Influenza A viruses (IAVs) are one of the most consequential pathogens of birds and terrestrial mammals. IAVs (family Orthomyxoviridae) are enveloped, segmented, negative-sense, single-stranded RNA viruses, subtyped based on the surface glycoproteins hemagglutinin (H) and neuraminidase (N) embedded in the virus envelope. With the exceptions of H17N10 and H18N11 IAVs in bats, all other existing combinations of H (n = 16) and N (n = 9) subtypes are found in wild aquatic and shore birds, their natural reservoir hosts (1).

Phylogenetically, IAVs can be separated into 2 genetically distinct lineages, Eurasia (EA) and North America (NA), because of the fidelity of migratory waterfowl to distinct flyways (2). However, in areas where migratory flyways overlap, there are, although rare, IAVs with gene segments from both NA and EA lineages. Surveillance studies in Alaska (United States) and Newfoundland (Canada) documented a preponderance of IAVs with whole-genome segments from EA lineage viruses and only rarely, reassortant IAVs with EA and NA gene segment constellations (3,4). In 2014, along the Pacific flyway corridor in Canada and the United States, goose/Guangdong/1/96 (Gs/GD) lineage H5N8 virus was detected in wild waterfowl (5). The Gs/GD virus reassorted with NA lineage IAVs, subsequently resulting in devastating losses among commercial poultry flocks in Canada and the United States.

Wild bird–origin IAVs can breach the host species barrier and cause outbreaks involving several terrestrial mammals. After crossing the species barrier, some IAVs can become established in new hosts and circulate independently of their reservoir hosts. Different species of marine mammals are susceptible to IAV infection through exposure at haul-out sites, where they might come into contact with wild birds, infected sympatric marine mammal species, terrestrial wildlife, including mink and river otters, and through direct exposure to infected humans or waterfowl in rehabilitation facilities (6). Previous outbreaks and individual case reports have shown that seals are susceptible to and have died as a result of infection with H10N7, H3N8, H7N7, H4N6, and other IAVs (6).

**The Study**

A live adult male harbor seal weighing 49 kg washed ashore at Combers Beach near Tofino, British Columbia (BC), Canada, but died before he could be captured alive. On June 1, 2021, the seal was brought to the Animal Health Center Laboratory (Abbotsford, BC, Canada) for necropsy. The seal was in moderate body and fair postmortem condition. The most

We isolated a novel reassortant influenza A(H10N7) virus from a harbor seal in British Columbia, Canada, that died from bronchointerstitial pneumonia. The virus had unique genome constellations involving lineages from North America and Eurasia and polymerase basic 2 segment D701N mutation, associated with adaptation to mammals.

Author affiliations: University of Manitoba Department of Animal Science, Winnipeg, Manitoba, Canada (Y. Berhane); Canadian Food Inspection Agency, Winnipeg (Y. Berhane, O. Lung, C. Embury-Hyatt, W. Xu); Western College of Veterinary Medicine, Saskatchewan, Canada (Y. Berhane, S. Raverty); Ministry of Agriculture, Food and Fisheries, Abbotsford, British Columbia, Canada (T. Joseph, S. Raverty); University of Manitoba Department of Biological Sciences, Winnipeg (O. Lung); Fisheries and Aquaculture Management, Vancouver, British Columbia, Canada (P. Cottrell); University of British Columbia Institute for Oceans and Fisheries, Vancouver (S. Raverty)

DOI: https://doi.org/10.3201/eid2807.212302
notable findings from gross examination were serosanguinous fluid within the thoracic cavity and focally extensive hemorrhage and edema from the skin to the visceral pleura along the right midlateral aspect of the thorax, with focal visceral to parietal pleural adhesion. Histopathology revealed necrotizing bronchitis and bronchiolitis, peribronchial lymphoid hyperplasia, alveolar histiocytosis, and perivascular lymphoplasmacytic cuffing. Immunohistochemistry testing identified IAV viral antigen within the bronchiolar-associated lymphoid tissue of 2 bronchioles in more severely affected areas of the lungs and in situ hybridization confirmed IAV RNA (Figure 1). Aerobic cultures of the lungs, hilar lymph node, brain, intercostal skeletal muscle, and small intestine yielded variable mixed growth of *Streptococcus phocae* and *Serratia liquefaciens* with no fungal growth from the lung. PCR testing of pooled tissues proved positive for consensus influenza virus and mollicutes (we confirmed *Mycoplasma* spp., but were unable to speciate) and negative for canine distemper virus.

We extracted RNA samples from the hilar lymph node, thymus, spleen, and lungs, and thoracic fluid tested positive on a real-time reverse transcription PCR assay based on IAV matrix (M) genes, as described elsewhere (7). We isolated all IAV PCR-positive samples in embryonated specific pathogen–free chicken eggs from lung tissue samples only. We amplified all 8 viral gene segments from original lung specimens and isolates as described elsewhere (8), then purified amplified reverse transcription PCR products using a QIAGEN QIAquick PCR purification kit (https://www.qiagen.com) and determined the concentration of the amplicons used for sequencing using an

![Figure 1. Immunohistochemistry testing for influenza A virus antigen in lung tissue of an adult male harbor seal, British Columbia, Canada. Viral antigen (arrow) was detected by immunohistochemistry within the bronchiolar-associated lymphoid tissue but not in adjacent lung parenchyma (*). Viral RNA could also be detected in bronchiolar-associated lymphoid tissue by in situ hybridization (inset, pink areas). Scale bar = 50 μm.](image-url)

| Table. Similarity of all 8 gene segments and their associated proteins for influenza A(H10N7) virus from a harbor seal in British Columbia, Canada, to closest matches in GenBank* |
|-------------------------------------------------|---------------------------------|-----------------|------------------|
| Segment | Similarity† (%) | Closest match | GenBank accession no. | Lineage |
|---------|-----------------|-----------------|------------------|---------|
| PB2 | 2,225/2,280 (97.59) | A/ruddy turnstone/New Jersey/A13-2822/2013 | MH500897 | North America |
| PB1 | 2,240/2,274 (98.50) | A/Mallard/WA/AH0042257S.2.A/2015(H6N8) | MN254524 | North America |
| PA | 2,117/2,151 (98.42) | A/blue-winged teal/Texas/A11-3220/2011(H3N8) | CY205900 | North America |
| H | 1,613/1,686 (95.67) | A/duck/Bangladesh/24035/2014 (H10N1) | KY616787 | Eurasia |
| NP | 1,467/1,497 (98.00) | A/duck/Hokkaido/W90/2007(H10N7) | LC121485 | Eurasia |
| N | 1,372/1,416 (96.89) | A/duck/Mongolia/742/2015(H10N7) | LC121446 | Eurasia |
| M | 973/982 (99.07) | A/Mallard duck/Alberta/486/2019(H1N1) | MT624434 | North America |
| NS1/NEP | 821/844 (97.27) | A/duck/Hokkaido/56/2017(H12N2) | MK929497 | Eurasia |

*H, hemagglutinin; M, matrix; N, neuraminidase; NP, nucleoprotein; NS1/NEP, nonstructural protein 1/nuclear export protein; PA, acidic polymerase; PB, basic polymerase.
†Nucleotide matches at identical sites.
Figure 2. Maximum-likelihood phylogenetic tree of influenza A virus subtype H10 hemagglutinin gene from an adult male harbor seal, British Columbia, Canada (red text), and reference sequences. Phylogenetic analyses were based on the full-length nucleotide sequence of the hemagglutinin gene of strains representing the H10 subtype (n = 1,512). The evolutionary relationship was inferred using RAxML (https://github.com/stamatak/standard-RAxML) based on the general time-reversible model with 1,000 bootstrap replicates. For purposes of clarity, some clades are collapsed, and colors are assigned to indicate the origin of the gene: blue for North America, green for Asia, and yellow for Europe. The tree was drawn to scale; branch lengths are measured in number of substitutions per site.
In our study of a harbor seal infected with a novel reassortment H10N7 IAV containing a unique constellation of NA and EA lineage gene segments, we found no conclusive evidence of where and when the reassortment occurred. However, the seal was recovered at a location within the Pacific avian flyway, where Gs/GD lineage H5N8 virus was detected in 2014 and later reassorted with NA lineage viruses to create novel reassortant H5N2 and H5N1 IAVs that were responsible for large outbreaks in domestic poultry in Canada and the United States (5). Harbor seals share habitat with seabirds and shorebirds, and the seal in our study may have been infected through spillover directly from infected birds as happened in cases reported elsewhere (6,8). In previous outbreaks in seals, the proximate cause of death was attributed to secondary or opportunistic Mycoplasma spp. (14), which were also detected in this case, or bacterial infection. Finally, because novel IAVs in marine mammals have been shown to be potentially zoonotic (15), IAV circulation among seals could have important public health implications for humans and other mammals.

Acknowledgments
The authors thank Tamiko Hisanaga, Matt Suderman, Estella Moffat, and Katherine Handel for their technical assistance. The authors also thank staff from the necropsy, bacteriology, virology, and molecular diagnostics sections of the British Columbia Animal Health Centre for their technical assistance.

This study was supported by the Public Health Agency of Canada and Canadian Food Inspection Agency to support genetic and antigenic monitoring of influenza A virus in animals.

About the Author
Dr. Berhane is a virologist with the National Centre for Genetic and Antigenic Monitoring of Influenza A Virus in Animals.

References
1. Wu Y, Wu Y, Tefsen B, Shi Y, Gao GF. Bat-derived influenza-like viruses H17N10 and H18N11. Trends Microbiol. 2014;22:183–91. https://doi.org/10.1016/j.tim.2014.01.010
2. Webster RG, Bean WJ, Gorman OT, Chambers TM, Kawakita Y. Evolution and ecology of influenza A viruses. Microbiol Rev. 1992;56:152–79. https://doi.org/10.1128/MI.56.1.152-179.1992
3. Huang Y, Wille M, Benkaroun J, Munro H, Bond AL, Fifield DA, et al. Perpetuation and reassortment of gull

Conclusions
In our study of a harbor seal infected with a novel reassortment H10N7 IAV containing a unique constellation of NA and EA lineage gene segments, we found no conclusive evidence of where and when the reassortment occurred. However, the seal was recovered at a location within the Pacific avian flyway, where Gs/GD lineage H5N8 virus was detected in 2014 and later reassorted with NA lineage viruses to create novel reassortant H5N2 and H5N1 IAVs that were responsible for large outbreaks in domestic poultry in Canada and the United States (5). Harbor seals share habitat with seabirds and shorebirds, and the seal in our study may have been infected through spillover directly from infected birds as happened in cases reported elsewhere (6,8). In previous outbreaks in seals, the proximate cause of death was attributed to secondary or opportunistic Mycoplasma spp. (14), which were also detected in this case, or bacterial infection. Finally, because novel IAVs in marine mammals have been shown to be potentially zoonotic (15), IAV circulation among seals could have important public health implications for humans and other mammals.

Acknowledgments
The authors thank Tamiko Hisanaga, Matt Suderman, Estella Moffat, and Katherine Handel for their technical assistance. The authors also thank staff from the necropsy, bacteriology, virology, and molecular diagnostics sections of the British Columbia Animal Health Centre for their technical assistance.

This study was supported by the Public Health Agency of Canada and Canadian Food Inspection Agency to support genetic and antigenic monitoring of influenza A virus in animals.

About the Author
Dr. Berhane is a virologist with the National Centre for Genetic and Antigenic Monitoring of Influenza A Virus in Animals.

References
1. Wu Y, Wu Y, Tefsen B, Shi Y, Gao GF. Bat-derived influenza-like viruses H17N10 and H18N11. Trends Microbiol. 2014;22:183–91. https://doi.org/10.1016/j.tim.2014.01.010
2. Webster RG, Bean WJ, Gorman OT, Chambers TM, Kawakita Y. Evolution and ecology of influenza A viruses. Microbiol Rev. 1992;56:152–79. https://doi.org/10.1128/MI.56.1.152-179.1992
3. Huang Y, Wille M, Benkaroun J, Munro H, Bond AL, Fifield DA, et al. Perpetuation and reassortment of gull

Conclusions
In our study of a harbor seal infected with a novel reassortment H10N7 IAV containing a unique constellation...
influenza A viruses in Atlantic North America. Virology. 2014; 456-457:353–63. https://doi.org/10.1016/j.virol.2014.04.009

4. Ramey AM, Reeves AB, Sonsthagen SA, TeSlaa JL, Nashold S, Donnelly T, et al. Dispersal of H9N2 influenza A viruses between East Asia and North America by wild birds. Virology. 2015;482:79–83. https://doi.org/10.1016/j.virol.2015.03.028

5. Ramey AM, DelLiberto TJ, Berhane Y, Swayne DE, Stallknecht DE. Lessons learned from research and surveillance directed at highly pathogenic influenza A viruses in wild birds inhabiting North America. Virology. 2018;518:55–63. https://doi.org/10.1016/j.virol.2018.02.002

6. Venkatesh D, Bianco C, Núñez A, Collins R, Thorpe D, Reid SM, et al. Detection of H3N8 influenza A virus with multiple mammalian-adaptive mutations in a rescued Grey seal (Halichoerus grypus) pup. Virus Evol. 2020;6:veaa016. https://doi.org/10.1093/ve/veaa016

7. Weingartl HM, Berhane Y, Hisanaga T, Neufeld J, Kehler H, Embury-Hyatt C, et al. Genetic and pathobiologic characterization of pandemic H1N1 2009 influenza viruses from a naturally infected swine herd. J Virol. 2010;84:2245–56. https://doi.org/10.1128/JVI.00786-11

8. Zhou B, Donnelly ME, Scholes DT, St George K, Hatta M, Kawakoa Y, et al. Single-reaction genomic amplification accelerates sequencing and vaccine production for classical and Swine origin human influenza a viruses. J Virol. 2009;83:10309–13. https://doi.org/10.1128/JVI.01109-09

9. Xu W, Weese JS, Ojkic D, Lung O, Handel K, Berhane Y. Phylogenetic inference of H3N2 canine influenza A outbreak in Ontario, Canada in 2018. Sci Rep. 2020;10:6309. https://doi.org/10.1038/s41598-020-63278-z

10. Bogs J, Kalthoff D, Veits J, Pavlova S, Schwemmler M, Mänz B, et al. Reversion of PB2-627E to -627K during replication of an H5N1 Clade 2.2 virus in mammalian hosts depends on the origin of the nucleoprotein. J Virol. 2011;85:10691–8. https://doi.org/10.1128/JVI.00786-11

11. Li J, Ishaq M, Prudence M, Xi X, Hu T, Liu Q, et al. Single mutation at the amino acid position 627 of PB2 that leads to increased virulence of an H5N1 avian influenza virus during adaptation in mice can be compensated by multiple mutations at other sites of PB2. Virus Res. 2009;144:123–9. https://doi.org/10.1016/j.virusres.2009.04.008

12. Gabriel G, Dauber B, Wolff T, Planz O, Klenk HD, Stech J. The viral polymerase mediates adaptation of an avian influenza virus to a mammalian host. Proc Natl Acad Sci U S A. 2005; 102:18590–5. https://doi.org/10.1073/pnas.0507415102

13. Fan S, Deng G, Song J, Tian G, Suo Y, Jiang Y, et al. Two amino acid residues in the matrix protein M1 contribute to the virulence difference of H5N1 avian influenza viruses in mice. Virology. 2009;384:28–32. https://doi.org/10.1016/j.virol.2008.11.044

14. van Kuppeveld FJ, van der Logt JT, Angulo AF, van Zoest MJ, Quint WG, Niesters HG, et al. Genus- and species-specific identification of mycoplasmas by 16S rRNA amplification. Appl Environ Microbiol. 1992;58:2606–15. https://doi.org/10.1128/AEM.58.8.2606-2615.1992

15. van den Brand JMA, Wolfsen P, Herfst S, Bodewes R, Pfankuche VM, van de Bildt MWG, et al. Influenza A (H10N7) virus causes respiratory tract disease in harbor seals and ferrets. PLoS One. 2016;11:e0159625. https://doi.org/10.1371/journal.pone.0159625

Address for correspondence: Stephen Raverty, Animal Health Center–Pathology, 1767 Angus Campbell Rd, Abbotsford, Vancouver, BC V3G 2M3, Canada; email: stephen.raverty@gov.bc.ca