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SARS-CoV-2 Infection in Beaver Farm, Mongolia, 2021

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We report an outbreak of COVID-19 in a beaver farm in Mongolia in 2021. Genomic characterization revealed a unique combination of mutations in the SARS-CoV-2 of the infected beavers. Based on these findings, increased surveillance of farmed beavers should be encouraged.

The COVID-19 pandemic that began in 2019 remains uncontained, and fatalities and multiple waves of infection continue to occur worldwide (1). The causative agent, SARS-CoV-2, has been detected in humans and several animal species, including domestic, wild, and laboratory animals (2,3). Because SARS-CoV-2 can be transmitted from humans to animals and back to humans, understanding the dynamics of infection in animals can contribute to the creation of more comprehensive response strategies.

We identified SARS-CoV-2 infection in beavers (Castor fiber) farmed for conservation reasons in Mongolia and report on serologic and whole genome sequence data from this outbreak. The beaver farm, located in the Bayanzurkh district in Ulaanbaatar, Mongolia, reared 32 adults and 16 kits in 2021. They were housed indoors in a large area separated by waist-high walls, with space for multiple animals. One of the 7 employees of the farm had influenzalike symptoms for several days and was diagnosed with COVID-19 on August 6, 2021. On August 9, the beaver farm reported the death of 2 beavers (one 6 months of age and one 2 years of age) after signs of coughing, nasal discharge, rasping on auscultation of the lungs and chest cavity, sluggish movement, and aversion to food. On August 13, research investigators collected nasal swabs, saliva, and 7 tissue samples

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(lung, kidney, liver, heart, spleen, larynx, and tongue from the 2 dead animals. Researchers also collected nasal swab specimens, saliva, and blood from 7 other beavers with notable clinical signs of coughing and purulent nasal discharge. Follow-up investigation on August 18 or 19 and on September 12 included collection of additional nasal swab specimens, saliva, and blood samples from the same animals as well as from 2 healthy animals (September 12 only).

All samples were transported to a Biosafety Level 3 facility in Ulaanbaatar and were screened by quantitative reverse transcription PCR according to the Peiris protocol (4). The results showed that 46 of 48 specimens from 9 animals with clinical signs, including the 2 dead animals, tested positive for SARS-CoV-2 RNA. Serum was separated from the blood samples by centrifugation (2,000 × *g* for 10 min) and stored at -20° C until required. The serum samples were then subjected to antibody screening by using a commercial ELISA kit (ID Screen SARS-CoV-2 Double Antigen Multi-species ELISA; Innovative Diagnostics, https://www.innovative-diagnostics.com). Fifteen of 23 samples tested positive and 1 was intermediate, indicating that all animals became antibody positive within 1 month of confirmation of SARS-CoV-2 RNA positivity. One clinically unremarkable beaver tested positive for SARS-CoV-2 antibodies, indicating a possible subclinical infection (Table).

We shipped 5 randomly selected quantitative reverse transcription PCR-confirmed SARS-CoV-2positive RNA samples to the Animal Production and Health Laboratory (Seibersdorf, Austria), a joint program of the International Atomic Energy Agency and the Food and Agriculture Organization of the United Nations, and subjected them to whole-genome sequencing (Appendix 1, https:// wwwnc.cdc.gov/EID/article/30/2/23-1318-App1. pdf; Appendix 2, https://wwwnc.cdc.gov/EID/ article/30/2/23-1318-App2.xlsx). Based on genotype analysis, all 5 genome sequences were assigned to the B.1.617.2 lineage, commonly referred to as the Delta variant. At the time of sampling, Alpha and Delta variants of SARS-CoV-2 were being identified in humans in Mongolia. The closest related sequences to those we identified in the beavers studied were from



Figure. Phylogenetic tree of SARS-CoV-2 identified from beavers and humans in Mongolia (gray boxes) and reference sequences. The circle size indicates the bootstrap values at the node. The vertical bar shows the genetic distance. SARS-CoV-2 lineages are identified at right. GenBank accession numbers and date identified are shown for reference sequences; the newly obtained sequence data were deposited in GenBank (accession nos. OR389473–7).

Table. Sampling date and results of serologic analysis of SARS-CoV-2 antibodies from	farmed beavers, Ulaanbaatar, Mongolia, 2021*
Date of swah sampling and aPT PCP results	Data of corum compling and ELISA recultet

		Date of swab sampling and qRT-PCR results		Date of serum sampling and ELISA results†			
Animal ID	Status	2021 Aug 13	2021 Aug 19	2021 Sep 12	2021 Aug 13	2021 Aug 18	2021 Sep 12
1	Died Sep 8	Positive	NT	NT	NT	NT	NT
2	Died 2021 Sep 8	Positive	NT	NT	NT	NT	NT
3	Sick	Positive	Positive	Positive	Negative	Negative	Positive
4	Sick	Positive	Positive	Positive	Negative	Negative	Positive
5	Sick	Positive	Positive	Positive	Negative	Positive	Positive
6	Sick	Positive	Positive	Positive	Positive	Positive	Positive
7	Sick	Positive	Positive	Positive	Positive	Positive	Intermediate
8	Sick	Positive	Positive	Negative	Positive	Positive	Positive
9	Sick	Positive	Positive	Negative	Negative	Positive	Positive
10	Healthy	NT	NT	NT	ŇT	NT	Positive
11	Healthy	NT	NT	NT	NT	NT	Negative

*ID, identification; qRT-PCR, quantitative reverse transcription PCR; NT, not tested.

†ID Screen SARS-CoV-2 Double Antigen Multi-species ELISA (Innovative Diagnostics, https://www.innovative-diagnostics.com). The interpretation is based on the signal-to-noise (S/N) ratio, (sample optical density [OD] 450/negative control OD450) × 100, according to instruction manual. Positive: S/N>100.0; intermediate: 100.0>S/N>50.0; negative: 50.0>S/N.

human SARS-CoV-2 in Mongolia (GenBank accession nos. ON008302, OM190617, and OM961234) identified during April–September 2021 (Figure). In addition to 4 mutations in the spike region, the sequences shared 7 amino acid substitutions in open reading frame [ORF] 1a, 4 amino acid substitutions in ORF1b, and 1 amino acid substitution in nucleocapsid genes. In the beaver sequences, 4 amino acid substitutions identified were not in the human isolates from Mongolia: S2500F, A3657V in ORF1a and H604Y, T1404M in ORF1b. Although those substitutions have been identified individually in SARS-CoV-2 sequences in GenBank and the GISAID database (https://www.gisaid.org), there are no records of sequences with all 4 mutations.

Several cases of SARS-CoV-2 transmission between humans and animals have already been reported (5-8). An alarming aspect of SARS-CoV-2 infection in animals is that host animals can maintain the virus and contribute to the emergence in humans of new variants that have accumulated multiple mutations (7-10). Indeed, the specific combination of mutations observed in the beavers we studied has not been found in other SARS-CoV-2 sequences in public databases (as of November 2023). This finding suggests that the mutations might have occurred or accumulated after the introduction of the virus into the beaver population. Because the emergence of viruses with mutations not targeted by current SARS-CoV-2 vaccines is a credible possibility, more active surveillance of SARS-CoV-2 infection in animals should be encouraged to identify the appearance of mutated viruses. In intensively farmed animals, species-species and species-humans contact is more frequent than in animals dwelling in other environments, which might increase the risk for zoonotic pathogen transmission (2). Thus, implementing more active surveillance and infection control strategies is critical to disease prevention and containment.

Samples used in this study were those submitted to the State Central Veterinary Laboratory, Mongolia, for emergency diagnosis of SARS-CoV-2. Ethics approval was not required.

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Severe Infective Endocarditis Caused by Bartonella rochalimae

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A 22-year-old man from Guatemala sought care for subacute endocarditis and mycotic brain aneurysm after living in good health in the United States for 15 months. *Bartonella rochalimae*, a recently described human and canine pathogen, was identified by plasma microbial cellfree DNA testing. The source of infection is unknown. A 22-year-old man with a history of an unrepaired congenital ventricular septal defect (VSD) experienced 3 months of progressive dyspnea on exertion, weight loss, and fatigue and 2 weeks of debilitating weakness. He had been born in Guatemala, where he worked in construction; he had had contact with goats, horses, cattle, and chickens but reported no contact with dogs or cats. Eighteen months before he sought care, he had migrated to the mid-Atlantic region of the United States, where he lived with his uncle and a few other adults in a suburban town. He continued to work in construction, did not use illicit drugs, and had 1 female sexual partner. Six months after he arrived, his uncle took in a stray dog.

The patient was afebrile, hypotensive, bradycardic, and thin. A systolic ejection murmur and a fourth heart sound were present. He had right upper quadrant abdominal tenderness and digital clubbing. Laboratory studies revealed anemia, unremarkable creatinine levels, and elevated liver enzymes (Table). Results of 3 sets of bacterial blood cultures and 1 set of fungal blood cultures were negative. Transthoracic and transesophageal echocardiograms demonstrated a VSD with bidirectional shunting and a mobile mitral valve echodensity. We

Table. Laboratory results for patient with infective endocarditis					
caused by Bartonella rochalimae, United States*					
Test	Result	Reference range			
Leukocytes, K/µL	4.8	4.5-11.0			
Hemoglobin, g/dL	10.9	12.6-17.4			
Platelets, K/μL	168	153–367			
Creatinine, mg/dL	0.77	0.66-1.25			
AST, units/L	84	17–59			
ALT, units/L	58	0–49			
Alkaline phosphatase, units/L	109	38–126			
Total bilirubin, mg/dL	0.6	0.3–1.2			
CRP, mg/dL	3.7	<u><</u> 1.0			
ESR, mm/h	81	0–15			
4th-generation HIV antigen	Nonreactive	Nonreactive			
and antibody test					
Coxiella burnetii Phase 2 IgM	1:32	Negative			
<i>C. burnetii</i> Phase 2 IgG	1:128	Negative			
<i>C. burnetii</i> Phase 1 IgM	Negative	Negative			
C. burnetii Phase 1 IgG	1:16	Negative			
Brucella antibody	<1:20	<1:20			
agglutination					
<i>Chlamydia pneumoniae</i> IgM	<1:20	<1:20			
<i>C. pneumoniae</i> IgG	1:512	<1:64			
C. trachomatis IgM	<1:20	<1:20			
C. trachomatis IgG	1:128	<1:64			
<i>C. psittaci</i> IgM	<1:20	<1:20			
<i>C. psittaci</i> IgG	1:512	<1:64			
<i>Bartonella henselae</i> IgG	>1:1024	Unknown			
<i>B. henselae</i> IgM	1:64	Unknown			
<i>B. quintana</i> IgG	>1:1024	Unknown			
<i>B. quintana</i> IgM	<1:16	Unknown			

*Bold text indicates abnormal values. ALT, alanine transaminase; AST, aspartate transferase; CRP, C-reactive protein; ESR, erythrocyte sedimentation rate.

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SARS-CoV-2 Infection in Beaver Farm, Mongolia, 2021

Appendix

Screening by RT-qPCR

The viral RNA was extracted using the NucleoSpin RNA Virus Mini kit (Macherey-Nagel, Germany; https://www.mn-net.com/us) from nasal swabs and saliva samples. The extracted RNA samples were screened by RT-qPCR (orf1b/nsp14 region) (*1*).

Whole genome sequencing analysis

First, RNA was reverse transcribed to cDNA using the Ion Torrent[™] NGS Reverse Transcription Kit per the manufacturer's instructions. Next, the amplification of the targeted genome and the multiplexed barcoded libraries construction was performed using the Ion AmpliSeq SARS-CoV-2 Insight Research Assay GS, Chef-ready kit (Thermo Fisher Scientific, https://www.thermofisher.com/us). The kit consists of two primer pools targeting 237 amplicons of 125–275 bp, covering more than 99% of the SARS-CoV-2 genome. The combined library pool thus generated was quantified and further sequenced on a Ion S5 system (Thermo Fisher Scientific).

The run was pre-processed using the torrent suite software to remove the primers and adapters. Then, the raw reads were quality-filtered using fastq-mcf v1.04.676 (ea-utils), and their quality was assessed with FastQC (v. 011.5). De Novo Assemblies were performed using SPAdes (v3.11.1). Using the Denovo assembly's contigs, BLAST searches identified a sequence from a Mongolian human SARS-CoV-2 isolate (GenBank # ON008302) as the most relevant reference. After mapping the cleaned raw reads against the reference sequence using BWA (v0.7.17), SAMtools (v1.11) was used to generate Mpileup files, and variant calling was

performed using BCFtools (v1.9). The consensus sequences produced with vcfutils.pl (VCFtools v0.1.16) and seqtk (v1.3.106) were compared to the Denovo assemblies using Mafft (ver. 7.467) and Aliview. We obtained five consensus sequences that were subjected to genotyping, mutation detection and phylogenetic analysis. The amino acid substitutions were analyzed using Nextclade and GISAID. After manually trimming the encoding sequences in both ends, a phylogenetic tree was constructed by the Maximum-likelihood method using MEGA 7.0 software with 100 bootstrap repeats. The phylogenetic tree was visualized and annotated using iTOL.

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