and died. Our findings suggest that more attention should be paid to the possibility of rabies virus transmission through organ transplantation for clinical and public health reasons.

In 2016, Zhou et al. reported a case of rabies virus transmission in China that was probably a result of organ transplantation (1). We report on rabies transmission that occurred among solid organ transplant recipients in Changsha, China, during December 2015–January 2016.

In November 2015, the donor, a previously healthy boy, showed development of fever, insomnia, and agitation. On day 6 of infection, these symptoms progressed, and he was sent to a healthcare center. At this time, he experienced weakness, no desire to drink water, poor appetite, and panic. One day later, he began vomiting, and was admitted to a local hospital (hospital A), where he exhibited anemophilia, convulsions, limb rigidity, and hypersalivation. The patient was moved to hospital B (days 7–14) in Changsha.

At admission, some examination findings indicated a possibility of viral encephalitis (online Technical Appendix, Table 1, https://wwwnc.cdc.gov/EID/article/23/9/16-1704-Techapp1.pdf). Subhypothermia hibernation therapy and assisted ventilation were administered within 72 hours of admission, and the patient’s vital signs became stable. On day 10, hyponatremia was observed, and on day 11, the patient again became febrile and tachycardic, with hypertensive abdominal distention and alimentary tract hemorrhage. On day 13, viral encephalitis was diagnosed, and rabies was suspected. However, rabies virus antibody tests performed on serum samples by using ELISA yielded negative results. On day 14, the patient was transferred to hospital C, where he became comatose and was declared brain dead. Permission was granted for organ donation, because no specific pathogen had been detected and China’s organ

1These authors contributed equally to this article.
2These authors contributed equally to this article.
transplant policy allows for organ donations in cases of viral encephalitides without laboratory-confirmed pathogens. Rabies virus–specific binding antibodies were also not detected in the serum samples. The donor had possibly been exposed to rabies but had not been vaccinated against rabies (online Technical Appendix). The patient’s kidneys and liver were removed for transplantation in hospital D in Changsha on the following day.

Kidney and liver transplantations were performed on December 10, 2015. Two female recipients in hospital D received the kidneys, and an 8-month-old girl in hospital E in Shanghai, China, received the liver. No other organs were collected or transplanted from this donor. The surgeries were all uneventful, and the recipients were discharged after transplantation. However, all 3 recipients were eventually readmitted to the hospital with complex symptoms (Figure).

On the 40th day after transplantation, the left kidney recipient (29 years of age) revisited hospital D, reporting worsening right quadrant abdominal pain and vomiting. In the next 3 days, she successively had anemophobia, shortness of breath, chest tightness, anxiety, insomnia, and hypersalivation. During hospitalization, her blood pressure was as high as 248/148 mm Hg. On her 7th day in the hospital, she became comatose and then died.

On the 43rd day after transplantation, the right kidney recipient (age 47 years) developed aches in her lower limbs, insomnia, and a decreased appetite. She was readmitted to hospital D the next day. On the 3rd day after admission, she exhibited hydrophobia, anemophobia, photesthesis, and fatigue. She showed agitation, dyspnea, and hypersalivation on the 4th day; she became comatose and died 1 day later.

The liver recipient was readmitted with pneumonia on the 34th day after transplantation and died of asphyxia and
multiple organ failure within 5 days. This patient did not show any signs or symptoms of rabies or encephalitis.

None of the recipients had been exposed to potentially rabid animals or had been vaccinated previously for rabies (online Technical Appendix). Both kidney recipients tested positive for rabies virus (online Technical Appendix Table 2). The genome sequences of the rabies virus isolates from the right kidney recipient (isolate no. CCS1501H) were ≈11 kb nucleotides in length and belonged to the China I lineage. No testing for rabies was done on the donor or on the liver recipient.

In the past 10 years, rabies transmission by solid organ transplantation has been described occasionally worldwide (2–4). Hence, rabies transmission through organ transplantation is a clinical and public health concern. To prevent future cases such as this, we recommend that patients with unexplained encephalitis or mental status changes should not be used as organ donors even if tests for some infectious causes of encephalitis are negative. In addition, if rabies is suspected in the donor after organs have been transplanted, the recipients should also not be used as organ donors. An antibody test is not the ideal choice for the diagnosis of rabies virus and by itself cannot reliably exclude rabies from the differential diagnosis. For this reason, a combination of multiple techniques, preferably direct fluorescent antibody test and reverse transcription PCR, should be used before organ transplantation, especially when the donor is suspected of having rabies or a potential exposure to rabies. In addition, if a patient has meningoencephalitis of unknown cause, a specific epidemiologic and laboratory evaluation should be performed to conclusively rule out rabies as a cause of illness before organ donation.

Dr. Shuilian Chen is a medical doctor in the Changsha Center for Disease Control and Prevention in Changsha, China. Her main field of research is emerging infectious disease control and prevention.

References

Identification of Clade E Avipoxvirus, Mozambique, 2016

Lourenço P. Mapaco, Zeiss Lacerda, Iolanda V.A. Monjane, Esayas Gelaye, Afonso H. Sussuro, Gerrit J. Viljoen, William G. Dundon, Sara J. Achá

Author affiliations: Agrarian Research Institute of Mozambique, Maputo, Mozambique (L.P. Mapaco, I.V.A. Monjane, A.H. Sussuro, S.J. Achá); Eduardo Mondlane University, Maputo (Z. Lacerda); National Veterinary Institute, Debre Zeit, Ethiopia (E. Gelaye); International Atomic Energy Agency, Vienna, Austria (G.J. Viljoen, W.G. Dundon)

DOI: https://doi.org/10.3201/eid2309.161981

Analysis of scab samples collected from poultry during outbreaks of fowlpox in Mozambique in 2016 revealed the presence of clade E avipoxviruses. Infected poultry were from flocks that had been vaccinated against fowlpox virus. These findings require urgent reevaluation of the vaccine formula and control strategies in this country.

Avipoxviruses are large, enveloped DNA viruses that belong to the genus *Avipoxvirus* in the *Chordopoxvirinae* subfamily of the family *Poxviridae*. These viruses cause disease in a large number of bird species and are generally named after the species from which the virus was first isolated and characterized (1). Fowlpox virus (FPV) has caused substantial economic losses in domestic poultry resulting from reduced egg production and growth, blindness, and death, with a death ratio that can reach as high as 50%.

Phylogenetic analyses of the *Avipoxvirus* genus are usually conducted with the segments of the genes encoding the 4b core-like protein (P4b) and the DNA polymerase, which are both highly conserved among poxviruses (2,3). Using these loci, researchers have seen that most strains cluster into 3 major clades, namely A, B, and C, with clade A being subdivided further into subclades A1–A7 and clade B into subclades B1–B3 (3–5). Two additional clades