

New Threats from H7N9 Influenza Virus: Spread and Evolution of High- and Low-Pathogenicity Variants with High Genomic Diversity in Wave Five

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ABSTRACT H7N9 virus has caused five infection waves since it emerged in 2013. The highest number of human cases was seen in wave 5; however, the underlying reasons have not been thoroughly elucidated. In this study, the geographical distribution, phylogeny, and genetic evolution of 240 H7N9 viruses in wave 5, including 35 new isolates from patients and poultry in nine provinces, were comprehensively analyzed together with strains from first four waves. Geographical distribution analysis indicated that the newly emerging highly pathogenic (HP) and low-pathogenicity (LP) H7N9 viruses were cocirculating, causing human and poultry infections across China. Genetic analysis indicated that dynamic reassortment of the internal genes among LP-H7N9/H9N2/H6Ny and HP-H7N9, as well as of the surface genes, between the Yangtze and Pearl River Delta lineages resulted in at least 36 genotypes, with three major genotypes (G1 [A/chicken/Jiangsu/SC537/2013-like], G3 [A/Chicken/Zhongshan/ ZS/2017-like], and G11 [A/Anhui/40094/2015-like]). The HP-H7N9 genotype likely evolved from G1 LP-H7N9 by the insertion of a KRTA motif at the cleavage site (CS) and then evolved into 15 genotypes with four different CS motifs, including PKG-KRTAR/G, PKGKRIAR/G, PKRKRAAR/G, and PKRKRTAR/G. Approximately 46% (28/61) of HP strains belonged to G3. Importantly, neuraminidase (NA) inhibitor (NAI) resistance (R292K in NA) and mammalian adaptation (e.g., E627K and A588V in PB2) mutations were found in a few non-human-derived HP-H7N9 strains. In summary, the enhanced prevalence and diverse genetic characteristics that occurred with mammalianadapted and NAI-resistant mutations may have contributed to increased numbers of human infections in wave 5.

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Address correspondence to George F. Gao, gaof@im.ac.cn, or Yuhai Bi, beeyh@im.ac.cn. Chuansong Quan, Weifeng Shi, and Yang Yang contributed equally to this article. **IMPORTANCE** The highest numbers of human H7N9 infections were observed during wave 5 from October 2016 to September 2017. Our results showed that HP-H7N9 and LP-H7N9 had spread virtually throughout China and underwent dynamic reassortment with different subtypes (H7N9/H9N2 and H6Ny) and lineages (Yangtze and Pearl River Delta lineages), resulting in totals of 36 and 3 major genotypes, respectively. Notably, the NAI drug-resistant (R292K in NA) and mammalian-adapted (e.g., E627K in PB2) mutations were found in HP-H7N9 not only from human isolates but also from poultry and environmental isolates, indicating increased risks for human infections. The broad dissemination of LP- and HP-H7N9 with high levels of genetic diversity and host adaptation and drug-resistant mutations likely accounted for the sharp increases in the number of human infections during wave 5. Therefore, more strategies are needed against the further spread and damage of H7N9 in the world.

KEYWORDS H7N9, HPAIV, wave five, avian influenza virus, dynamic reassortment, evolution, genetic diversity, origin

rior to 2013, the H7 subtype avian influenza viruses (AIVs) were mainly found in birds and were known to cause occasional human infections with mild illness (1). In March 2013, a novel H7N9 AIV emerged and caused severe disease in humans in China (2-7), with a case fatality rate of 39.1% (612 deaths/1,564 cases) as of 30 October 2017 (http://www.who.int/influenza/human_animal_interface/HAI_Risk_Assessment/ en/). Previous studies have shown that the novel H7N9 was low-pathogenicity (LP) avian influenza virus (LPAIV) and had an increased ability to bind to human-like sialic acid receptors (8–11). In addition, they also possessed several host adaptation mutations (e.g., T271A, K526R, E627K, or D701N) in the PB2 protein, which were associated with increased virus replication rates and disease severity in mammals and/or humans (12–14). So far, poultry-to-human transmission has been the primary route for human H7N9 infections (15, 16). The majority of human cases had histories of exposure to live-poultry or AIV-contaminated environments, such as live-poultry markets (LPMs) (http://www.who.int/influenza/human_animal_interface/HAI_Risk_Assessment/en/) (5, 15–17). Due to potential for further adaptation to mammals and possible human-tohuman transmission, the World Health Organization (WHO) previously classified H7N9 as "an unusually dangerous virus for humans" (http://www.sciencemag.org/news/2013/04/ h7n9-unusually-dangerous-virus-international-group-experts-concludes). Several asymptomatic and mild infections with the H7N9 LPAIVs, which may increase mammalian adaptation and transmissibility between humans, have also been reported in humans (18, 19).

There have been five infection waves since H7N9 emerged in March 2013. Notably, the fifth wave started earlier than the previous four, with a sudden and steep increase in the number of human cases (http://www.who.int/influenza/human _animal_interface/avian_influenza/riskassessment_AH7N9_201702/en/). More importantly, a high-pathogenicity (HP) avian influenza virus (HPAIV) H7N9 variant emerged during the fifth wave (wave 5) and has been identified as the causative pathogen in several human infections (20, 21). Furthermore, there have been reports of H7N9 infections in several family clusters, as well as in patients in the same hospital ward (http://www.who.int/influenza/human_animal_interface/avian_influenza/archive/en/). A comprehensive study of the geographical distribution and genetics of H7N9 viruses in wave 5 is needed to understand the underlying causes for the increased numbers of cases.

In this study, we sequenced 35 H7N9 AIVs collected from patients and poultry in LPMs or poultry farms in nine Chinese provinces during wave 5. Together with publicly available data, full-length genome sequences from a total of 240 H7N9 AIVs in wave 5 from 26 provinces and municipalities were systematically analyzed, and the results are presented here.



FIG 1 The temporal and spatial distributions of H7N9 human infections during wave 5. (A) The numbers of human H7N9 cases in each wave are listed by month. The fatality rates in January of wave 2 to 5 are shown. (B) Three provinces (QH, NX, and HaiN) without human or poultry H7N9 infections during wave 5 are colored in dark gray. The other provinces reporting human or poultry H7N9 infections are colored in red or faint yellow. Red indicates that HP-H7N9 was found in these provinces. The blue border indicates the new H7N9 viruses isolated from poultry and humans by our group. The numbers of HP- and LP-H7N9 strains analyzed in the present study are listed within the brackets as indicated in the provinces. Abbreviations in the map are as follows: AH, Anhui; BJ, Beijing; CQ, Chongqing; FJ, Fujian; GD, Guangdong; GS, Gansu; GX, Guangxi; GZ, Guizhou; HaiN, Hainan; HB, Hebei; HK, Hong Kong; HLJ, Heilongjiang; HN, Henan; HuB, Hubei; HuN, Hunan; IM, Inner Mongolia; JL, Jilin; JS, Jiangsu; JX, Jiangsu; JX, Jiangxi; LN, Liaoning; MC, Macao; NX, Ningxia; QH, Qinghai; SA, Shaanxi; SC, Sichuan; SD, Shandong; SH, Shanghai; SX, Shanxi; TB, Tibet; TJ, Tianjin; TW, Taiwan; XJ, Xinjiang; YN, Yunnan; ZJ, Zhejiang.

RESULTS

A sudden increase in the number of human H7N9 cases in wave 5. Human infections with H7N9 AIV emerge in waves during the winter months, with peaks around January, and there have been five waves to date (Fig. 1A). The fifth wave began

in October 2016, and the number of infections suddenly increased and peaked in January 2017 but remained at high levels through May 2017. Over 760 human cases were reported between October 2016 and September 2017, which is substantially higher than the rates in waves 1 to 4 (Fig. 1A).

Similarly to previous waves, almost all cases of human H7N9 infections corresponded to a history of exposure to poultry or AIV-contaminated environments during wave 5, with the cases starting at the centers of the Yangtze River Delta (Zhejiang and Jiangsu) and the Pearl River Delta (Guangdong). Over 71% of cases were reported after December 2016 were found in areas around those two centers (including Zhejiang, Jiangsu, Shanghai, Anhui, Guangdong, Hong Kong, Macao, Fujian, Hunan, Jiangxi, and Guangxi). Approximately 140 cases were reported in Guizhou, Hubei, Yunnan, Sichuan, Chongqing, Shandong, Henan, and Taiwan provinces or municipalities, where are geographically distant from the two centers. Over 80 human cases were reported in even more distant regions, such as Liaoning, Jilin, Inner Mongolia, Hebei, Tianjin, Beijing, Shanxi, Shaanxi, Gansu, Xinjiang, and Tibet provinces or municipalities, after February 2017. In contrast to previous waves (22), human infections with H7N9 during wave 5 were widespread throughout virtually all of China, although the reports from Heilongjiang Province merely noted an H7N9 outbreak in poultry. Only three provinces (Ningxia, Qinghai, and Hainan) did not report H7N9 infections during wave 5 (Fig. 1B).

Importantly, HP-H7N9 variants have emerged in poultry or environmental samples from LPMs or have caused outbreaks in poultry farms from several provinces. These include Guangdong, Guangxi, Fujian, Hunan, Hebei, Tianjin, Beijing, Henan, Shandong, Shaanxi, Inner Mongolia, and Heilongjiang, and human infections were reported in Guangdong, Taiwan, Guangxi, Hunan, Shaanxi, Hebei, Henan, Fujian, and Yunnan (Fig. 1B). LP-H7N9 was additionally found in Jiangsu, Zhejiang, Jiangxi, and Liaoning provinces in LPMs or poultry farms. The prevalence of HP-H7N9 and LP-H7N9 in poultry farms and LPMs from the south (Guangdong) to the north (Inner Mongolia) of China closely matched the geographical distribution of reported human cases (Fig. 1; see also Table S1 in the supplemental material).

LP- and HP-H7N9 are cocirculating in China during wave 5. A total of 35 whole-genome sequences of H7N9 viruses in wave 5 were obtained by our AIV surveillance network (23). These sequences originated from nine provinces, including Guangdong (n = 12, 9 from human cases and 3 from poultry in LPMs), Jiangsu (n = 4, from poultry in LPMs), Jiangxi (n = 4, from poultry in LPMs), Liaoning (n = 2, from poultry in poultry farms), Shandong (n = 3, from poultry in poultry farms), Yunnan (n =2, from human cases), Zhejiang (n = 3, from poultry in LPMs), Fujian (n = 4, 3 from human cases and 1 from a poultry farm), and Hebei (n = 1, from a poultry farm) (Fig. 1) (Table 1). Genome sequences from 827 additional H7N9 AIVs were also obtained from the Global Initiative on Sharing All Influenza Data (GISAID), including 205 viruses isolated from wave 5 and the remaining from waves 1 to 4. In total, our data set included 240 H7N9 strains from 26 provinces or municipalities in wave 5 (Fig. 1B; see also Table S1), with 200 strains from humans. By inspecting the hemagglutinin (HA) protein alignment, a total of 61 strains (Guangdong, n = 29; Guangxi, n = 17; Hunan, n = 4; Shaanxi, n = 4; Hebei, n = 2; Beijing, n = 1; Fujian, n = 1; Inner Mongolia, n = 11; Shandong, n = 1; Taiwan, n = 1) were found to contain an insertion of multiple basic amino acids at the cleavage site (CS) (Fig. 1B) (Table 1) and thus should be regarded as HP-H7N9 variants. The remaining H7N9 sequences were found to be LP. Despite the emergence and spread of HP-H7N9, LP-H7N9 has still been dominant in China during wave 5. Taking into account all reported regions with HP-H7N9 or outbreaks in poultry with human cases, there were a total of 14 provinces and municipalities (including 4 provinces and municipalities without HP-H7N9 strain information) from the north (Inner Mongolia and Heilongjiang) to the south (Guangdong) of China which has reported HP-H7N9 (Fig. 1), suggesting that LP- and HP-H7N9 were cocirculating and causing human infections across China.

TABLE 1 Molecular characteristics of the H7N9 viruses isolated from poultry and humans^a

				NA (N2							
	HA (H3 numbe	ering)		numberii	ng)	M2	PB2				
		Residue	Residues	Residue	Stalk	Residue	Residue	Residue	Residue	Residue	Residue
H7N9 virus	Cleavage site	186	224–228	292	deletion	31	526	588	591	627	701
A/GD/Th008/2017	PKRKRTARG	V	NGQSG	К	69–73	Ν	K	V	Q	E	D
A/SZ/Th001/2016	PKGRG	V	NGLSG	R	69–73	N	К	V	Q	К	D
A/SZ/Th002/2016	PKGRG	V	NGLSG	R	69–73	N	К	V	Q	К	D
A/SZ/Th003/2017	PKGRG	V	NGLSG	R	69–73	N	Κ	V	Q	E	D
A/SZ/Th004/2017	PKGRG	V	NGLSG	K	69–73	N	Κ	V	Q	E	D
A/GD/Th005/2017	PKGKRIARG	V	NGQSG	R	69–73	N	R	A	Q	К	D
A/SZ/Th006/2017	PKGRG	V	NGLSG	R	69–73	Ν	К	V	Q	К	D
A/SZ/Th007/2017	PKRKRTARG	V	NGQSG	K	69–73	Ν	К	V	Q	К	D
A/SZ/Th008/2017	PKRKRTARG	V	NGHSG	К	69–73	Ν	R	Α	Q	E	Ν
A/FJ/QZ-Th001/2017	PKGRG	V	NGLSG	R	69–73	Ν	К	V	Q	К	D
A/FJ/QZ-Th002/2017	PKGRG	V	NGLSG	R	69–73	Ν	К	V	Q	К	D
A/FJ/QZ-Th005/2017	PKGRG	V	NGLSG	R	69–73	Ν	К	V	Q	К	D
A/YN/YN001/2017	PKGRG	V	NGLSG	R	69–73	Ν	К	V	Q	К	D
A/YN/YN002/2017	PKGRG	V	NGLSG	R	69–73	Ν	К	V	Q	К	D
CK/GD/SZBJ0011-O/2017	PKRKRTARG	V	NGQSG	К	69–73	Ν	К	V	Q	E	D
EN/GD/SZBA-E1/2017	PKGKRTARG	V	NGQSG	R	69–73	Ν	Κ	V	Q	К	D
CK/JS/KS001-O/2016	PKGRG	V	NGLSG	R	69–73	Ν	Κ	V	Q	E	D
CK/JS/KS004-O/2016	PKGRG	V	NGLSG	R	69–73	Ν	Κ	V	Q	E	D
CK/JS/KS008-O/2016	PKGRG	V	NGLSG	R	69–73	Ν	Κ	V	Q	E	D
DK/JX/ShRSX002-O/2017	PKGRG	V	NGLSG	R	69–73	Ν	Κ	V	Q	E	D
EN/JX/ShRXZ038/2017	PKGRG	V	NGLSG	R	69–73	Ν	Κ	V	Q	E	D
CK/JX/ShRGF080-P/2017	PKGRG	V	NGLSG	R	69–73	Ν	К	V	Q	E	D
CK/JX/ShRGF219-O/2017	PKGRG	V	NGLSG	R	69–73	Ν	Κ	V	Q	E	D
CK/ZJ/HZ142/2017	PKGRG	V	NGLSG	R	69–73	Ν	Κ	V	Q	E	D
CK/ZJ/HZ293/2016	PKGRG	V	NGLSG	R	69–73	Ν	К	I	Q	E	D
CK/ZJ/HZ298/2016	PKGRG	V	NGLSG	R	69–73	Ν	Κ	V	Q	E	D
CK/GD/DGCPLB032/2016	PKGRG	V	NGQSG	R	69–73	Ν	Κ	V	Q	E	D
CK/LN/LN-083/2017	PKGRG	V	NGLSG	R	69–73	Ν	R	А	Q	E	D
CK/SD/QD2-758/2017	PKGRG	V	NGLSG	R	69–73	Ν	К	V	Q	E	D
CK/SD/QD4-739/2017	PKGRG	V	NGLSG	R	69–73	Ν	К	V	Q	E	D
CK/SD/DZ056/2017	PKRKRTARG	V	NGQSG	R	69–73	Ν	R	А	Q	E	D
CK/LN/SY059/2017	PKGRG	V	NGLSG	R	69–73	Ν	К	V	Q	E	D
CK/HB/XT001/2017	PKGRG	V	NGLSG	R	69–73	Ν	К	V	Q	E	D
CK/JS/DT001/2017	PKGRG	V	NGLSG	R	69–73	Ν	К	V	Q	Е	D
CK/FJ/NP001/2017	PKRKRTARG	V	NGQSG	R	69–73	Ν	R	А	Q	E	D

^aA total of 35 H7N9 viruses were isolated and sequenced in the present study. CK, chicken; DK, duck; EN, environment; FJ, Fujian; GD, Guangdong; HB, Hebei; JS, Jiangsu; JX, Jiangsu; JX, Jiangsu; JX, Jiangsu; SD, Shandong; SZ, Shenzhen; YN, Yunnan; ZJ, Zhenjiang.

Molecular characterizations of H7N9 AIVs in wave 5. Similarly to viruses in previous waves, H7N9 viruses in wave 5 possessed the amantadine resistance mutation (S31N) in M2. The neuraminidase inhibitor (NAI)-resistant R292K mutation was also found in four new human strains. Interestingly, CK/GD/SZBJ0011-O/2017 isolated from LPM also possessed K292. To our knowledge, this is the first report that the NAIresistant H7N9 strain was isolated from poultry (Table 1). All of the new human isolates possessed at least one of the four mammalian adaptation mutations, i.e., K526R, A588V, E627K, or D701N, in PB2. Several human strains (A/SZ/Th001/2016, A/SZ/Th002/2016, A/SZ/Th006/2017, A/SZ/Th007/2017, A/YN/YN001/2017, A/YN/YN002/2017, A/FJ/QZ-Th001/2017, A/FJ/QZ-Th002/2017, and A/FJ/QZ-Th005/2017) possessed both the A588V and E627K mutations. A/GD/Th005/2017 had both K526R and E627K mutations, while A/SZ/Th008/2017 had both the K526R and D701N mutations. Importantly, apart from CK/ZJ/HZ293/2016, which possesses I588, all of the avian-origin isolates possessed either of the mammalian adaptive residues V588 and R526 in PB2. Furthermore, the E627K mutation was found in EN/GD/SZBA-E1/2017, an environmental strain which also possessed the A588V mutation (Table 1).

Consistent with data from previous waves, V186 and L226 in HA were found in all of the LP-H7N9 AIVs from human samples during wave 5. In contrast, the majority of HP-H7N9 viruses from humans and poultry possessed Q226 in HA, except for five

		НА													NA											
		1	Cleavage site																							
Consensus-Wave 1		Ι	А	к	L	Ρ	Е	I	Ρ	к	G	-	-	-	-	R	G	L	Е	А	Ρ	н	Е	R	А	R
Consensus-Wave 2		I	A	к	L	P	Е	I	Ρ	к	G	-	-	-	-	R	G	L	Е	А	Р	н	Е	R	А	R
Consensus-Wave 3		I	А	к	L	P	Е	I	Р	к	G	-	-	-	-	R	G	L.	Е	А	Р	н	Е	R	А	R
Consensus-Wave 4		I	т	к	L	P	Е	I	P	к	G	-	-	-	-	R	G	L	Е	А	P	н	Е	R	А	R
Consensus-Wave 5 (LP)		T	т	к	Ē	P	E	T	P	к	G	-	-	-	_	R	G	Ē	F	Δ	P	н	F	R	Δ	R
CK/HY/16876/2016	(2)	T	P	F	0	P	F	v	P	ĸ	G	к	R	т	Δ	R	G	1	ĸ	т	S	R	ĸ	R	D	ĸ
EN/GD/C16283222/2016	(3)	Ť	P	E	0	P	E	v	P	ĸ	G	к	R	÷	A	R	G	Ē	ĸ	÷	P	R	ĸ	R	D	к
CK/HZ/HZ04/2016	(3)	т	P	к	Q	P	Е	v	P	к	R	к	R	т	А	R	G	L	к	т	s	R	к	R	D	к
A/GD/17SF003/2016	(9)	т	P	Е	Q	P	Е	v	P	к	R	к	R	т	А	R	G	L	к	т	s	R	к	к	D	к
CK/GD/SZBJ0011-O/2017	(3)	т	Р	Е	Q	Р	Е	v	Р	к	R	к	R	т	А	R	G	L	к	А	Р	н	Е	к	A	R
CK/HZ/HZX/2017	(1)	т	s	Е	Q	Р	Е	v	Р	к	G	к	R	т	А	R	G	L.	к	А	Р	R	к	R	D	к
CK/Zhongshan/ZS/2017	(25)	т	Р	Е	Q	Р	Е	v	Р	к	R	к	R	т	А	R	G	L	к	т	s	R	к	R	D	к
A/GD/Th005/2017	(1)	т	Р	Е	Q	Р	Е	v	Р	к	G	к	R	I	А	R	G	L	к	А	Р	R	к	R	D	к
A/GD/17SF037/2017	(1)	т	Ρ	к	Q	Ρ	Е	v	Р	к	R	к	R	А	А	R	G	L	к	т	s	R	к	R	D	к
A/GD/17SF039/2017	(2)	т	Р	Е	Q	Р	Е	v	Р	к	R	к	R	т	А	R	G	L	к	А	Р	н	Е	R	А	R
A/GD/17SF033/2017	(2)	т	Р	Е	Q	Ρ	Е	v	Р	к	R	к	R	т	А	R	G	L	Е	т	S	R	к	R	D	к
A/GX/18891/2017	(2)	т	Р	Е	L	Р	Е	v	Р	к	R	к	R	т	А	R	G	L	к	т	S	R	к	R	D	к
A/GD/17SF064/2017	(1)	т	Ρ	Е	Q	Р	Е	v	Ρ	к	R	к	R	т	А	R	G	L	к	А	Р	н	Е	×	А	R
A/GX/18902/2017	(1)	т	Р	Е	×	Р	Е	v	Ρ	к	R	к	R	т	А	R	G	L	Е	т	S	R	к	R	D	к
EN/GD/SZBA-E1/2017	(1)	т	Р	Е	Q	Ρ	Е	v	Р	к	G	к	R	т	А	R	G	L	к	А	Р	н	Е	R	А	R
A/SZ/Th008/2017	(1)	т	Ρ	Е	н	Ρ	Е	v	Ρ	к	R	к	R	т	Α	R	G	L	к	т	S	R	к	к	D	к
A/HuN/25351/2017	(1)	т	Ρ	Е	н	Ρ	Е	v	Ρ	к	R	к	R	т	Α	R	G	L	к	А	Р	н	Е	R	А	R
A/HuN/17875/2017	(1)	т	Р	Е	Q	Ρ	Е	G	Р	к	R	к	R	т	А	R	G	L	к	т	S	R	к	R	D	к
CK/GD/SD027/2017	(1)	Ι	S	к	L	Р	Е	I	Ρ	к	G	к	R	т	А	R	G	L	к	А	Ρ	н	к	R	D	к
H7 and N9 numbering			130	182	235	333	334	335	336	337	338	339	340	341	342	343	344	345	407	21	39	45	280	289	354	427
H3 and N2 numbering		48	122	173	226	324	325	326	327	328	328+1	328+2	328+3	328+4	328+5	329	330	331	393	21	39	46	283	292	358	430

FIG 2 Distinctive amino acid variations in HA and NA of HP-H7N9 compared to LP-H7N9. The amino acid sequences of HA and NA were aligned between HP-H7N9 and the consensus sequences of LP-H7N9. The specific residues in HA and NA are listed. "X" of A/GX/18902/2017 at site 235 indicates an H or L residue, and "X" of A/GD/17SF064/2017 at site 289 indicates a K or R residue. The numbers in brackets represent the numbers of HP-H7N9 strains. CK, chicken; DK, duck; EN, environment; GD, Guangdong; GX, Guangxi; HuN, Hunan; HY, Heyuan; HZ, Huizhou; SZ, Shenzhen.

human and one poultry strains (A/SZ/Th008/2017 and A/HuN/25351/2017 possessed H226, which had not been seen before in H7N9; two A/GX/18891/2017-like human strains and the poultry-origin CK/GD/SD027/2017 strain possessed L226; A/GX/18902/ 2017 possessed either residue H or residue L at position 226 of HA). There were four motif patterns at the CS in the HA protein of the HP H7N9 viruses, PKR<u>KRTA</u>R/G, PKG<u>KRTA</u>R/G, PKG<u>KRTA</u>R/G, and PKR<u>KRAA</u>R/G (Fig. 2). The majority of the HP strains possessed PKR<u>KRTA</u>R/G. Interestingly, the PKG<u>KRTA</u>R/G motif was just found in poultry-derived viruses and the last two motifs named above were possessed by only a single human strain each, A/GD/Th005/2017 and A/GD/17SF037/2017, respectively (Fig. 2) (Table 1). All of the H7N9 viruses from waves 1 to 5 possessed the same five-amino-acid-deletion (amino acids 69 to 73; N2 numbering) in the NA stalk.

Distinctive amino acid substitutions in the HA and NA of HPAIV H7N9. Apart from the four-amino-acid insertion, several specific amino acid substitutions between the HP- and LP-H7N9 AIVs were also identified (Fig. 2). For example, amino acid residues at positions 56, 130, 182, 235 (H3 numbering as 226), 335, and 407 in the HA protein of the HP strains were different from those of the LP strains (Fig. 2). In addition, amino acids at positions 21, 39, 45, 280, 354, and 427 were also specific to NA of HP-H7N9, although a few HP-H7N9 strains, e.g., CK/GD/SZBJ0011-O/2017-like and A/GD/17SF039/2017-like, possessed the same amino acids as LP-H7N9. Interestingly, A/GD/17SF064/2017 was associated with either a K or R residue mutation caused by a single nucleotide polymorphism (SNP) at site 289 (N2 numbering as 292) of NA (Fig. 2).

Origin and dynamic reassortment of H7N9 AIVs in wave 5. On the basis of the phylogenies, both the HA and NA of all the H7N9 strains identified in wave 5 clustered



FIG 3 Phylogenies of HA and NA genes of H7N9 viruses from different waves. All of the H7N9 viruses from waves 1 to 5, including our 35 new isolates, were used to perform the phylogenetic analysis. Panel A shows the HA gene tree, and the HP-H7N9 viruses in panel A were magnified as shown in panel B. Panel C shows the NA gene tree. The HP strains with different NA lineages are marked as stars with different colors in panels B and C; stars with the same color represent the same strain. H7N9 viruses from waves 1, 2, 3, 4, and 5 are colored blue, red, orange, light green, and lime green, respectively.

within the Yangtze River Delta lineage (n = 229) and the Pearl River Delta lineage (n = 229) 7), while four strains possessed a hybrid combination with HA from the Yangtze River Delta lineage and NA from the Pearl River Delta lineage (Fig. 3; see also Fig. S1 in the supplemental material). Although the 61 HP-H7N9 AIVs have four different amino acid motifs at the CS, all of the HP viruses were closely related, forming a single cluster in the Yangtze River Delta lineage in the HA tree (Fig. 3A and B; see also Fig. S1A). It should be noted that the Guangxi strains formed an independent clade (Fig. 3B), suggesting that there might have been more than one virus introduction event from Guangdong to Guangxi. However, the NA genes of the 61 HP strains were not clustered together in the phylogenetic tree (Fig. 3C; see also Fig. S1B). A total of 58 strains fell within the Yangtze River Delta lineage, among which 52 HP strains clustered together and shared the same origin. CK/FJ/NP001/2017, EN/GD/SZBA-E1/2017, A/GD/17SF039/2017, A/GX/ 18910/2017, A/HuN/25351/2017, and A/Taiwan/1/2017 did not fall within this cluster (Fig. 3C; see also Fig. S1B). Remarkably, the NA gene sequences of three HP strains, A/SZ/Th007/2017, A/GD/17SF064/2017, and CK/GD/SZBJ0011-O/2017, fell within the Pearl River Delta lineage (Fig. 3C; see also Fig. S1B). Therefore, the HA genes of the HP strains shared a single origin in the Yangtze River Delta lineage, whereas the NA gene had multiple different origins.

Phylogenetic analyses of the internal genes revealed high diversity compared to the HA and NA genes (Fig. 4; see also Fig. S1C to H) (Table 2; see also Table S1). After designating different H7N9 lineages for each internal gene, we classified the 240 isolates of wave 5 into 36 genotypes, 27 of which had emerged for the first time, suggesting continuing dynamic reassortment of H7N9 AIVs in wave 5 (Fig. 4A) (Table 2; see also Table S1). Nine genotypes, G1, G2, G3, G5, G9, G11, G17, G23, and G25,



FIG 4 Schematic representation of the evolutionary pathway of the H7N9 viruses during wave 5. (A) The schematic evolutionary pathway of H7N9 in wave 5 on the basis of the timeline of the first and last detection for each genotype during wave 5. Three dominant genotypes (G1, G3, and G11) were expressed as bigger "virions." The eight gene segments (shown as horizontal bars starting from top to bottom of the "virion") are PB2, PB1, PA, HA, NP, NA, M, and NS. Different colors of gene segments represent different virus lineages. (B) The dominant genotypes, the genotypes that included more than eight strains, and the transient genotypes are listed along with their isolation times. Symbols represent the corresponding genotypes of the H7N9 viruses.

emerged before wave 5 and were still circulating in the influenza season current at the time of writing. In particular, G1 (n = 38), known as CK/JS/SC537/2013 (JS537) in a previous study, emerged in wave 1 and continued to circulate during wave 5. Similarly, G11 (n = 98), represented by A/AH/40094/2015 (AH40094), emerged in wave 3 and still circulating in wave 5. Additionally, the strains in G3 (n = 28, represented by CK/ Zhongshan/ZS/2017) that were detected during wave 5 all belonged to HP-H7N9 and were widely spread in nine provinces or municipalities (Guangdong, Guangxi, Hunan, Fujian, Hebei, Beijing, Shandong, Shaanxi, and Inner Mongolia). These three genotypes caused the majority of human and poultry infections and seemed to be the major

									No. of	strains			
Genotype	HA	NA	PB2	PB1	PA	NP	MP	NS	LPAIV	LPAIV HPAIV First detection		Reference strain ^a	Accession no.
G1	Yangtze	Yangtze	1	1	1	1	1	1	29	9	16 April 2013	CK/JS/SC537/2013	EPI_ISL_142930
G2	Yangtze	Yangtze	1	1	1	1	1	2	0	1	1 March 2016	CK/GD/DG16419/2016	EPI_ISL_249139
G3	Yangtze	Yangtze	1	1	1	1	4	1	0	28	26 February 2015	A/GD-92/2015	EPI_ISL_198746
G4	Yangtze	Yangtze	1	1	1	9	1	1	0	1	15 January 2017	A/CK/GuZ/HD621/2017	EPI_ISL_248816
G5	Yangtze	Yangtze	1	1	4	1	1	1	2	0	28 October 2014	EN/JS/98337/2014	EPI_ISL_192340
G6	Yangtze	Yangtze	1	1	4	1	4	1	0	1	22 February 2017	A/GD/17SF033/2017	EPI_ISL_267759
G7	Yangtze	Yangtze	1	1	4	9	1	1	1	1	7 October 2016	A/ZJ/6/2016	EPI_ISL_242866
G8	Yangtze	Yangtze	1	1	4	9	3	1	1	0	3 March 2017	A/GX/6/2017	EPI_ISL_268525
G9	Yangtze	Yangtze	1	1	6	1	1	1	0	1	31 December 2014	A/JS/18828/2014	EPI_ISL_192278
G10	Yangtze	Yangtze	1	1	6	7	4	1	0	1	27 February 2017	A/GX/18892/2017	EPI_ISL_268502
G11	Yangtze	Yangtze	2	1	4	1	1	1	97	1	23 May 2015	A/AH/40094/2015	EPI_ISL_192471
G12	Yangtze	Yangtze	2	1	4	6	4	1	1	0	16 February 2017	CK/JX/ShRGF219-O/2017	EPI_ISL_259763
G13	Yangtze	Yangtze	2	1	4	9	1	1	9	0	26 October 2016	A/JS/60467/2016	EPI_ISL_242873
G14	Yangtze	Pearl	2	1	4	9	4	1	1	1	10 December 2016	CK/GD/DGCPLB032/2016	EPI_ISL_259747
G15	Pearl	Pearl	2	1	4	9	4	1	1	0	13 December 2016	A/GD/60061/2016	EPI_ISL_242888
G16	Pearl	Pearl	2	1	4	9	4	2	1	0	28 December 2016	A/HK/VB16189623/2016	EPI_ISL_240520
G17	Yangtze	Yangtze	2	1	6	1	1	1	0	8	15 January 2015	A/GD-20/2015	EPI_ISL_198726
G18	Yangtze	Yangtze	2	1	6	6	1	1	3	0	5 January 2017	A/HK/125/2017	EPI_ISL_259269
G19	Pearl	Pearl	2	1	6	7	1	1	1	0	9 December 2016	A/GD/60060/2016	EPI_ISL_242871
G20	Pearl	Pearl	2	1	6	7	4	1	3	0	28 December 2016	A/GD/60923/2016	EPI_ISL_242870
G21	Yangtze	Pearl	2	1	6	9	1	2	0	2	12 March 2017	A/GD/17SF064/2017	EPI_ISL_267764
G22	Yangtze	Yangtze	2	1	10	9	4	1	1	0	16 January 2017	A/SZ/Th003/2017	EPI_ISL_250313
G23	Yangtze	Yangtze	2	2	1	1	1	1	3	0	13 January 2016	A/FJ/1/2016	EPI_ISL_233628
G24	Yangtze	Yangtze	2	2	4	1	1	1	12	2	1 January 2017	A/FJ/02151/2017	EPI_ISL_242842
G25	Yangtze	Yangtze	2	2	4	9	1	1	1	3	28 February 2016	CK/GD/CZ9/2016	EPI_ISL_249282
G26	Yangtze	Yangtze	2	2	4	9	4	1	3	0	11 January 2017	A/HuN/02287/2017	EPI_ISL_242845
G27	Yangtze	Yangtze	2	6	4	9	4	1	1	0	30 December 2016	A/SZ/Th002/2016	EPI_ISL_250425
G28	Yangtze	Yangtze	2	6	6	6	1	1	1	0	19 December 2016	A/HK/VB16184091/2016	EPI_ISL_239994
G29	Pearl	Pearl	2	6	6	9	1	2	1	0	1 January 2017	CK/GD/JMS26/2017	b
G30	Yangtze	Yangtze	2	H6	H6	H6	1	H6	1	0	16 February 2017	DK/JX/ShRSX002-O/2017	EPI_ISL_259751
G31	Yangtze	Yangtze	2	2	5	9	1	1	1	0	4 February 2017	A/HuN/25360/2017	EPI_ISL_285569
G32	Yangtze	Yangtze	2	1	5	1	4	1	1	0	22 February 2017	A/GZ/18980/2017	EPI_ISL_285148
G33	Yangtze	Yangtze	1	1	5	9	1	2	1	0	22 March 2017	A/CQ-Yuzhong/1500/2017	EPI_ISL_285024
G34	Yangtze	Yangtze	1	6	4	9	1	1	1	0	17 June 2017	A/SC/29764/2017	EPI_ISL_285299
G35	Yangtze	Yangtze	1	1	4	9	6	1	1	0	25 June 2017	A/YN/32293/2017	EPI_ISL_285307
G36	Yangtze	Yangtze	2	1	6	9	1	1	0	1	11 April 2017	A/HuN/25351/2017	EPI_ISL_269517

^aAH, Anhui; CK, chicken; CQ, Chongqing; DK, duck; EN, environment; FJ, Fujian; GD, Guangdong; GuZ, Guangzhou; GX, Guangxi; GZ, Guizhou; HK, Hong Kong; HuN, Hunan; JS, Jiangsu; JX, Jiangxi; SC, Sichuan; SZ, Shenzhen; YN, Yunnan; ZJ, Zhenjiang.

^b—, not available.

genotypes in wave 5 (Fig. 4B). In detail, 229 isolates from wave 5 and the Yangtze River Delta HA/NA lineage belonged to 29 genotypes, seven strains of the Pearl River Delta HA/NA lineage clustered into five genotypes (G15, G16, G19, G20, and G29), and the four hybrid HA/NA strains belonged to two genotypes (G14 and G21) (Table 2; see also Table S1).

The 61 HP-H7N9 strains with full-length genomes belonged to 15 genotypes (Fig. 4) (Table 2; see also Table S1); among them, G1, G7, G11, G14, G24, and G25 were also seen in LP-H7N9 AIVs and G2, G3, G4, G6, G9, G10, G17, G21, and G36 were found only in the HP strains that emerged in wave 5. Therefore, the internal genes of HP-H7N9 viruses also underwent a dynamic reassortment process, although G1 (n = 9, CK/ Heyuan/16876/2016-like) and G3 (n = 28, CK/Zhongshan/ZS/2017-like) accounted for the ~61% of the HP strains.

DISCUSSION

H7N9 AIVs have caused the largest number of human cases in wave 5. Similarly to previous waves, human infections were first identified in the Yangtze River Delta and Pearl River Delta regions and then were gradually identified in other regions (22). However, in contrast with previous waves, over 100 H7N9 human cases were reported in the regions around the two centers as early as December 2016. Additionally, over 220 human cases were reported during wave 5 in regions far from

the two centers, some of which were important poultry breeding regions. Meanwhile, HP- and LP-H7N9 viruses were also identified in poultry from the south (Guangdong) to the north (Inner Mongolia and Heilongjiang) of China (Fig. 1; see also Table S1 in the supplemental material) (http://www.oie.int/en/animal-health -in-the-world/update-on-avian-influenza/2017/). These results indicated a much broader distribution of H7N9 viruses in China in wave 5, and the potential infection and transmission of both LP- and HP-H7N9 in poultry farms warrant further investigation.

The positivity rates for H7N9 were 15.7%, 6.5%, and 0.5% in LPMs from the Yangtze River Delta (Jiangsu, Shanghai and Zhejiang), the South Central region (Hubei, Hunan, Anhui and Fujian), and the Pearl River Delta (Guangdong), respectively, between November 2014 and April 2016 (24). In December 2016, the positivity rates for H7N9 were 15.8% and 9.4% in the LPMs of Jiangsu and Guangdong provinces, respectively (http://www.who.int/influenza/human_animal_interface/avian_influenza/riskassessment_AH7N9_201702/en/). These results indicated that the positivity rate of H7N9 in Jiangsu Province was consistent with previous observations from the start of wave 5, whereas a higher infection rate was seen in Guangdong Province than had been seen in previous waves (24) (http://www.who.int/influenza/human_animal_interface/avian_influenza/riskassessment_AH7N9_201702/en/). An increased positivity rate in provinces across China was probably one of the underlying reasons for the increased numbers of human infections in wave 5.

The emergence of HP-H7N9 undoubtedly poses a serious threat to both agriculture and public health. Although there was no evidence that HP-H7N9 was more virulent than LP-H7N9 for humans (20) or more transmissible as a consequence of enhanced infectivity of the respiratory epithelium in nasal passages (25), poultry infected by HP-H7N9 become sick and may shed high concentrations of viruses that would increase the risk of human infections. Human infections caused by HP-H7N9 AIVs have been identified in the Guangdong, Guangxi, Fujian, Hunan, Shaanxi, Hebei, Henan, and Yunnan provinces of mainland China and also in Taiwan (http://www.who.int/influenza/ human_animal_interface/HAI_Risk_Assessment/en/) (26), but several additional provinces, such as Heilongjiang, Inner Mongolia, Shandong, Beijing, and Tianjin, have also reported HP-H7N9 outbreaks in poultry or detection in LPMs (Fig. 1; see also Table S1) (http://www.oie.int/en/animal-health-in-the-world/update-on-avian-influenza/ 2017/). The majority of the circulating H7N9 strains in wave 5 were LP, suggesting that LP-H7N9 AIVs are still dominant in China. However, it seems that HP-H7N9 has spread into the north (Inner Mongolia and Heilongjiang) from the south (Guangdong) of China, and continued transmission and cocirculation with LP-H7N9 in China warrant further investigation.

HP-H7N9 AIVs contain the characteristic insertion of multiple basic amino acids at the CS of the HA protein (20). At least four different motifs were identified at the CS of the HA protein, suggesting various levels of potential virulence of the HP-H7N9 strain with respect to poultry and mammals (1, 27, 28). In addition, apart from the insertion of multiple basic amino acids, several amino acid substitutions in HA have also been found only in the novel HP-H7N9 AIVs. Interestingly, almost all of the HP-H7N9 viruses have Q226 (H3 numbering) in HA, indicating preferential binding to avian virus-type receptors (8, 29). Therefore, the HP-H7N9 variant likely first emerged in poultry and has not yet adapted to human as well as LP-H7N9. However, two human-origin HP-H7N9 isolates possessed H226, which had not been found before, and thus the receptor binding properties associated with this mutation need further characterization. Furthermore, two human-derived strains and one poultry-derived HP-H7N9 strain also possessed L226 (Fig. 2), indicating the ability to bind to human-type receptors (30). All these mutations suggested that HP-H7N9 was adapting to humans, and genetic mutations associated with pathogenicity need to be closely monitored.

The functional balance between the HA and NA proteins has been previously described (31–36). Along with the occurrence of specific mutations in the HA protein, there are several residues that have been identified specifically in the NA protein of the HP-H7N9 strains, but the potential biological functions of these mutations require

further studies. However, it should be noted that eight HP-H7N9 strains possessed an LP-H7N9-like NA, which is indicative of multiple origins.

The HA and NA genes of LP-H7N9 isolates shared the same origin and were relatively stable (37). Similarly to the HA gene of LP-H7N9 AIVs, the HP strains clustered together and shared a single origin Yangtze or Pearl River Delta lineage. However, the NA gene sequences of HP-H7N9 failed to cluster together. Alternatively, they fell within different parts of the Yangtze River Delta lineage, as well as within the Pearl River Delta lineage, suggesting that the NA gene might have also undergone a dynamic reassortment, which has not been seen in LP-H7N9.

The evolution of internal genes of LP-H7N9 was characteristic of a dynamic reassortment process (37–39). Our phylogenetic analyses showed that the LP-H7N9 AIVs were still undergoing dynamic reassortment in wave 5. The HP strains also underwent dynamic reassortment and recruited internal genes from the LPAIV reservoir, such as LP-H7N9/H9N2. On the basis of the previous classification method (37), the viruses in waves 1 to 5 were systematically analyzed. At least 36 genotypes were assembled via dynamic reassortment and were cocirculating in wave 5 (Fig. 4) (Table 2; see also Table S1). Three major genotypes (G1, G3, and G11) were identified. The G1 (JS537-like) viruses have been prevalent since wave 1 and may be ancestors for the HP variants emerging in wave 5 (Fig. 4), while the derived G3 genotype was dominant for HP strains. G11 (AH40094-like) viruses were also dominant and emerged as early as wave 3; however, only 1 of 98 G11 strains belonged to HP.

The R292K amino acid substitution in NA conferring NAI resistance was mainly found in clinics after treatments with NAIs (e.g., oseltamivir) (20, 40). Fourteen HP-H7N9 isolates in the present study possessed the R292K mutation. Notably, one was isolated from poultry (CK/GD/SZBJ0011-O/2017). Moreover, one NAI-resistant HP-H7N9 strain with an R292K mutation was reported that could transmit among ferrets (41). The E627K mutation in the PB2 protein is associated with increased replication efficiency and pathogenicity of AIV in mammalian hosts (12–14) and was seen in one environmental HP strain (EN/GD/SZBA-E1/2017). Furthermore, mammalian adaptation mutations A588V and K526R in PB2 were also found in the avian-origin isolates. These results suggest that there is increasing NAI resistance and adaptation of HP/LP-H7N9 to mammals.

In summary, we performed a systematic analysis of the genetic evolution of H7N9 AIVs in wave 5 compared to previous waves in China. Our results revealed a much broader distribution of H7N9 with high genetic diversity in China than previously known, which may account for the sharp increase in human infections with this virus. Although LP-H7N9 was still dominant in China, the rapid spread of and variations seen in HP-H7N9 are causes for concern. Intensive surveillance of H7N9 AIV in China is required to monitor the continuing dynamic reassortment of both LP- and HP-H7N9, as well as the spread of HP-H7N9 across China.

MATERIALS AND METHODS

H7N9 infection confirmation and whole-genome sequencing. Oropharyngeal or sputum samples were collected from suspected human cases at Shenzhen Third People's Hospital and Yunnan Center for Disease Control and Prevention. Environmental samples and oropharyngeal and cloacal swabs from apparently healthy or diseased poultry in LPMs and poultry farms located close to the suspected patients were also collected. Viral RNA was extracted using a Magabio plus virus RNA purification kit (automatic nucleic acid purification system NPA-32+; Bioer, China). Clinical samples were detected by quantitative reverse transcription-PCR (qRT-PCR) using avian influenza H7N9 detection kits (Mabsky Bio-tech Co, Ltd. and Zhengzhou Zhongdao Biotechnology Co., Ltd.), following manufacturer instructions. Whole-viral-genome amplification and Sanger sequencing were performed using specific primers (see Table S2 in the supplemental material).

Geographical distribution analysis. On the basis of data from the National Health and Family Planning Commission of the People's Republic of China (http://www.nhfpc.gov.cn/jkj/new_index.shtml), FluTrackers (https://flutrackers.com/forum/forum/china-h7n9-outbreak-tracking), and WHO (http://www .who.int/influenza/human_animal_interface/avian_influenza/archive/en/), human H7N9 cases in mainland China were analyzed by month since 2013. In addition to the H7N9-positive regions identified in the present study, all provinces and municipalities reporting human and poultry infections during wave 5 were systematically analyzed.

Genetic analysis. The gene sequences were edited using DNASTAR (Version 5.01) and BioEdit (Version 7.0.5.2). Protein sequences of 35 new H7N9 viruses were obtained using DNASTAR (Version 5.01) and confirmed by the use of the Annotation Program from GenBank (https://www.ncbi.nlm.nih.gov/genomes/FLU/annotation/). Nucleotide and amino acid sequences for all 240 H7N9 viruses in wave 5, including 205 whole genomes downloaded from GISAID (http://platform.gisaid.org), and for 35 new isolates were aligned using DNAMAN (Version 6.0). Biologically important amino acid residue mutations associated with pathogenicity, replication, transmissibility, host adaptation, and drug resistance (12–14, 42–51) were inspected.

Phylogenetic and genotyping analysis. Full-length H7N9 genome sequences from previous waves were also downloaded from GenBank and GISAID. For internal genes, the reference strains were retrieved via BLAST comparisons of the H7N9 gene sequences to those obtained from GISAID. For each gene, multiple-sequence alignment was performed using Muscle (52). Phylogenetic analysis was performed using RAxML (53). GTRGAMMA was applied as the nucleotide substitution model, and 1,000 bootstrap replicates were run. For phylogenetic trees of the internal genes, different lineages of H7N9 AIVs were classified if they had a common ancestor from no later than 2013. Generally, on the basis of the classification of different lineages of each internal gene and bootstrap values of >80%, the genotype of each H7N9 AIV from wave 2 to wave 5 was designated and genotypes from wave 5 were analyzed in detail.

Ethics statement. The study was performed in accordance with guidelines approved by the Ethics Committees from Shenzhen Third People's Hospital (SZTHEC2016001) and Yunnan Center for Disease Control and Prevention Ethics Committee (YNCDC2017001).

Accession number(s). Full-length genome sequences of the 35 H7N9 viruses from our study have been deposited in the GISAID EpiFlu database with accession numbers EPI_ISL_250311 to -17, EPI _ISL_250424 and -25, EPI_ISL_259747 to -63, EPI_ISL_266936, EPI_ISL_266938, and EPI_ISL_276781 to -87.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/JVI .00301-18.

SUPPLEMENTAL FILE 1, PDF file, 0.9 MB.

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We declare that we have no conflicts of interest.

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