We report a case of natural infection with severe acute respiratory syndrome coronavirus 2 transmitted from an owner to a pet ferret in the same household in Slovenia. The ferret had onset of gastroenteritis with severe dehydration. Whole-genome sequencing of the viruses isolated from the owner and ferret revealed a 2-nt difference.

Natural infections with severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) in domestic animals living in infected households have been reported (1). Because of their increased popularity as a pet (2), domestic ferrets (Mustela putorius furo) pose a high risk for transmitting anthropozoonotic infections. A recent study in Spain showed that natural SARS-CoV-2 infections can occur in ferrets kept as working animals for rabbit hunting, especially if a high viral circulation is present in the human population (3). Further, ferrets are common laboratory animal models and, at least in experimental conditions, have been shown to be highly susceptible to SARS-CoV-2 infection and likely to transmit the virus to other ferrets without apparent clinical signs (4).

The Study
On November 20, 2020, a 5-year-old neutered male domestic ferret had signs of acute gastroenteritis, including apathy, anorexia, vomiting, and profuse mucous diarrhea. Another ferret in the same household appeared healthy. Because the ferret’s condition did not improve, the owner took it to a veterinary hospital for clinical examination on November 23. The ferret was lethargic and, on the basis of skin turgor, was >5% dehydrated with low body temperature (36.4°C, reference range 37.8–40°C) and slow heart rate (180 beats/min, reference 200–400 beats/min). The body condition of the ferret was good, with a bodyweight of 1.3 kg. Several hematology and serum biochemistry results were elevated: red blood cell count (12.36 × 10⁶/µL, reference 7.01–9.65 × 10⁶/µL), hemoglobin concentration (21.2 g/dL, reference 12.2–16.5 g/dL), and hematocrit (0.57%, reference 0.36%–0.48%); blood urea nitrogen (>129.94 mg/dL, reference 18–32 mg/dL), hyperproteinemia (8.5 g/dL, reference 4.5–6.2 g/dL), hyperglobulinemia (4.4 g/dL, reference 2.8–3.6 g/dL), and borderline hyperalbuminemia (4.0 g/dL, reference 2.5–4.0 g/dL) were consistent with dehydration and possible infection. The results of all other hematologic and biochemical values were within reference ranges. Whole-body radiographs (Appendix Figure, https://wwwnc.cdc.gov/EID/article/27/9/21-0774-App1.pdf) showed splenomegaly and gas accumulation in intestinal loops. Interstitial and alveolar patterns of cranial lung lobes were present, indicating possible lobar pneumonia. The ferret was hospitalized and initially given fluid therapy, amoxicillin, esomeprazole, maropitant, and dexamethasone. Three days later, the clinical status of the ferret improved, hematologic and biochemical values normalized, and the ferret was scheduled for discharge. However, on the same day, the owner informed the veterinary hospital of having positive results for SARS-CoV-2 RNA tested on November 24, after 9 days of malaise. Additional rectal and oropharyngeal swab specimens and blood samples were taken from the ferret for further diagnostic procedures, and the ferret was discharged from the hospital and put into isolation at the owner’s home. Samples were not taken from the other pet ferret at the residence, but the rest of household members tested negative for SARS-CoV-2 RNA on November 25.

We tested the ferret’s samples for SARS-CoV-2 RNA (Appendix) and ferret-specific enteric
Transmission of SARS-CoV-2 from Human to Ferret

 coronavirus (FERCV) (5) by real-time reverse transcription PCR; influenza A and B viruses (6) by reverse transcription PCR; and herpesviruses (7), adenoviruses (8), and circoviruses (9) and by PCR. The only positive result was the detection of SARS-CoV-2 RNA in the rectal and oropharyngeal swab specimens. In the oropharyngeal swab specimen, all 3 targeted genes (envelope, cycle threshold [Cₜ] 27.7; RNA dependent RNA polymerase, Cₜ 28.5; and nucleocapsid, Cₜ 32.1) were detected, and in the rectal swab specimen only envelope gene (Cₜ 35.6) was detected, a finding probably attributable to a lower virus concentration. To compare the SARS-CoV-2 detected in the owner and the ferret, we conducted whole-genome sequencing on Illumina MiSeq (Illumina, https://www.illumina.com) on the basis of the ARTIC protocol (https://artic.network/ncov-2019/ncov2019-bioinformatics-sop.html). The complete genome sequences were deposited in the GISAID database (https://www.gisaid.org; accession nos. EPI_ISL_1490186 and EPI_ISL_1490187). According to the pangolin nomenclature, the sequences belonged to the B.1.258 Pango lineage, which was on the rise in Slovenia in November 2020 (Figure 1). The comparison of both sequences showed ≈100% identity, differing by 2 nucleotides (position/owner/ferret: 2,097/G/T; 22,832/C/A).

We also confirmed the SARS-CoV-2 infection in the ferret on the basis of SARS-CoV-2 IgG seroconversion and development of neutralizing antibodies. We tested the ferret’s acute and convalescent serum samples with an in-house immunofluorescent assay (Appendix). The first serum sample obtained on day 6 after disease onset tested negative; however, seroconversion was observed on day 19, when the IgG titer was 1:1,024 (Figure 2, panels A, B). In addition, we detected a high neutralizing antibody titer of 1:320 in the second serum sample (Figure 2, panel C).

Conclusions

 SARS-CoV-2 originated in animals, jumped into humans, and is now easily transmitted among humans. In addition to spreading from animals to humans, the virus can be transmitted back into animals, as observed in farmed mink (Neovison vison) (10). Most experimentally infected ferrets do not exhibit clinical signs or have only mild fever, lethargy, loss of appetite, and occasional cough (4,11). Also, among working ferrets naturally infected with SARS-CoV-2 in Spain, no signs of illness were reported (3).

In our study, the infected ferret had onset of severe disease with gastroenteritis, pneumonia, and dehydration and required aggressive fluid therapy and supportive care with antibiotics, antacids, antimetics, and parenteral dexamethasone. The ferret responded to the therapy promptly and fully recovered in 3 days. Acute epizootic catarrhal enteritis caused by coronaviruses (CoronaVirus-like enteritis virus [Cov-Enteritis] or porcine coronavirus [PCV] (12) can cause a similar disease in ferrets (13). However, diarrhea and viremia were not detected in the ferret in this study, suggesting that the infection was not caused by a coronavirus. The diagnosis of SARS-CoV-2 infection in the ferret was confirmed by both reverse transcription real-time PCR and whole-genome sequencing, which is consistent with the clinical, serological, and histopathological findings from this study.
by FERCV was one of the plausible differential diagnoses in the initial treatment plan for the ferret. For this reason, dexamethasone was used parenterally because additional treatment with a short course of steroids might speed the recovery and reduce future problems of malabsorption attributable to villi destruction caused by fulminate FERCV infection (12).

In humans, the effective drugs against coronavirus disease are poorly defined, yet dexamethasone in combination with supportive therapy is frequently used (13). However, the risk for unnecessary use and adverse effects must be considered before treatment attempts with corticosteroids.

We assume that SARS-CoV-2 likely spread from the infected owner to the ferret living in the same household. Symptoms appeared in the owner 4 days before the ferret became ill. All other household members tested negative for SARS-CoV-2 RNA, ruling out asymptomatic infected persons in the family. Another close contact ferret in the same household appeared healthy. Likewise, no disease among staff or animals at the veterinary hospital was reported during or after the hospitalization of the ferret. Nevertheless, ferrets as laboratory models were shown to shed SARS-CoV-2 up to 8 days postinfection in nasal swab, saliva, urine, and fecal samples. Ferrets can effectively transmit the infection to other animals (14) or possibly humans, thus highlighting the importance of recognizing the infection in pets early to prevent spread to other animals or humans in the same household or elsewhere (15).

In the mink farm outbreak in Denmark, SARS-CoV-2 transmission was shown to spill over from minks to humans accumulating mutations that are resistant to neutralizing antibodies or vaccines along the way (10). In our study, whole-genome sequencing of the virus detected in the owner and ferret revealed only a 2-nt difference, and neither of those was present in the spike protein gene. Nonetheless, retaining the One Health approach is crucial for early detection and monitoring of emerging zoonoses in humans.

Acknowledgments
We thank all members of the COVID-19 Next Generation Sequencing team for great technical assistance in sequencing SARS-CoV-2 genomes and the staff at the Toplica Veterinary Hospital for caring for the ferret. We thank the staff at the Institute for Poultry, Birds, Small Mammals, and Reptiles for their assistance in performing PCR assays. Thanks to the owners of the ferret for their kind support and for allowing publication of this report.

This work was supported by the Slovenian Research Agency (grant nos. P3-0083 and V3-2034 at the Institute of Microbiology and Immunology, Faculty of Medicine, University of Ljubljana and P4-0092 at the Faculty of Veterinary Medicine, University of Ljubljana) and by the European Virus Archive Global project, which received funding from the European Union’s Horizon 2020 research and innovation program under grant no. 871029.

About the Author
Dr. Račnik is an associate professor at the Faculty of Veterinary Medicine, University of Ljubljana, Slovenia, and diplomate of the European College of Zoological Medicine, wildlife population health specialty. His primary research interests include clinical veterinary medicine and emerging diseases of exotic pets and wild birds.

References
Transmission of SARS-CoV-2 from Human to Ferret

in cats and dogs from households in Italy. Nat Commun. 2020;11:6231. https://doi.org/10.1038/s41467-020-20097-0


Address for correspondence: Jožko Račnik, Institute for Poultry, Birds, Small Mammals, and Reptiles, Faculty of Veterinary Medicine, University of Ljubljana, Gerbičeva 60, 1000 Ljubljana, Slovenia; email: josko.racnik@vf.uni-lj.si

EID Podcast Developing Biological Reference Materials to Prepare for Epidemics

Having standard biological reference materials, such as antigens and antibodies, is crucial for developing comparable research across international institutions. However, the process of developing a standard can be long and difficult.

In this EID podcast, Dr. Tommy Rampling, a clinician and academic fellow at the Hospital for Tropical Diseases and University College in London, explains the intricacies behind the development and distribution of biological reference materials.

Visit our website to listen: https://go.usa.gov/xyfJX
Transmission of SARS-CoV-2 from Human to Domestic Ferret

Appendix

Materials and Methods

SARS-CoV-2 real-time reverse transcription—polymerase chain reaction (rRT-PCR)

Viral RNA was isolated with EZ1 Advanced XL using Virus Mini Kit v2.0 (Qiagen) following the manufacturer’s instructions. For real-time RT-PCR amplification, we used LightMix® Modular SARS-CoV E-, RdRp-, and N-gene (Tib-Molbol, Berlin Germany) in combination with TaqMan® Fast Virus 1-Step MasterMix (Thermo Fisher Scientific, Grand Island, NY, USA) on QuantStudio 7 Pro Real-Time PCR System (Applied Biosystems, Thermo Fisher Scientific). The cycling conditions were as follows: reverse transcription at 50°C for 5 min and 95°C for 20 sec, followed by 40 cycles of PCR at 95°C for 3 sec and 60°C for 30 sec.

Whole genome sequencing

RNA was used for cDNA synthesis and PCR amplicon generation according to the nCoV-2019 sequencing protocol designed by Artic Network (https://artic.network/ncov-2019). One ng of cleaned PCR amplicon was used for synthesis of the NGS library with the Nextera XT DNA Library Preparation Kit (Illumina) following the manufacturer’s instructions with an increasing number of PCR cycles in step Amplify Libraries from 12 (instructions) to 14. The NGS libraries were cleaned using a 1.8 ratio sample: Agencourt ampure XP beads (Beckman Coulter) as per instructions for short amplicons. We measured the NGS libraries’ concentration with Qubit 3 (Thermo Scientific) using the Qubit dsDNA HS Assay. We measured the average fragment size and checked the NGS libraries for primer dimers with Agilent 2100 Bioanalyzer (Agilent) using the Agilent High Sensitivity DNA Kit (Agilent). NGS libraries were normalized following the manufacturer’s instructions for standard normalization for the MiSeq system. In brief, all libraries were diluted to 4 nM with nuclease-free water (Qiagen), and 10 μL of each NGS library was pooled together. The pooled NGS libraries were vortexed, and 5 μL of pool was used for library denaturation and sequencing.
**Immunofluorescent test**

The sera were diluted from 1:16 to 1:1,024 and tested using indirect immunofluorescence assay on spot slides containing Vero E6 cells infected with SARS-CoV-2 (strain Slovenia/SI-4265/20, D614G; EVA GLOBAL Ref-SKU: 005V-03961). In brief, serum dilutions were dispensed on the antigen spot on the slide. The slide was incubated for 30 min at room temperature and washed in phosphate-buffered saline (PBS), and afterward goat anti-ferret IgG (H+L) FITC conjugate (KPL, USA) dilution at a ratio of 1:10 was added. Another incubation at room temperature followed. Then the slide was washed in PBS and dried again. The fluorescence was examination under a fluorescent microscope (Eclipse 80i; Nikon).

**Neutralization test**

VERO E6 cells were seeded on a 96-well plate in a concentration of $10^5$ cells/mL 1 day before the neutralization test was performed. The plasma sample was incubated at 56°C for 30 min. The sample was diluted from 1:10 to 1:1,280 in DMEM with 1% FBS. SARS CoV-2 virus (strain Slovenia/SI-4265/20, D614G; EVA GLOBAL Ref-SKU: 005V-03961; 100 TCID50/mL) was added to serial dilutions in a ratio of 1:1 and incubated for 1 hour. Then 50 µL of a mixture of diluted plasma and virus was inoculated into the VERO E6 cells. The inoculated cells were incubated for 4 days at 37°C and 5% CO₂. On day 5, the cells were observed for CPE and the antibody titer was detected as the highest dilution that CPE was inhibited on for at least 2 of 3 wells.
Appendix Figure. Whole-body radiograph in laterolateral (A) and ventrodorsal (B) projections of a ferret presenting splenomegaly and gas accumulation in intestinal loops. Moreover, interstitial and alveolar patterns of cranial lung lobes are present, indicating possible lobar pneumonia.