possibility that dromedaries may not be the ultimate natural reservoir (i.e., the long-term host of a pathogen of an infectious disease). Topography (i.e., mountain chains) may limit camel movements from the Middle East or Africa to Central Asia, although such interchange certainly occurred centuries ago as a consequence of the silk-trade routes through southern Kazakhstan. The only known recent imports to Kazakhstan are dromedaries (Arvana breed), brought from Turkmenistan for cross-breeding with Bactrians to improve milk production (8). The findings that MERS-CoV is not universally endemic in dromedaries raises the hypothesis that certain species of bats or some other animal, the environment, or both, may constitute a maintenance community and be the true natural reservoir of MERS-CoV and that the virus spills over to camels and is maintained within camels for varying periods of time. Further studies on the epidemiology of MERS-CoV infection among camelids from central Asia are warranted.

This work was supported by a research grant from the US National Institute of Allergy and Infectious Diseases, National Institutes of Health, Department of Health and Human Services (contract no. HHSN272201500006C) and a commissioned grant from the Health and Medical Research Fund, Food and Health Bureau, Government of the Hong Kong Special Administrative Region.

References

Address for correspondence: Eve Miguel, CIRAD, AGIRs, Avenue Agropolis, 34398 Montpellier CEDEX 5, France; email: eve.miguel@cirad.fr

Novel Reassortant Avian Influenza A(H5N1) Virus in Human, Southern Vietnam, 2014

Ikuyo Takayama,1 Nguyen Trung Hieu,1 Masayuki Shirakura, Mina Nakauchi, Seiichiro Fujisaki, Hitoshi Takahashi, Shioi Nagata, Nguyen Thanh Long, Takato Odagiri, Masato Tashiro, Tsutomu Kageyama

Author affiliations: National Institute of Infectious Diseases, Tokyo, Japan (I. Takayama, M. Shirakura, M. Nakauchi, S. Fujisaki, H. Takahashi, S. Nagata, T. Odagiri, M. Tashiro, T. Kageyama); Pasteur Institute, Ho Chi Minh City, Vietnam (N.T. Hieu, N.T. Long)

DOI: http://dx.doi.org/10.3201/eid2203.151360

To the Editor: The first case of human infection with highly pathogenic avian influenza A(H5N1) virus in Vietnam was reported in December 2003 (1), and >120 human cases were confirmed through 2013, with a high case-fatality rate (2). In 2013, clade 2.3.2.1a/c H5N1 viruses circulated widely in poultry across the country, although clade 1.1.1/1.1.2 H5N1 viruses predominated in poultry from the Mekong Delta region to central Vietnam (3,4).

In 2014, two cases of human infection with A(H5N1) virus were identified in southern Vietnam. One case was associated with a clade 1.1.2 reassortant virus, A/Vietnam/14012902/2014 (Global Initiative on Sharing

1These first authors contributed equally to this article.
All Influenza Data [GISAID; http://www.gisaid.org] accession nos. EPI624919–EPI624926, which had been previously detected in Cambodia and Vietnam (5,6). We isolated the virus from the other case, performed phylogenetic analysis to identify the clade of this virus, and identified a novel virus that had undergone gene reassortment.

The case-patient was a 52-year-old man who lived in Binh Phuoc Province (140 km northeast of Ho Chi Minh City). On January 11, 2014, he experienced mild fever and general fatigue; high fever developed on January 13. He was hospitalized with dyspnea on January 16 and died 2 days later. He was not given antiviral drug treatment. Dead poultry infected with H5N1 viruses were found scattered near his house during January 1–16, and he buried his 2 dead chickens on January 5. H5N1 virus infection was detected in the patient’s throat swab specimen by real-time reverse transcription PCR at the Pasteur Institute in Ho Chi Minh City. Virus was isolated by inoculating the throat swab specimens into 10-day-old embryonated chicken eggs; the resulting isolate, A/Vietnam/14011801/2014 (GISAID accession nos. EPI624911–EPI624918), then underwent gene sequencing. The 8 viral genes were amplified with SuperScript III Reverse Transcriptase Kit (Fisher Scientific, Pittsburg, PA, USA) and Phusion High-Fidelity DNA Polymerase (New England BioLabs, Ipswich, MA, USA) with specific paired primers, according to the manufacturer’s instructions, and sequenced on an ABI 3730 automated sequencer with Big-Dye Terminator Cycle Sequencing reagents (Applied Biosystems, Foster City, CA, USA). Whole genome sequence was determined.

By gene sequencing analysis, A/Vietnam/14011801/2014 was found to have the multibasic cleavage site of hemagglutinin (HA) protein, which indicates highly pathogenic avian influenza A(H5N1) viruses, and was shown to predict binding specificity to an avian α2,3 sialic acid receptor. The neuraminidase gene possessed no amino acid substitutions associated with decreased antiviral activity, nor did the virus have amino acid substitutions associated with increased adaptation, virulence, infectivity, or transmissibility in mammalian hosts, including the E627K and D701N mutations in polymerase basic protein 2 (7).

Phylogenetic analyses of the 8 viral genes of A/Vietnam/14011801/2014 were performed by using databases (GISAID and the Influenza Virus Resource, National Center for Biotechnology Information, Bethesda, MD, USA; http://www.ncbi.nlm.nih.gov/genomes/FLU/FLU.html) that contained complete sequences of viral genomes belonging to clades 1.1.1, 1.1.2, and 2.3.2.1 a/b/c, most of which were collected in Vietnam, particularly after 2012. Neighbor-joining and Kimura 2-parameter methods were implemented by using MEGA version 5.0 software (http://www.megasoftware.net). Reliability of the phylogenetic analysis was tested by using 1,000 bootstrap replications. Lineages of the HA gene were defined by using previously described criteria (8). Lineages of the other 7 genes were defined by using criteria and nomenclature of Nguyen et al. (9).

The HA of A/Vietnam/14011801/2014 belonged to clade 2.3.2.1c (online Technical Appendix Figure, panel A, http://wwwnc.cdc.gov/EID/article/23/3/15-1360-Techapp.pdf). The neuraminidase, polymerase basic proteins 1 and 2, and polymerase acid protein genes of this virus were also derived from respective lineages of ancestor clade 2.3.2.1c (online Technical Appendix Figure, panels B–E). However, nucleoprotein, matrix, and nonstructural genes were classified as lineages of ancestor clade 2.3.2.1a (online Technical Appendix Figure, panels F–H) and differed from the gene lineages of almost all clade 2.3.2.1c viruses isolated from poultry in Vietnam. As reported in the Influenza Virus Resource, 2 viruses collected in Vietnam in December 2013 (A/muscovy duck/Long An/43/2013 and A/muscovy duck/Long An/46/2013) were similar reassortant viruses of clade 2.3.2.1c (Figure). However, the ancestor of the nonstructural gene lineage of these 2 viruses is clade 2.3.2.1c, which differs from A/Vietnam/14011801/2014.
The differences indicate that A/Vietnam/14011801/2014 is a novel reassortant virus between clades 2.3.2.1a and 2.3.2.1c, between clades 1.1.2 and 2.3.2.1c, or both (Figure). This novel reassortant virus has not been reported in poultry in Vietnam, although novel reassortants between clade 1.1.2 and clade 2.3.2.1a viruses have been detected in Vietnam since 2013 (i.e., A/Vietnam/VP13-28H/2013, GISAID accession nos. EPI624927–EPI624934; and A/Vietnam/14012902/2014) (6). These novel reassortment viruses were first identified in human, animal, and environmental samples in Cambodia in 2013 (5). Other novel gene reassortments in clade 2.3.2.1 viruses have been previously reported (10), and new clade 2.3.4.4 viruses have been observed in Vietnam since 2014.

As multiple clade viruses co-circulate, reassortment events occur frequently in Vietnam. Continuous surveillance of avian influenza A(H5N1) viruses, not only in humans but also in poultry and wild birds, is needed for infection control measures during epidemics of these viruses.

This study was partially supported by Grants-in-Aid for Emerging and Reemerging Infectious Diseases from the Ministry of Health, Labour and Welfare, Japan.

References


Address for correspondence: Tsutomu Kageyama, Influenza Virus Research Center, National Institute of Infectious Diseases, 4-7-1 Gakuen, Musashimurayama-shi, Tokyo 208-0011, Japan; email: tkage@nih.go.jp

Mycobacterium arupense as an Emerging Cause of Tenosynovitis

Fiorella Krapp Lopez, Madeline Miley, Babafemi Taiwo

Author affiliations: Northwestern University, Chicago, Illinois, USA (F. Krapp Lopez, B. Taiwo); Northwestern Memorial Hospital, Chicago (M. Miley)

DOI: http://dx.doi.org/10.3201/eid2203.151749

To the Editor: Mycobacterium arupense was identified as a novel species within the M. terrae complex with close similarity to M. nonchromogenicum (1). Since then, 8 cases describing clinically notable disease have been published (2–8), including 5 cases of tenosynovitis. We report M. arupense tenosynovitis in an immunocompromised person who received the selective interleukin (IL) 1β-inhibitor canakinumab.

In July 2014, a 62-year-old man sought treatment at the emergency department, Northwestern Memorial Hospital (Chicago, Illinois, USA), after 1 week of pain and swelling in the right hand. During the previous 5 years, he had received multiple immunomodulatory drugs for treatment of natural killer cell deficiency, hyper–IL-6 syndrome, recurrent polychondritis, and Sweet syndrome. His medications were prednisone (42.5 mg/d), intravenous immunoglobulin (400 mg/kg monthly), and subcutaneous canakinumab (180 mg every 8 weeks, which began 3 weeks before onset of symptoms).

His first symptom was a tender red nodule on the right palm that increased in size and became extremely tender over the following week (Figure, panels A, B). He did not recall any trauma and denied fever or chills. No improvement was seen after he received oral linezolid for 5 days. A
Novel Reassortant Avian Influenza A(H5N1) Virus in Human, Southern Vietnam, 2014

Technical Appendix
**Technical Appendix Figure.** Phylogenetic analyses of the genes of the avian influenza A(H5N1) virus described in this study: A) hemagglutinin (HA) gene; B) neuraminidase (NA) gene; C) polymerase basic (PB) 2 gene; D) PB1 gene; E) polymerase acid (PA) protein gene; F) nucleoprotein (NP) gene; G) matrix (M) gene; and H) nonstructural (NS) gene. Trees were constructed by using the neighbor-joining method. The evolutionary distances were computed by using the Kimura 2-parameter method. Bootstrap values were calculated from 1,000 replicates and values >60% are shown next to branches. Sequences of the A(H5N1) virus isolated in this study are in bold font and inside a box. Viruses are colored on the basis of their hemagglutinin clade: 1.1.1 in light blue; 1.1.2 in green; 2.3.2.1a in red; 2.3.2.1b in purple; and 2.3.2.1c in dark blue. Scale bars indicate nucleotide substitutions per site.