converged toward the same result: the convalescent-phase serum from the cat contained immunoglobulins against SARS-CoV-2, which were absent from the serum from control cats. These antibodies target several distinct viral proteins, and they caused a total neutralizing effect up to a much higher dilution than those from the owner’s serum. This household cat was therefore productively infected with the SARS-CoV-2 virus excreted by its owner, and the infection caused a nonfatal but nevertheless severe disease, mainly of the respiratory system (Videos 2–6).

Public health officials are still learning about SARS-CoV-2, but no current evidence indicates that pets play a role in spreading the virus. Therefore, taking measures against companion animals that may compromise their welfare is not justified.

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Naturally occurring human-to-animal transmission of severe acute respiratory syndrome (SARS) coronavirus was reported during 2003 when viral RNA was detected in oropharyngeal and rectal swab specimens from healthy domestic cats in a housing estate at the center of a large SARS cluster in Hong Kong, China; infections were confirmed serologically (1). Susceptibility of cats to infection with this virus and transmission between cats were demonstrated experimentally (2). Pulmonary pathologic changes, similar to those for humans with SARS, developed in infected cats, but the cats remained asymptomatic (2,3).

These findings informed the current precautionary strategy of the Agriculture, Fisheries and Conservation Department of Hong Kong to quarantine mammalian pets from households with confirmed human coronavirus disease (COVID-19) or their close contacts (defined as a person who had face-to-face contact for >15 min with a person who had confirmed SARS Coronavirus-2 [SARS-CoV-2] infection) in a holding facility, when alternative care was unavailable. Pets are swabbed for SARS-CoV-2 testing and confined until reverse transcription PCR (RT-PCR) results are negative on 2 consecutive oc-

SARS-CoV-2 in Quarantined Domestic Cats from COVID-19 Households or Close Contacts, Hong Kong, China

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We tested 50 cats from coronavirus disease households or close contacts in Hong Kong, China, for severe acute respiratory syndrome coronavirus 2 RNA in respiratory and fecal samples. We found 6 cases of apparent human-to-feline transmission involving healthy cats. Virus genomes sequenced from 1 cat and its owner were identical.

1These authors contributed equally to this article.
Findings for 2 infected dogs have been reported (4). We report testing results for cats.

Swab (nasal, oral, rectal) specimens and feces collected from nonsedated cats after admission were tested for SARS-CoV-2 RNA at the Agriculture, Fishries and Conservation Department Veterinary Laboratory by using a commercial RT-PCR targeting the partial envelope and RNA-dependent RNA polymerase genes (Molbiol Lightmix; TIB MOLBIOL, https://www.tib-molbiol.com). This PCR does not

Figure. Phylogenetic analysis of SARS-CoV-2 full genome from an infected cat and the human index case-patient, Hong Kong, China. A virus sequenced directly from a tiger in a zoo in United States was included in this analysis. Virus genome alignment was prepared and manually trimmed at genome 5′ and 3′ ends for low-alignment quality. A resulting alignment of 29,655 nt was analyzed by using PhyML (http://www.atgc-montpellier.fr) and the generalized time reversible nucleotide substitution model. Branch support identified by using the fast approximate likelihood ratio test are shown at major nodes. The Hong Kong feline virus from cat 1 and that of its owner are shown in red. Canine and human viruses from Hong Kong, including from the dogs’ owners (HK_case 163 and HK_case 85) are shown in blue. Numbers along branches are bootstrap values. Scale bar indicates nucleotide substitutions per site. SARS-CoV-2, severe acute respiratory syndrome coronavirus 2.
show cross-reactivity with these genes from enteric coronavirus (5).

For samples with positive or equivocal results, confirmatory quantitative RT-PCRs targeting non-structural protein 4, nonstructural protein 16, nucleoprotein, and membrane genes were performed at the World Health Organization COVID-19 Reference Laboratory at the University of Hong Kong (4). Animals with positive results were evaluated by repeated sampling to monitor viral shedding by RT-PCR. Serologic analysis was used selectively.

We sampled 50 cats during February 11–August 11, 2020. Time from onset of COVID-19 symptoms in owners to first sampling of their cats was available for 21 owners of 35 cats and ranged from 3 to 15 (median 8, interquartile range 4) days. SARS-CoV-2 RNA was detected in samples from 6 (12%) of 50 cats (Table, https://wwwnc.cdc.gov/EID/article/26/12/20-2786-T1.htm).

The first positive case (cat 1) was from a household that had 3 persons with confirmed cases of COVID-19; their symptoms (fever, cough, or shortness of breath) started on March 20, 29, and 30, 2020. Their 7-year-old, female, domestic shorthair cat was examined by a veterinarian at admission on day 1 (March 30), and reported to be clinically healthy. Nasal, oral, and rectal swab specimens collected on day 1 were positive for SARS-CoV-2 RNA; viral nucleoprotein gene copy numbers were log_{10} 6.3/mL, log_{10} 5.6/mL, and 3.2 log_{10}/mL, respectively.

Attempts to culture virus from day 1 samples on Vero E6 (ATCC CRL-1586) cells as described (4) were unsuccessful. Viral RNA was detected in oral swab specimens for 8 days and nasal swab specimens for 11 days, but rectal swab specimens were negative after day 1 (Table).

We performed serologic analysis to detect neutralizing antibodies by using a 90% plaque reduction neutralization test for SARS-CoV-2 (4). The result was positive for the only serum sample collected (on day 19) and had titer ≥1:320 (4).

Viral genomes from cat 1 and 1 owner were sequenced by using a MiSeq Sequencing Platform (Illumina, https://www.illumina.com) after reverse transcription of viral RNA and multiple, overlapping, ≥2-kb PCRs that targeted the viral genome (4). Both genome sequences (29,830 nt sequenced; 99.8% of the genome) were identical (Figure) and deposited in GenBank (accession no. MT628701).

Four of the other 5 positive cats were from confirmed COVID-19–infected households, and 1 indoor-only cat belonged to a close contact who was not confirmed to be infected. Time from onset of COVID-19 symptoms in owners to animal sampling was known for 3 cats (5, 11, and 8 days); 1 had equivocal envelope gene PCR results but was positive by a novel surrogate virus neutralization test (6). Signs of disease did not develop in any infected cats, consistent with experimental feline infections, which are also usually subclinical (7; A. Bosco-Lauth et al., unpub. data).

COVID-19–like signs have been reported in domestic cats naturally infected with SARS-CoV-2 in other countries (8). In addition, 4 tigers and 3 lions with respiratory signs in a zoo in New York, New York, USA, were confirmed to be shedding SARS-CoV-2 in feces (S.L. Bartlett et al., unpub. data). Susceptibility to SARS-CoV-2 might differ between felid species.

SARS-CoV-2 RNA persisted longest in nasal secretions in 1 case for 11 days at low levels. Viral RNA was detected in nasal washes from kittens experimentally inoculated with SARS-CoV-2 for 8–9 days, after which sampling was stopped (7). Cats acquiring infection from being cohoused with experimentally infected cats shed virus in respiratory secretions longer (7 days) than directly inoculated cats (5 days) (8).

Although feline-to-human transmission is theoretically possible, we did not find any evidence of this transmission. The timeline of infection in cat 1 and the finding of an identical SARS-CoV-2 genome sequence in a human from the same household is consistent with human-to-animal transmission. In support of these findings, the cat had no outdoor access.

More widespread serologic surveillance of cats in contact with COVID-19 patients is warranted to determine the prevalence of human-to-cat transmission. Some infected cats might have stopped shedding virus before being quarantined because viral shedding periods as short as 3 days have been reported in experimentally infected cats (7).

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Lack of Susceptibility to SARS-CoV-2 and MERS-CoV in Poultry

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We challenged chickens, turkeys, ducks, quail, and geese with severe acute respiratory syndrome coronavirus 2 or Middle East respiratory syndrome coronavirus. We observed no disease and detected no virus replication and no serum antibodies. We concluded that poultry are unlikely to serve a role in maintenance of either virus.

Coronaviruses of animals periodically transmit to humans (1), as recently occurred with severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). SARS-CoV-2 was recognized in December 2019 in cases of atypical pneumonia in hospitalized patients in Wuhan, China. The virus is a novel betacoronavirus, related to the now-eradicated severe acute respiratory syndrome coronavirus (SARS-CoV) from 2003, with which SARS-CoV-2 has 82% identity across the genome (2). SARS-CoV-2 is highly transmissible among humans and particularly virulent for elderly persons and those with certain underlying health conditions. Multiple studies have examined the susceptibility of domestic animals to SARS-CoV-2 to establish the risk for zoonotic transmission; 2 studies have shown chickens and Pekin ducks were not susceptible to infection (3,4).

Middle East respiratory syndrome coronavirus (MERS-CoV), another coronavirus of high concern associated with zoonotic infection, was first detected in patients with severe acute lower respiratory tract disease in Saudi Arabia in 2012. MERS-CoV causes lower respiratory disease, similar to the SARS-CoVs (5). Unlike SARS-CoV-2, MERS-CoV transmits poorly to humans and does not exhibit sustained human-to-human transmission; however, it has a high case fatality rate of ≈30%. Although the MERS-CoV case count is low, human cases continue to be reported, therefore there is a possibility for the virus to adapt to humans.

Based on sequence similarity, the closest relatives of SARS-CoV-2 and MERS-CoV are believed to be bat betacoronaviruses (6); the sequence difference...