New Reassortant Clade 2.3.4.4b Avian Influenza A(H5N6) Virus in Wild Birds, South Korea, 2017–18

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DOI: https://doi.org/10.3201/eid2410.180461

We isolated new reassortant avian influenza virus (H5N6) viruses from feces of wild waterfowl in South Korea during 2017–18. Phylogenetic analysis suggested that reassortment occurred between clade 2.3.4.4b H5N8 and Eurasian low pathogenicity avian influenza viruses circulating in wild birds. Dissemination to South Korea during the 2017 fall migratory season followed.

Clade 2.3.4.4 H5 highly pathogenic avian influenza viruses (HPAIVs) have evolved by reassortment with different neuraminidase (NA) and internal genes of prevailing low pathogenicity avian influenza viruses (LPAIVs) and other HPAIVs to generate new genotypes and further evolved into genetic subgroups A–D since 2014 (1). Among these, subgroups A and B viruses were disseminated over vast geographic regions by migratory wild birds (2,3). Subgroup B influenza A(H5N8) viruses were detected in Qinghai Lake, China, and Uvs-Nuur Lake, Russia, during May–June 2016 (Qinghai/Uvs-like), followed by the identification of reassortant viruses in multiple Eurasian countries (4–6). Recently, subgroup B H5N6 viruses were isolated from birds in Greece during February 2017 and England, Germany, the Netherlands, Japan, and Taiwan during winter 2017–18 (7,8).

During December 2017–January 2018 in South Korea, we isolated 6 H5N6 HPAIVs from 231 fecal samples of wild birds collected from the banks of the Cheongmi-cheon River (37°06′56.9″N, 127°25′18.3″E) and 34 from 222 fecal samples collected from the banks of the Gokgyo-cheon River (36°45′12.3″N, 127°07′12.7″E) (online Technical Appendix 1, https://wwwnc.cdc.gov/EID/article/24/10/18-0461-Techapp1.pdf). These wild bird habitats are wintering sites of migratory waterfowl, including mallard (Anas platyrhynchos), spot-billed duck (Anas poecilorhyncha), Mandarin duck (Aix galericulata), and common teal (Anas crecca). The Gokgyo-cheon River is a major habitat site for Mandarin ducks, and numerous HPAIVs were detected in fecal samples from Mandarin ducks during 2011, 2015, and 2016 (9). We identified avian influenza virus–positive fecal samples from 38 Mandarin ducks and 2 mallards, based on DNA barcoding technique (10). We performed full-length genome sequencing and comparative phylogenetic analysis on 19 of the 40 isolates (online Technical Appendix 1; online Technical Appendix 2, https://wwwnc.cdc.gov/EID/article/24/10/18-0461-Techapp2.xlsx).

All H5N6 isolates shared high nucleotide sequence identities in all 8 gene segments (99.58%–100%) and were identified as HPAIVs based on the presence of multiple basic amino acids at the HA proteolytic cleavage site (PLREKRRKR/G). Searches of the GISAID (https://www.gisaid.org) and BLAST (https://blast.ncbi.nlm.nih.gov/Blast.cgi) databases indicated that all 8 genotypes had the highest nucleotide identity with A/Great Black-backed Gull/Netherlands/1/2017 (Netherlands/1) clade 2.3.4.4 subgroup B H5N6 strain from December 2017 (99.17%–99.79%), rather than subgroup B H5N6 viruses from Japan and Taiwan collected during December 2017 (97.18%–99.27%).

In phylogenetic analysis, we identified 2 genotypes of subgroup B H5N6 viruses (online Technical Appendix 1 Figures 1, 2): genotypes B.N6.1 and B.N6.2. The genotype B.N6.1 viruses were identified from South Korea, Japan, Taiwan, Greece, and the Netherlands (Netherlands/1 strain), and the genotype B.N6.2 viruses were detected from England, Germany, and the Netherlands. For genotype B.N6.1, all genes except NA clustered with H5N8 HPAIV of previously reported genotypes, H5N8-NL cluster I in the Netherlands (6), Ger-11-16 in Germany (5), and Duck/Poland/82a/16-like in Italy (4). The NA gene clustered with LPAIVs circulating in wild birds in Eurasia and separated into 2 clusters, suggesting the potential for >2 independent reassortment events between H5N8 virus and unidentified wild bird origin N6 segments. Consistent clustering of South Korea isolates with the Netherlands/1 strain in maximum-likelihood (ML) phylogenies for each gene supported by high ML bootstrap values (86–100) suggests their close relationship. The genotype B.N6.2 viruses
had different polymerase basic 2 (PB2) and polymerase acidic (PA) genes from genotype B.N6.1. The polymerase basic 2 gene probably originated from other LPAIVs, and a polymerase acidic gene originated from H5N8-NL cluster II genotype (6,7). The phylogenetic network and ML analysis suggest that H5N6 viruses have evolved from subgroup B H5N8 viruses into 3 independent pathways, detected in Greece, Europe/South Korea, and Japan/Taiwan (online Technical Appendix, Figures 2,3).

The time of most recent common ancestry (tMRCA) for each gene of genotype B.N6.1 influenza A(H5N6) viruses isolated during winter 2017–18 in Eurasia, except for the NA gene, ranged from January 2016 to October 2016, suggesting that genotype B.N6.1 viruses diverged during the previous year. The tMRCA of the NA gene was September 2015 (95% highest posterior density August 2014–August 2016). The tMRCA of the NA gene has wide 95% highest posterior density range because only a few recent N6 genes of LPAIVs were available in databases for analysis. The tMRCA for each gene of H5N6 HPAIVs identified in South Korea ranged from July through September 2017, suggesting that ancestors of these viruses emerged among wild birds during or after summer 2017, possibly at the breeding and molting sites in the Palearctic region (Table; online Technical Appendix Figure 4). Detection of H5N6 HPAIV from fecal samples of wild birds in South Korea during the 2017–18 wintering season and our phylogenetic analysis suggest that the viruses had moved through wild birds during the fall migration season.

On the basis of our data and migratory pattern of birds, we estimate that H5N6 viruses possibly descended from H5N8 viruses circulating during 2016–17, reaching breeding regions of wild birds during early 2017, followed by dissemination into Europe and East Asia during the fall migration. Enhanced surveillance in wild birds is needed for early detection of new introductions of HPAIV and to trace the transmission route of HPAIV.

This work was funded by Konkuk University in 2015.

### About the Author

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### References


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### Table. Time to most recent common ancestor for each gene segment of genotype B.N6.1 influenza A(H5N6) viruses isolated in South Korea, December 2017–January 2018*

<table>
<thead>
<tr>
<th>Gene</th>
<th>South Korea isolates, node 1</th>
<th>South Korea and Europe isolates, node 2</th>
<th>South Korea, Europe, Japan, Taiwan, and Greece isolates, node 3</th>
</tr>
</thead>
</table>

*Nodes of the temporally structured maximum clade credibility phylogenetic tree (online Technical Appendix Figure 4, http://wwwnc.cdc.gov/EID/article/24/10-0461-Techapp1.pdf). HA, hemagglutinin; HPD, highest posterior density; M, matrix; NA, neuraminidase; NP, nucleoprotein; NS, nonstructural; PA, polymerase acidic; PB, polymerase basic.
Clinical Isolation and Identification of Haematospirillum jordaniae

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DOI: https://doi.org/10.3201/eid2410.180548

A clinical case study involving a man (35–49 years of age) with wounds to his lower right extremity. An isolate was sent to the Delaware Public Health Laboratory for confirmatory testing by genetic analysis of the 16S gene. Testing identified the isolate as a novel genus and species, Haematospirillum jordaniae.

The Centers for Disease Control and Prevention (CDC) recently recognized a human pathogen, Haematospirillum jordaniae, as a new bacterial genus and species (1). H. jordaniae is a gram-negative, spiral-shaped aerobe in the family Rhodospirillaceae (1). Before the pathogen’s identification, this organism was isolated 14 times from 10 different states. These organisms were received by CDC’s Special Bacteriology Reference Laboratory, Division of High-Consequence Pathogens and Pathology, National Center for Emerging and Zoonotic Infectious Diseases, over the course of 10 years (February 2003–October 2012) before the initial publication. Here we report the clinical identification of H. jordaniae, a potential emerging infectious pathogen, by the Delaware Public Health Laboratory (DPHL) and describe possible routes of infection.

1Location redacted for patient confidentiality.

A man (35–49 years of age) sought care at a hospital emergency department in September 2016 with worsening pain, redness, and swelling of the right lower extremity, which had an open wound on the right shin. The patient stated that he had finished a project cutting stones 3 weeks earlier where the water and stone chips were hitting his right leg. The patient ignored swelling, erythema, and fever he experienced around the same period when the stonecutting occurred and decided not to seek medical intervention until he was no longer able to ignore the symptoms.

The patient was found to have cellulitis of the right leg and sepsis, and he was admitted to the hospital for treatment. On the same day, he was started on intravenous clindamycin, and blood cultures were collected. The infectious disease physician discontinued clindamycin and placed the patient on vancomycin, ciprofloxacin, and aztreonam because of concern for potential infection with Pseudomonas, Aeromonas, or methicillin-resistant Staphylococcus aureus. Growth was observed in both sets of aerobic bottles, the first after 2 days, 15 hours, and the second after 4 days, 18 hours. Blood cultures grew gram-negative bacilli, leading the infectious disease physician to remove vancomycin but continue with ciprofloxacin and aztreonam. The patient improved before identification of the pathogen and was discharged on oral ciprofloxacin.

Gram stain of the organism revealed small, curved, gram-negative bacilli, resembling Campylobacter. Subcultures from the bottles did not grow aerobically until 48 hours after incubation on sheep’s blood agar and chocolate agar. The organisms did not grow on MacConkey agar or when incubated anaerobically on blood agar. Although it was suspected that the organism might be a Campylobacter species, it did not grow in a microaerophilic environment. Biochemical testing revealed the organism as oxidase-positive, indole-negative, catalase-positive, and urease-negative (using urea agar). The isolate was sent to the DPHL for identification.

DPHL received a grown isolate from the submitting hospital 8 days after the patient’s admission. The organism was cultured in the same manner as described previously; phenotypic tests were consistent with the findings of the submitting hospital. Initial identification was performed at DPHL, but the pathogen could not be determined because the organism was not part of the instrument’s stored database.

The isolate’s DNA was extracted and amplified by using general methods described in numerous publications. The sample generated an 845-bp portion of the 16S region. The assembled sequence was uploaded to open-source rRNA databases for comparison, including GenBank BLAST (https://blast.ncbi.nlm.nih.gov/Blast.cgi), the Ribosomal Database Project (https://rdp.cme.msu.edu/), and MicrobeNet (https://microbenet.cdc.gov). Samples most closely matched to Novispirillum itersonii strain...
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Technical Appendix

Methods

Virus Isolation

Fecal samples were tested for influenza A virus by egg inoculation using 9–1-day-old specific-pathogen-free embryonating chicken eggs. After a 72-h incubation period, allantoic fluids were harvested from an axenic medium, and undiluted allantoic fluids were tested for hemagglutinin (HA) activity. For the definitive diagnosis of influenza A virus, RNA was extracted from HA-positive allantoic fluid using an RNeasy Kit (QIAGEN, Valencia, CA, USA) according to the manufacturer’s instructions. The extracted RNA was tested for the influenza A virus matrix gene by real-time reverse transcription PCR (RT-PCR) as described previously (1). The hosts of the positive fecal sample were identified as Mandarin ducks (Aix galericulata) and mallards (Anas platyrhynchos) using DNA barcoding techniques as previously described (2).

Full-genome Sequencing

Full-length genome sequencing was performed for 19 of 40 isolates (5 from Cheongmi-cheon River collected on 22 December 2017, 13 from Gokgyo-cheon River collected on 23 December 2017, and 1 from Gokgyo-cheon River collected on 18 January 2018). For molecular analysis, RNA was extracted using the RNeasy kit (QIAGEN) according to the manufacturer’s instructions. The 8 genes of each virus were amplified using 1-step RT-PCR. The RT-PCR amplicons (2 µg) of all 8 gene segments was used to prepare Ion Fragment sequencing libraries (Life Technologies, Carlsbad, CA, USA) according to the manufacturer’s instructions. Briefly, amplicons were loaded onto beads, and emulsion PCR was conducted before sequencing with an Ion 318 chip on an Ion Torrent Personal Genome Machine. De novo and directed assembly of genome sequences were performed using Geneious R9 software (http://www.geneious.com). We
provided the genome sequences in the EpiFlu database of Global Initiative on Sharing All Influenza Data (GISAID; https://www.gisaid.org), under accession nos. EPI1190336–40, EPI1190342–61, EPI1190381, EPI1190386, EPI1190392, EPI1190410, EPI1190418, EPI1190426, EPI1190426, EPI1190433, EPI1190446, EPI1190498, EPI1190503, EPI1190508, EPI1190514–19, EPI1200474–509, EPI1200511–87.

**Phylogenetic Analysis**

For phylogenetic analysis, complete coding regions were aligned using MAFFT (https://mafft.cbrc.jp/alignment/software/), and manual editing and tree reconstruction were conducted using Geneious 8 software (https://www.geneious.com/). The sequence of clade 2.3.4.4 H5 highly pathogenic avian influenza viruses representing each subgroup and low pathogenic avian influenza virus strains related to new reassortment of H5N6 highly pathogenic avian influenza viruses in 2017–18 were selected from GISAID Epiflu and GenBank (https://www.ncbi.nlm.nih.gov/genomes/FLU) database as a reference for phylogenetic tree analysis. A maximum-likelihood tree was estimated by the MEGA7 software using the Hasegawa-Kishino-Yano model (3) of nucleotide substitution with gamma-distributed rate variation among sites with 4 rate categories. Statistical analysis of the phylogenetic tree was performed by bootstrap analysis on 1,000 replicates. The percentages of replicate trees (>70%) in which the associated taxa clustered together in the bootstrap test are shown next to the branches.

To better visualize the genetic relatedness of viruses, HA gene was analyzed using the median joining method implemented by NETWORK ver. 5.0 with epsilon set to 0. Nucleotide sequences of complete protein coding sequence region of subgroup B H5N6 and H5N8 identified from Asia and Europe in 2016 and 2017 (n = 250) were retrieved from GISAID Epiflu database on February 13, 2018. The sequences were pruned by using the online software cd-hit (4) at 97% homology to filter-out 66 representative sequences. The nucleotide sequences of HA segment obtained in this study (n = 19) were added to the dataset. A total of 85 nt sequences were aligned using MAFFT, and alignments were manually edited in Geneious 8 software (https://www.geneious.com/).

For estimation of the time to the most recent common ancestor, Bayesian analysis was performed for all 8 gene segments using BEAST version 1.8.4 (http://beast.bio.ed.ac.uk). We used an Hasegawa-Kishino-Yano substitution model with 4 gamma categories and specified an
uncorrelated lognormal relaxed clock and GMRF Bayesian skyride tree prior for each segment. A Markov chain Monte Carlo method was used with 50 million chain lengths to draw inference under this model. A maximum clade credibility tree with common ancestor height was generated for each dataset using TreeAnnotator version 1.8.4 after 10% burn-in. The maximum clade credibility trees were visualized using the program FigTree v1.4.2.

References


**Technical Appendix 1 Figure 1.** Proposed reassortment events leading to the new subgroup B influenza A(H5N6) viruses. Eight gene segments in each of the schematic virus particles are arranged from top to bottom to represent the PB2, PB1, PA, HA, NP, NA, M, and NS genes. The circled HPAI viruses are colored in red, and unknown LPAI viruses are colored in black. For each gene segment, different phylogenetic groupings are in different colors. The mean time to the most recent common ancestor of HA gene are shown at the node. HA, hemagglutinin; HPAI, highly pathogenic avian influenza; HPD, highest posterior density; LPAI, low pathogen avian influenza; M, matrix; NA, neuraminidase; NP, nucleoprotein; NS, nonstructural; PA, polymerase acidic; PB, polymerase basic; tMCRA, time to the most recent common ancestor.
Technical Appendix 1 Figure 2. Maximum-likelihood phylogenetic tree of the PB2, PB1, PA, HA, NP, NA, M, and NS genes. The new subgroup B H5N6 isolates are colored according to the isolated location (South Korea, red; Europe, blue; Japan/Taiwan, green; Greece, purple). The percentages of replicate trees (>70%) in which the associated taxa clustered together in the bootstrap test (1,000 replicates) are shown next to the branches. Each cluster is labeled according to classification of Pohlmann et al. (5) and Beerens et al. (6). HA, hemagglutinin; M, matrix; NA, neuraminidase; NP, nucleoprotein; NS, nonstructural; PA, polymerase acidic; PB, polymerase basic. Scale bars indicate nucleotide substitutions per site.
Technical Appendix 1 Figure 3. Median-joining phylogenetic network of subgroup B H5 highly pathogenic avian influenza viruses. The median-joining network was constructed from the HA gene. It includes all the most parsimonious trees linking the sequences. Each unique sequence is represented by a circle sized relative to its frequency in the dataset. Branch length is proportional to the number of mutations. Isolates are colored according to the location. HA, hemagglutinin.
Technical Appendix 1 Figure 4. Temporally structured maximum clade credibility phylogenetic tree (years on the horizontal axis) of the PB2, PB1, PA, HA, NP, NA, M, and NS genes of subgroup B HPAIV. The new subgroup B H5N6 Isolates are colored according to the isolated location (South Korea, red;
Europe, blue; Japan/Taiwan, green; and Greece, purple). The horizontal bars indicate the 95% highest posterior density (HPD) intervals of the most recent common ancestor. HA, hemagglutinin; HPD, highest posterior density; M, matrix; NA, neuraminidase; NP, nucleoprotein; NS, nonstructural; PA, polymerase acidic; PB, polymerase basic. Scale bars indicate nucleotide substitutions per site.