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Address for correspondence: Janina Trauth, University Hospital Gießen and Marburg GmbH, Section of Infectious Diseases, Klinikstr 33, 35392 Gießen, Germany; email: janina.trauth@innere.med.uni-giessen.de

Natural Reassortment of Eurasian Avian-Like Swine H1N1 and Avian H9N2 Influenza Viruses in Pigs, China

Wanying Sun,¹ Samuel S.M. Cheng,¹ Kristy N.T. Lam, Tsz C. Kwan, Ricky W.K. Wong, Leo H.K. Lau, Gigi Y.Z. Liu, Leo L.H. Luk, John K.C. Li, Haogao Gu, Malik Peiris, Leo L.M. Poon

Author affiliations: School of Public Health, The University of Hong Kong, Hong Kong, China (W. Sun, S.S.M. Cheng, K.N.T. Lam, T.C. Kwan, R.W.K. Wong, L.H.K. Lau. G.Y.Z. Liu, L.L.H. Luk, J.K.C. Li, H. Gu, M. Peiris, L.L.M. Poon); Centre for Immunology and Infection, Hong Kong (M. Peiris, L.L.M. Poon); HKU-Pasteur Research Pole, The University of Hong Kong, Hong Kong (M. Peiris, L.L.M. Poon)

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Several zoonotic influenza A viruses detected in humans contain genes derived from avian H9N2 subtypes. We uncovered a Eurasian avian-like H1N1 swine influenza virus with polymerase basic 1 and matrix gene segments derived from the H9N2 subtype, suggesting that H9N2 viruses are infecting pigs and reassorting with swine influenza viruses in China.

Swine are regarded as a mixing vessel for influenza A viruses (IAVs) (1). Avian, swine, and human IAVs can co-infect pigs and generate novel reassortants of zoonotic or pandemic potential. The emergence of pandemic H1N1 IAV (pH1N1), containing viral segments from avian, swine, and human viruses, highlighted the key role of pigs in contributing to IAV reassortment and evolution (2). Research in China also showed evidence of avian H5, H7, H9, and H10 influenza infections in pigs (3). Avian IAVs linked to human infection in this region contained internal genes derived from avian H9N2 viruses, indicating that the internal genes of the H9N2 virus might aid zoonotic transmission (4). We report detection of a swine IAV with polymerase basic (PB) 1 and matrix (M) gene segments of avian H9N2 origin.

In April 2021, we resumed monthly influenza surveillance program of imported pigs in a local slaughterhouse, which had been interrupted by COVID-19 outbreaks (5). We collected individual nasal swab samples (≈75 samples per visit), which we kept chilled in virus transport medium until they reached the laboratory. We then subjected swab samples to IAV isolation by using MDCK cells, as previously described (2). We identified cultures with cytopathic effect and tested them using a standard hemagglutination assay with turkey red blood cells. We tested hemagglutination-positive cultures with a universal influenza reverse transcription PCR assay specific for M segments (6). We studied samples that were positive for this reaction by using next-generation sequencing to deduce the full virus genomes (6).

During April 2021-February 2022, we collected a total of 829 porcine nasal swab samples (Table). We isolated 8 IAVs: 7 from August 2021 and 1 from September 2021. Virus sequences deduced from this study are available from GISAID (isolate nos. EPI_ISL_12471293-300). We compared those sequences with reference sequences (Appendix Table, https://wwwnc.cdc.gov/EID/article/28/7/22-0642-App1.pdf). IAVs detected in August 2021 were H3N2 viruses. The hemagglutinin (HA) and neuraminidase (NA) segments of those viruses were associated with human-like H3N2 swine influenza A virus; however, their internal gene segments all were derived from the pH1N1 lineage (Figure; Appendix Figures 1-6). These viruses were genetically not identical but highly similar. The influenza-positive pigs came from farms located in 2 provinces across southern China. Because this slaughterhouse followed a daily clearance policy requiring that all imported live pigs be slaughtered within 24 hours of admittance, our results suggest influenza transmission between pigs in the preslaughter transport chain outside Hong Kong. This

¹These authors contributed equally to this article.

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Year and month	No. nasal swabs	No. virus isolates	Isolation rate, %
2021			
Apr	60	0	0
May	75	0	0
Jun	75	0	0
Jul	75	0	0
Aug	75	7*	9.3
Sep	75	1†	1.3
Oct	79	0	0
Nov	85	0	0
Dec	80	0	0
2022			
Jan	75	0	0
Feb	75	0	0
Total	829	8	0.97

Table Swing influenza viruges detected in imported pige Ching April 2021 Echrugry 2022

†H1N1, pig was from imported from a province in southern China

H3N2 genotype was previously detected in pigs from Guangxi, China (7).

The swine H1N1 IAV that we isolated in September 2021, A/swine/HK/NS419/2021, a reassortant between multiple swine influenza lineages (Figure; Appendix Figures 1-6). The PB1 and M gene segments of this virus are of avian H9N2 virus subtype. This virus contains PB2, polymerase acidic, and NP gene segments derived from the pH1N1 lineage. Its HA and NA gene segments are of Eurasian avian-like H1N1 lineage, and its nonstructural gene segment is of a triple reassortant lineage. We further purified the isolated virus by using plaque assays to exclude the possibility of a mixed infection. We confirmed that all plaque-purified viral clones had an identical genotype.

The A/swine/HK/NS419/2021 isolate featured a PB1 gene segment of SH/F/98-like lineage and an M gene segment of G1-like H9N2 lineage (Figure). Similar PB1 and M sequences have been detected in zoonotic viruses in humans (Figure), PB1 in H10N8 and M in H7N9, but we did not find mutations known for mammalian host adaptation in these 2 segments. The encoded proteins of the PB1 and M gene segments that we isolated featured amino acid sequences rarely observed in mammalian and avian IAVs, including H9 (PB1, 97K, 156N, 397V, 535V, 688I, and 704T; M1, 31I and 46V; and M2, 25S). We could not determine whether these were random or adaptive mutations. The PB1 segment of avian H9N2 is highly compatible to other polymerase genes from mammalian IAVs (8). Such results suggest the need for further characterization of these mutations, particularly those in the PB1 gene.

A recent report in China discussed multiple Eurasian avian-like H1N1 swine influenza reassortants with internal genes derived from pH1N1 and triple reassortant lineages (9). One group of these reassortants (genotype 4) displayed a genotype similar to A/swine/HK/NS419/2021, the only exception being that the virus's PB1 and M gene segments were of pH1N1 lineage. That report showed that genotype 4 Eurasian avian-like swine IAVs can bind to human sialic acid receptors (i.e., α 2,3), enabling efficient virus replication in human airway epithelial cells, and achieve efficient aerosol transmission in ferrets (9). Serologic surveillance further showed that 10% of studied swine workers were positive for the genotype 4 reassortant (9). Our own sequence analyses suggest that some of the genotype 4 viruses and our Eurasian avian-like H1N1 viruses might share a common ancestry (e.g., A/swine/Shandong/1207/2016; Appendix Figures 1-6). Further risk assessment on the pandemic potential of this genotype and its reassortants is needed (10).

In summary, many zoonotic IAVs in humans have genes derived from H9N2 subtypes. Our results suggest that avian H9N2 IAVs are infecting swine and reassorting with swine IAVs, which indicates the need for continued monitoring of swine IAVs in both China and outlying regions.

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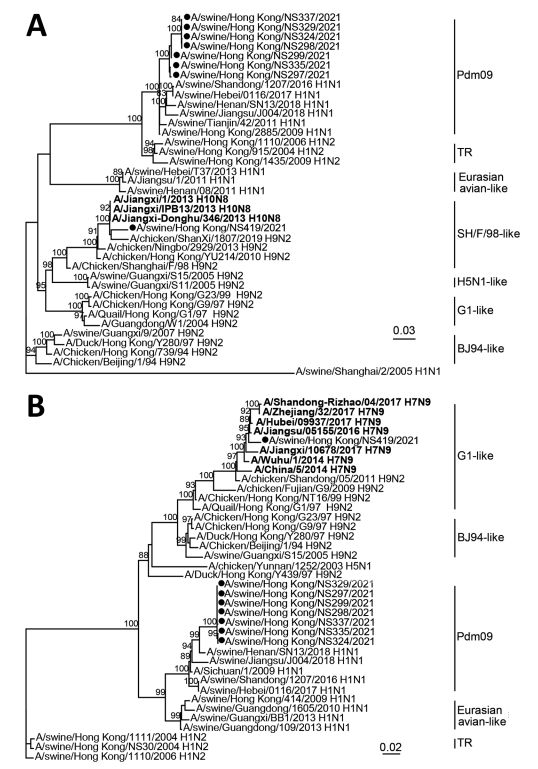


Figure. Phylogenetic tree of polymerase basic 1 (A) and matrix (B) gene sequences of swine influenza viruses from China and reference sequences. Bold indicates human H7N9 and H10N8 sequences. Viral sequences generated in this study (black circles) and those downloaded from public domains (Appendix Table, https://wwwnc.cdc.gov/EID/article/28/7/22-0642-App1.pdf) were aligned by using Muscle version 3.8 (http://www.drive5.com/muscle). Phylogenetic trees were constructed by IQ-TREE 1.6.12 (http://www.iqtree. org) by using the generalized time reversible plus gamma model. Major animal viral lineages are as shown. Bootstrap values ≥80% are shown. Scale bar indicates estimated genetic distance.

About the Author

Miss Sun is a postgraduate student at The University of Hong Kong, Hong Kong, China. Her primary research interest is bioinformatics.

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Address for correspondence: Malik Peiris or Leo Poon, School of Public Health, The University of Hong Kong, 7 Sassoon Road, Hong Kong, China; email: malik@hku.hk or llmpoon@hku.hk

Persistent SARS-CoV-2 Alpha Variant Infection in Immunosuppressed Patient, France, February 2022

Slim Fourati, Guillaume Gautier, Myriam Chovelon, Alexandre Soulier, Melissa N'Debi, Vanessa Demontant, Céline Kennel, Christophe Rodriguez, Jean-Michel Pawlotsky

Author affiliations: Hôpital Henri Mondor (AP-HP), Université Paris-Est, Créteil, France (S. Fourati, A. Soulier, M. N'Debi, V. Demontant, C. Rodriguez, J.-M. Pawlotsky); William Morey General Hospital, Chalon-sur-Saône, France (G. Gautier, M. Chovelon, C. Kennel)

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We describe persistent circulation of SARS-CoV-2 Alpha variant in an immunosuppressed patient in France during February 2022. The virus had a new pattern of mutation accumulation. The ongoing circulation of previous variants of concern could lead to reemergence of variants with the potential to propagate future waves of infection.

Immunosuppressed patients can have prolonged SARS-CoV-2 infection (1). Studies have reported the occurrence and selection of multiple mutations in the spike glycoprotein sequence in immunosuppressed patients with persistent SARS-CoV-2 infections (2–6). To date, intrahost mutations have been described essentially in the ancestral wild-type SARS-CoV-2 virus (3,5–8), especially during prolonged infection with variants of concern (VOCs) (9). Additional SARS-CoV-2 mutations in immunocompromised persons could enable increased virus transmissibility and immune evasion, shaping the emergence of new VOCs. We describe a new mutation accumulation pattern in SARS-CoV-2 Alpha virus in an immunosuppressed patient.

An 84-year-old woman with evolutive mantle cell lymphoma who was receiving maintenance rituximab and lenalidomide treatment was admitted to the hospital on May 17, 2021. She had asthenia, fever, and hypoxia (93% oxygen saturation). At admission (day 0), she tested positive for SARS-CoV-2 RNA (Figure). She had received 2 vaccine doses 84 and 66 days before admission. She did not have respiratory symptoms, but a chest computed tomography scan showed ground-glass opacities in her lungs. The patient was hospitalized and treated with corticosteroids for 10 days. She tested SARS-CoV-2-positive again on August 26, day 101 after her initial

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Appendix

egment	Strain name	Accession number
B2	A/swine/Fujian/204/2007	FJ536816
B2	A/swine/Wisconsin/30954/1976	CY036806
B2	A/swine/Shanghai/2/2005	FJ789824
B2	A/swine/Hong Kong/915/2004	GQ229273
B2	A/swine/Hong Kong/78/2003	GQ229313
B2	A/swine/Hong Kong/1562/2005	GQ229337
B2	A/swine/Hong Kong/1110/2006	GQ229377
B2	A/swine/Hong Kong/1435/2009	CY061650
B2	A/swine/Hong Kong/NS1583/2009	CY061722
B2	A/swine/Hong Kong/2299/2009	CY061730
B2	A/swine/Hong Kong/2314/2009	CY061738
32	A/swine/Hong Kong/NS1809/2009	CY061746
32	A/swine/Hong Kong/NS1810/2009	CY061754
32	A/swine/Hong Kong/2885/2009	CY061762
32	A/swine/Hong Kong/2886/2009	CY061770
32	A/swine/Hong Kong/2974/2009	CY061786
32	A/swine/Hong Kong/189/2010	CY061802
32	A/swine/Hong Kong/1795/1994	CY085102
32	A/swine/Hong Kong/8278/2001	CY085371
32	A/swine/Hong Kong/227/2002	CY085491
32	A/swine/Hong Kong/1248/2002	CY085523
32	A/swine/Hong Kong/1578/2003	CY085603
32	A/swine/Hong Kong/1111/2004	CY085659
2	A/swine/Hong Kong/729/2005	CY085731
32	A/swine/Hong Kong/72/2007	CY085811
32	A/swine/Hong Kong/NS252/2009	CY085995
32	A/swine/Guangdong/1605/2010	JN375017
32	A/swine/Guangdong/1623/2010	JN375023
32	A/swine/Shaanxi/s2/2012	JX963608
32	A/Jiangsu/1/2011	KF057091
B2	A/swine/Guangdong/109/2013	KP404209
B2	A/swine/Hunan/30/2013	KP404289
B2	A/swine/Hong Kong/299/1993	CY084982
32	A/swine/Hong Kong/1223/1993	CY085054
32	A/swine/Hebei/0116/2017	MN416337
B2	A/swine/Henan/SN13/2018	MN416362
B2	A/swine/Jiangsu/J004/2018	MN416364
B2	A/swine/Shandong/1207/2016	MN416383
31	A/Chicken/Hong Kong/G9/97	AF156416
31	A/Chicken/Hong Kong/G23/99	AF156417
31	A/Duck/Hong Kong/Y280/97	AF156419
31	A/Quail/Hong Kong/G1/97	AF156421
B1	A/Chicken/Hong Kong/739/94	AF156422
B1	A/Chicken/Beijing/1/94	AF156423
B1*	A/Jiangxi-Donghu/346/2013	EPI ISL 152846
31*	A/Jiangxi/1/2013	EPI_ISL_174265

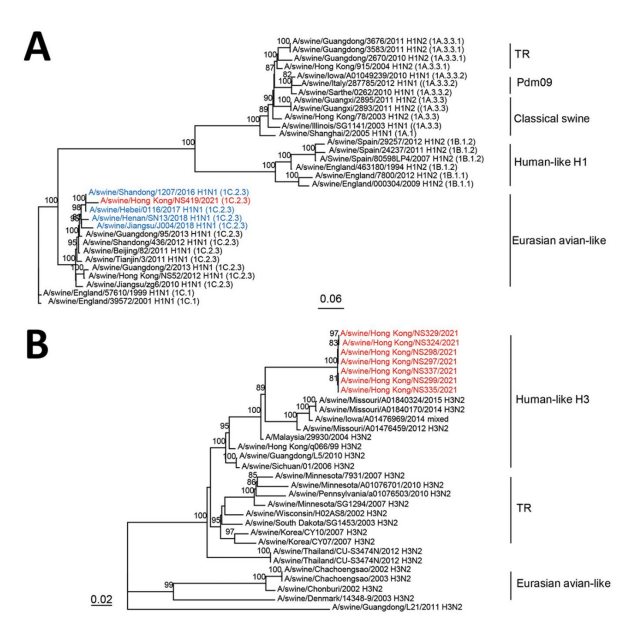
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PB1	A/swine/Guangxi/S15/2005	EU086331
°B1	A/swine/Hong Kong/915/2004	GQ229268
°B1	A/swine/Hong Kong/1110/2006	GQ229372
°B1	A/swine/Hong Kong/1435/2009	CY061651
°B1	A/swine/Hong Kong/2885/2009	CY061763
°B1	A/swine/Guangxi/9/2007	CY075044
'B1	A/Jiangsu/1/2011	KF057098
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B1	A/swine/Shandong/1207/2016	MN416470
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A 2A		CY061772
A 2A	A/swine/Hong Kong/2886/2009 A/swine/Hong Kong/2974/2009	
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Segment	Strain name	Accession number
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łA	A/swine/Jiangsu/zg6/2010	JN809152
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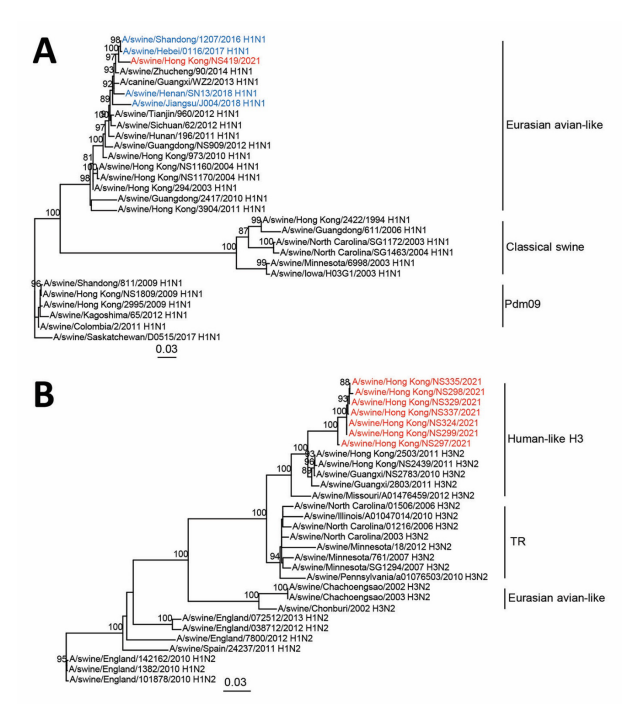
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IP	A/swine/Hong Kong/729/2005	CY085735
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IA	A/swine/Hong Kong/294/2003	CY086546
IA	A/swine/Hong Kong/NS1160/2004	CY086752
IA	A/swine/Minnesota/6998/2003	CY098486
IA	A/swine/Shandong/811/2009	JQ695889
IA	A/swine/Colombia/2/2011	KC703346
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Segment	Strain name	Accession number
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NA	A/swine/Hong Kong/973/2010	KM029513
NA	A/swine/Kagoshima/65/2012	AB911619
NA	A/swine/Hunan/196/2011	KP404286
NA	A/swine/Sichuan/62/2012	KP404382
NA	A/swine/Zhucheng/90/2014	KX264371
NA	A/swine/Hong Kong/2422/1994	CY087058
NA	A/canine/Guangxi/WZ2/2013	MG254094
NA	A/swine/Saskatchewan/D0515/2017	MK462308
NA	A/swine/Tianjin/960/2012	MN416762
NA	A/swine/Chachoengsao/2003	AB434346
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NA	A/swine/Chonburi/2002	AB573015
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M*	A/Hubei/09937/2017 H7N9	EPI ISL 285591
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M*	A/Jiangxi/10678/2017_H7N9	EPI_ISL_285246
M*	A/Shandong-Rizhao/04/2017_H7N9	EPI_ISL_285268
M*	A/wuhu/1/2014_H7N9	EPI_ISL_399682
M*	A/Zhejiang/32/2017_H7N9	EPI_ISL_285320
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M	A/Quail/Hong Kong/G1/97	AF156463
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NS	A/swine/Hong Kong/NS1583/2009	CY061729
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NS	A/swine/Hong Kong/2314/2009	CY061745
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NS	A/swine/Hong Kong/2885/2009	CY061769
NS	A/swine/Hong Kong/2886/2009	CY061777
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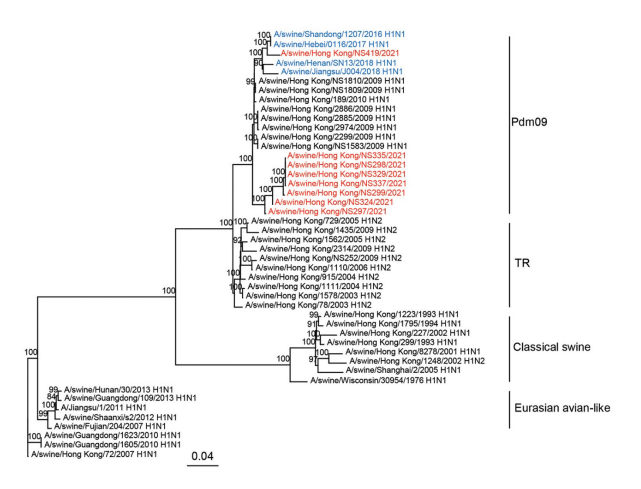
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NS	A/swine/Hong Kong/1248/2002	CY085530
NS	A/swine/Hong Kong/1578/2003	CY085610
NS	A/swine/Hong Kong/729/2005	CY085738
NS	A/swine/Hong Kong/NS252/2009	CY086002
NS	A/swine/Hubei/01/2009	CY091741
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NS	A/swine/Hubei/104/2009	CY091773
NS	A/swine/Guangdong/275/2010	JN375253
NS	A/swine/Guangdong/278/2010	JN375254
NS	A/swine/Guangdong/297/2010	JN375257
NS	A/swine/Guangdong/1437/2010	JN375267
NS	A/swine/Guangdong/1604/2010	JN375269
NS	A/swine/Guangdong/1611/2010	JN375271
NS	A/swine/Shaanxi/s2/2012	JX963611
NS	A/Jiangsu/1/2011	KF057140
NS	A/swine/Guangdong/109/2013	KP404216
NS	A/swine/Hunan/30/2013	KP404296
NS	A/swine/Hong Kong/299/1993	CY084989
NS	A/swine/Hong Kong/1223/1993	CY085061
NS	A/swine/Hong Kong/1795/1994	CY085109
NS	A/swine/Hong Kong/8278/2001	CY085378
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NS	A/swine/Hebei/0116/2017	MN418667
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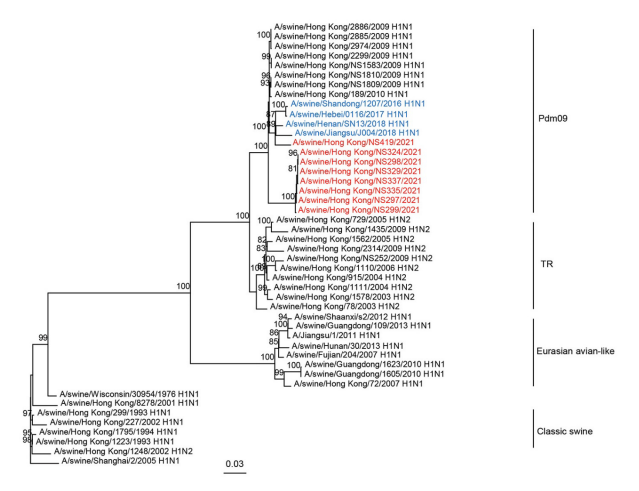
Appendix Figure 1. Phylogenetic tree of H1 (A) or H3 (B) gene sequences. Viral sequences generated from this study (red) and those downloaded from public domains (Appendix Table) were aligned using muscle v3.8 (http://www.drive5.com/muscle/). Virus sequences from G4 swine viruses (Main text) are highlighted in blue. Phylogenetic trees were then constructed by IQ-tree v1.6.12 (http://www.iqtree.org/release/v1.6.12) using the GTR+G model. Major animal viral lineages are as shown. The phylogenetic clades of the studied H1 sequences are shown in brackets (Anderson et al., mSphere 2016; https://doi.org/10.1128/mSphere.00275-16). Bootstrap values ≥80% are shown. Scale bar indicates estimated genetic distance.



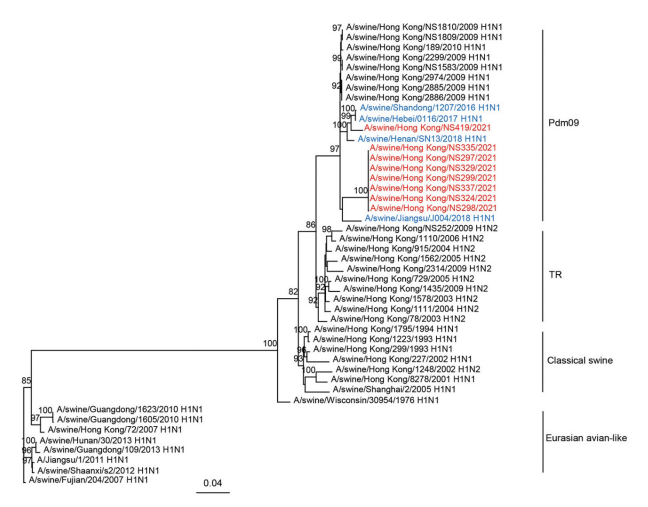
Appendix Figure 2. Phylogenetic tree of N1 (A) or N2 (B) gene sequences. Viral sequences generated from this study (red) and those downloaded from public domains (Appendix Table) were aligned using muscle v3.8 (http://www.drive5.com/muscle/). Virus sequences from G4 swine viruses (Main text) are highlighted in blue. Phylogenetic trees were then constructed by IQ-tree v1.6.12 (http://www.iqtree.org/release/v1.6.12) using the GTR+G model. Major animal viral lineages are as shown. Bootstrap values ≥80% are shown. Scale bar indicates estimated genetic distance.



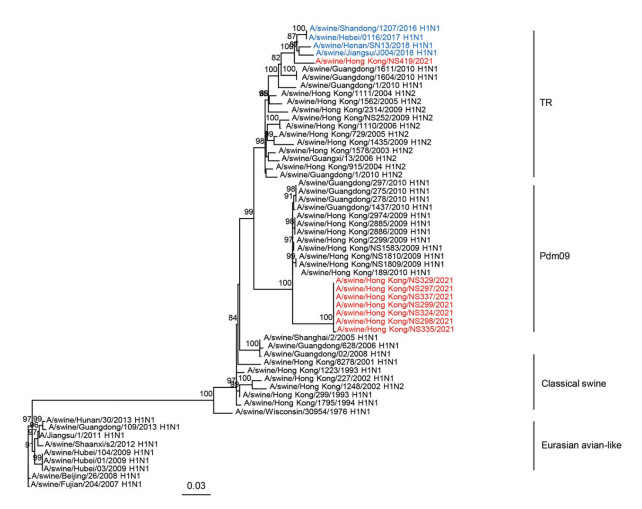
Appendix Figure 3. Phylogenetic tree of PB2 gene sequences. Viral sequences generated from this study (red) and those downloaded from public domains (Appendix Table) were aligned using muscle v3.8 (http://www.drive5.com/muscle/). Virus sequences from G4 swine viruses (Main text) are highlighted in blue. Phylogenetic trees were then constructed by IQ-tree v1.6.12 (http://www.iqtree.org/release/v1.6.12) using the GTR+G model. Major animal viral lineages are as shown. Bootstrap values ≥80% are shown. Scale bar indicates estimated genetic distance.



Appendix Figure 4. Phylogenetic tree of PA gene sequences. Viral sequences generated from this study (red) and those downloaded from public domains (Appendix Table) were aligned using muscle v3.8 (http://www.drive5.com/muscle/). Virus sequences from G4 swine viruses (Main text) are highlighted in blue. Phylogenetic trees were then constructed by IQ-tree v1.6.12 (http://www.iqtree.org/release/v1.6.12) using the GTR+G model. Major animal viral lineages are as shown. Bootstrap values ≥80% are shown. Scale bar indicates estimated genetic distance.



Appendix Figure 5. Phylogenetic tree of M gene sequences. Viral sequences generated from this study (red) and those downloaded from public domains (Appendix Table) were aligned using muscle v3.8 (http://www.drive5.com/muscle/). Virus sequences from G4 swine viruses (Main text) are highlighted in blue. Phylogenetic trees were then constructed by IQ-tree v1.6.12 (http://www.iqtree.org/release/v1.6.12) using the GTR+G model. Major animal viral lineages are as shown. Bootstrap values ≥80% are shown. Scale bar indicates estimated genetic distance.



Appendix Figure 6. Phylogenetic tree of NS gene sequences. Viral sequences generated from this study (red) and those downloaded from public domains (Appendix Table) were aligned using muscle v3.8 (http://www.drive5.com/muscle/). Virus sequences from G4 swine viruses (Main text) are highlighted in blue. Phylogenetic trees were then constructed by IQ-tree v1.6.12 (http://www.iqtree.org/release/v1.6.12) using the GTR+G model. Major animal viral lineages are as shown. Bootstrap values ≥80% are shown. Scale bar indicates estimated genetic distance.