**Human Adenovirus 7d Strains Associated with Influenza-Like Illness, New York, USA, 2017–2019**

Daryl M. Lamson, Adriana Kajon, Michael Popowich, Meghan Fuschino, Kirsten St. George

Author affiliations: New York State Department of Health, Albany, New York, USA (D.M. Lamson, M. Popowich, M. Fuschino, K. St. George); Lovelace Respiratory Research Institute, Albuquerque, New Mexico, USA (A. Kajon)

DOI: https://doi.org/10.3201/eid2605.200116

Human adenovirus 7d is a respiratory pathogen capable of causing acute respiratory disease of variable severity. Phylogenetic analysis of whole-genome sequences of 15 strains isolated from cases of influenza-like-illness during 2017–2019 demonstrated the circulation of 2 distinct clades of genomic variant 7d in colleges in New York, USA.

Human adenovirus genome (HAdV) type 7d (HAdV-7d) was first detected in the United States in December 2013 in Oregon, in association with acute respiratory disease (ARD) requiring hospitalization (1). In 2014, it was detected in more persons with ARD in Oregon and in Illinois in 2 adults with severe pneumonia (2). Circulation of this genome type, probably imported from East Asia, where its re-emergence was first documented in 2009 (3), has been detected since 2013 in other locations and settings in the United States in association with ARD of variable severity in children and adults, including in a long-term care facility in New Jersey (4); the US Marine Corps Officer Candidates School in Quantico, Virginia (5); and the University of Maryland (https://www.cnn.com/2018/11/21/health/university-of-maryland-death-adenovirus).

The recent increased detection of this distinct genomic variant of HAdV-7 in the United States and its association with severe disease manifestations has prompted public health laboratories to be more vigilant about detection of HAdVs in association with ARD. The Wadsworth Center of the New York State Department of Health (Albany, NY, USA), in collaboration with the Lovelace Respiratory Research Institute (Albuquerque, NM, USA), has monitored the prevalence of respiratory HAdVs detected in New York state since 2012. We characterized 15 HAdV-7 strains isolated from influenza virus–negative respiratory specimens collected from students with influenza-like illness at colleges in Tompkins, Albany, and Clinton counties in New York during the 2016–17, 2017–18, and 2018–19 influenza seasons. We used next-generation whole-genome sequencing and phylogenetic analysis to investigate possible epidemiologic connections among the New York college outbreaks and to monitor the dispersion of this reemerging variant within the United States.

Clinical specimens were initially tested for influenza viruses using a real-time reverse transcription PCR diagnostic panel and subsequently tested for HAdV as previously described (6). Samples testing positive for HAdV were processed for molecular typing by amplification and sequencing of hyper-variable regions 1–6 of the hexon gene (7) and for virus isolation by conventional culture with standard techniques. Intracellular genomic HAdV DNA was purified from infected A549 cells exhibiting cytopathic effect and used for initial genetic characterization by restriction enzyme analysis and for next-generation sequencing with Illumina MiSeq (Illumina, https://www.illumina.com), as previously described (6). We aligned genomic sequences generated in this study and reference sequences from GenBank in Geneious Pro R11 using MAFFT (https://www.geneious.com). We constructed a maximum-likelihood tree using MEGA6 (8). We also generated in silico digests of the genomic sequences in Geneious Pro R11. We annotated all sequences using VAPiD (9) and uploaded to GenBank (accession nos. MH921831–42, MK405661, and MN638755–56).

Initial digestion of viral DNA with endonucleases BamHI, BstEII, HpaI and subsequent in silico digestion of the corresponding complete genomic sequences identified all isolated strains as corresponding to genome type 7d (data not shown). Phylogenetic analysis of whole-genome sequencing demonstrated the co-circulation of 2 distinct clades of HAdV-7d strains in New York, even within the same county, in the sampled time period (Figure). The first clade comprised 11 strains isolated during March 2017–February 2019. Nine of these were genetically related to strains isolated in 2016–2017 in New Jersey (NJ/5644/2016 and NJ/6295/2017), and 2 were more closely related to a 2014 strain isolated in Oregon (OR/CDC2014012.949/2014). The second clade comprised 4 strains isolated during March and April 2017 that were closely related to strain DG01 isolated in 2011 in China and genetically similar to a sample isolated in 2017 from Virginia (VA/5677/2017). Both clades share a common ancestor, strain GZ6965, isolated in Guangdong, China, in 2011.
Our phylogenetic analysis provides strong evidence of ≥2 introductions of genomic variant 7d into New York state. Spatiotemporal analysis of a larger genomic sequence dataset with better representation of strains isolated in other US states, as well as in other countries, over an extended period is necessary to more accurately track the introduction of lineages and follow their dispersion.

As in our previous studies and those of others (4–6,10), these findings highlight the importance of HAdV as a causative agent of ARD in civilian communities and the value of college student populations for sentinel surveillance of HAdV activity. These data also provide another example of the power of whole-genome sequencing analysis for the epidemiologic investigation of HAdV-associated disease (9,10).

All HAdV-7d strains examined in this study were isolated from persons with influenza-like illness. Numerous recent publications report severe and even fatal cases of ARD in association with this genomic variant. All HAdV-7d strains sequenced thus far are indistinguishable by restriction enzyme analysis but not identical when examined at the whole-genome level. Characterization of more strains of diverse origins and associated disease and thorough mining of sequence data is needed to identify candidate determinants of virulence for HAdV-7. Host-related and other environmental risk factors are likely to contribute to the level of susceptibility, clinical presentation, and outcome of the associated disease.

Acknowledgments
We thank the staff of the Wadsworth Center Virology Laboratory for performing initial viral culture and molecular testing on clinical specimens. The specimens analyzed in this study were obtained as part of the surveillance program of the Enhanced Laboratory Capacity program of the Centers for Disease Control and Prevention under Cooperative Agreement no. NU50CK000423; this work was therefore partly supported by the Centers for Disease Control and Prevention.
About the Author
Mr. Lamson is Assistant Director of Special Projects in the Laboratory of Viral Diseases, Wadsworth Center, New York State Department of Health. His research focuses on molecular epidemiology and sequence analysis of adenoviruses and enteroviruses with next-generation whole-genome and di-deoxy Sanger sequencing.

References
1. Scott MK, Chommanard C, Lu X, Appelgate D, Grenz L, Schneider E, et al. Human adenovirus associated with severe respiratory infection, Oregon, USA, 2013–2014. Emerg Infect Dis. 2016;22:1044–51. https://doi.org/10.3201/eid2206.151898
2. Kajon AE, Ison MG. Severe infections with human adenovirus 7d in 2 adults in family, Illinois, USA, 2014. Emerg Infect Dis. 2016;22:730–3. https://doi.org/10.3201/eid2204.151403
3. Zhao S, Wan C, Ke C, Seto J, Dehghan S, Zou L, et al. Re-emergent human adenovirus genome type 7d caused an acute respiratory disease outbreak in southern China after a twenty-one year absence. Sci Rep. 2014;4:7365. https://doi.org/10.1038/srep07365
4. Killerby ME, Rozwadowski F, Lu X, Cauldric-Grimes M, McHugh L, Haldeman AM, et al. Respiratory illness associated with emergent human adenovirus genome type 7d, New Jersey, 2016–2017. Open Forum Infect Dis. 2019;6:ofz17. https://doi.org/10.1093/ofid/ofz17
5. Bautista-Gogel J, Madsen CM, Lu X, Sakhitvel SK, Froh I, Kamau E, et al. Outbreak of respiratory illness associated with human adenovirus type 7 among persons attending Officer Candidates School, Quantico, Virginia, 2017. J Infect Dis. 2019. https://doi.org/10.1093/infdis/jizk60
6. Lamson DM, Kajon A, Shudt M, Girouard G, St George K. Detection and genetic characterization of adenovirus type 14 strain in students with influenza-like illness, New York, USA, 2014–2015. Emerg Infect Dis. 2017;23:1194–7. https://doi.org/10.3201/eid2307.161730
7. Okada M, Ogawa T, Kubonoya H, Yoshizumi H, Shinozaki K. Detection and sequence-based typing of human adenoviruses using sensitive universal primer sets for the hexon gene. Arch Virol. 2007;152:1–9. https://doi.org/10.1007/s00705-006-0842-8
8. Tamura K, Stecher G, Peterson D, Filipski A, Kumar S. MEGA6: Molecular Evolutionary Genetics Analysis version 6.0. Mol Biol Evol. 2013;30:2725–9. https://doi.org/10.1093/molbev/mst197
9. Shean RC, Makhsous N, Stoddard GD, Lin MJ, Greninger AL. VAPiD: a lightweight cross-platform viral annotation pipeline and identification tool to facilitate virus genome submissions to NCBI GenBank. BMC Bioinformatics. 2019;20:48. https://doi.org/10.1186/s12859-019-2606-y
10. Kajon AE, Lamson DM, Bair CR, Lu X, Landry ML, Menegus M, et al. Adenovirus type 4 respiratory infections among civilian adults, northeastern United States, 2011–2015. Emerg Infect Dis. 2018;24:201–9. https://doi.org/10.3201/eid2402.171407

Address for correspondence: Daryl M. Lamson, Wadsworth Center—Viral Diseases, 120 New Scotland Ave, Albany, NY 12201-0509, USA; email: daryl.lamson@health.ny.gov

Risk for Transportation of Coronavirus Disease from Wuhan to Other Cities in China

Zhanwei Du,1 Lin Wang,1 Simon Cauchemez, Xiaoke Xu, Xianwen Wang, Benjamin J. Cowling, Lauren Ancel Meyers

Author affiliations: University of Texas at Austin, Austin, Texas, USA (Z. Du, L.A. Meyers); Institut Pasteur, Paris, France (L. Wang, S. Cauchemez); Dalian Minzu University, Dalian, China (X. Xu); Dalian University of Technology, Dalian (X. Wang); The University of Hong Kong, Hong Kong (B.J. Cowling); Santa Fe Institute, Santa Fe, New Mexico, USA (L.A. Meyers)

DOI: https://doi.org/10.3201/eid2605.200146

On January 23, 2020, China quarantined Wuhan to contain coronavirus disease (COVID-19). We estimated the probability of transportation of COVID-19 from Wuhan to 369 other cities in China before the quarantine. Expected COVID-19 risk is >50% in 130 (95% CI 89–190) cities and >99% in the 4 largest metropolitan areas.

In December 2019, a novel coronavirus, since named severe acute respiratory syndrome coronavirus 2, emerged in Wuhan, China (1), causing a respiratory illness that the World Health Organization has named coronavirus disease (COVID-19). On January 30, 2020, the World Health Organization declared the outbreak a public health emergency of international concern (2). By January 31, 2020, a total of 192 fatalities and 3,215 laboratory-confirmed cases had been reported in Wuhan; 8,576 additional cases were spread across >300 cities in mainland China, and 127 exported cases were reported in 23 countries spanning Asia, Europe, Oceania, and North America. The rapid global expansion, rising fatalities, unknown animal reservoir, and evidence of person-to-person transmission potential (3,4) initially resembled the 2003 SARS epidemic and raised concerns about global spread.

On January 22, 2020, China announced a travel quarantine of Wuhan and by January 30 expanded the radius to include 16 cities, encompassing a population of 45 million. At the time of the quarantine, China was already 2 weeks into the 40-day Spring Festival, during which residents and visitors make several billion trips throughout China to celebrate

1These first authors contributed equally to this article.