Influenza A Virus Acquires Enhanced Pathogenicity and Transmissibility after Serial Passages in Swine

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ABSTRACT
Genetic and phylogenetic analyses suggest that the pandemic H1N1/2009 virus was derived from well-established swine influenza lineages; however, there is no convincing evidence that the pandemic virus was generated from a direct precursor in pigs. Furthermore, the evolutionary dynamics of influenza virus in pigs have not been well documented. Here, we subjected a recombinant virus (rH1N1) with the same constellation makeup as the pandemic H1N1/2009 virus to nine serial passages in pigs. The severity of infection sequentially increased with each passage. Deep sequencing of viral quasispecies from the ninth passage found five consensus amino acid mutations: PB1 A469T, PA 1129T, NA N329D, NS1 N205K, and NEP T48N. Mutations in the hemagglutinin (HA) protein, however, differed greatly between the upper and lower respiratory tracts. Three representative viral clones with the five consensus mutations were selected for functional evaluation. Relative to the parental virus, the three viral clones showed enhanced replication and polymerase activity in vitro and enhanced replication, pathogenicity, and transmissibility in pigs, guinea pigs, and ferrets in vivo. Specifically, two mutants of rH1N1 (PB1 A469T and a combination of NS1 N205K and NEP T48N) were identified as determinants of transmissibility in guinea pigs. Crucially, one mutant viral clone with the five consensus mutations, which also carried D187E, K211E, and S289N mutations in its HA, additionally was able to infect ferrets by airborne transmission as effectively as the pandemic virus. Our findings demonstrate that influenza virus can acquire viral characteristics that are similar to those of the pandemic virus after limited serial passages in pigs.

IMPORTANCE
We demonstrate here that an engineered reassortant swine influenza virus, with the same gene constellation pattern as the pandemic H1N1/2009 virus and subjected to only nine serial passages in pigs, acquired greatly enhanced virulence and transmissibility. In particular, one representative pathogenic passaged virus clone, which carried three mutations in the HA gene and five consensus mutations in PB1, PA, NA, NS1, and NEP genes, additionally was able to confer respiratory droplet transmission as effectively as the pandemic H1N1/2009 virus. Our findings suggest that pigs can readily induce adaptive mutational changes to a precursor pandemic-like virus to transform it into a highly virulent and infectious form akin to that of the pandemic H1N1/2009 virus, which underlines the potential direct role of pigs in promoting influenza A virus pathogenicity and transmissibility.

In March and early April 2009, a novel H1N1 influenza A virus (IAV) emerged in Mexico and the United States and rapidly triggered the first human pandemic of the 21st century (1). Phylogenetic and genetic studies revealed that the eight-gene segments of the H1N1/2009 virus were generated through reassortment between well-established swine influenza lineages, the Eurasian avian-like lineage, and the North American triple-reassortant lineage (1–3). Furthermore, structural and serological studies of its hemagglutinin (HA) have demonstrated that the H1N1/2009 virus is antigenically similar to 1918-like and classical swine H1N1 viruses (4, 5). Phylogenetically, the H1N1/2009 virus corresponds to a genetic ancestry of swine viruses, suggesting either an increased evolutionary rate or a long but unnoticed period of circulation in pigs prior to its 2009 pandemic emergence. Bayesian molecular clock analysis demonstrated that the evolutionary rate preceding the H1N1/2009 virus pandemic was typical for swine influenza (2). Thus, the reassortment of Eurasian avian-like and North American triple-reassortant swine lineages may not have occurred just before the 2009 pandemic; instead, a single reassortant (pandemic H1N1-like) virus may have been cryptically circulating in an unidentified host species for years before the outbreak in humans (2, 3). However, the reassortment dynamics of H1N1/2009 virus have not been determined in swine or humans by epidemiological surveillance (2, 6).

In our earlier study, we constructed a reassortant swine H1N1 influenza virus (rH1N1) with the same phylogenetic gene combination as the pandemic H1N1/2009 virus. The neuraminidase (NA) and matrix (M) gene segments were from
a Eurasian avian-like H1N1 swine influenza virus, and the other six genes were from a triple-reassortant H1N2 swine influenza virus (7). Unlike the pandemic H1N1/2009 virus, we found that this H1N1/2009-like virus is not able to confer virus transmissibility in guinea pigs, and that additional amino acid mutations are needed to make the virus as transmissible as the pandemic IAV (7). Consequently, the question remains: can IAVs acquire the characteristics of H1N1/2009 virus, including specific amino acid mutations of the H1N1/2009 virus, after undergoing adaptive changes in a specific host?

Thus far, adaptive changes of IAVs have been studied mainly by serial viral passages in laboratory species (8, 9). Mice have been used extensively for investigating pathogenic mechanisms and host range determinants (10, 11). Guinea pigs and ferrets support IAV transmission and have been used as models for IAV adaptation studies (12–16). Pigs, on the other hand, can be naturally or experimentally infected with IAVs. They can serve as mixing vessels or intermediate hosts for the generation of novel reassortant viruses (17, 18). Therefore, pigs most likely perform key roles in the evolutionary process of influenza viruses and in their cross-species transmission (19, 20). The emergence of the H1N1/2009 virus provides further evidence of the role of pigs in the influenza virus ecosystem. However, there is no direct supporting evidence to show that the pandemic H1N1/2009 virus was derived from pigs. Thus, it is important to investigate the adaptive evolution of IAVs in pigs.

To explore the evolutionary genesis of the pandemic H1N1/2009 virus, we performed serial passages of the rH1N1 construct in pigs and examined its sequential replication, pathogenicity, and transmissibility. By the ninth passage, the resulting viral population with five consensus mutations had acquired marked pathogenicity and in-contact transmissibility in swine and guinea pigs. Moreover, one particular representative viral clone, with three additional HA mutations, also acquired the ability for efficient airborne transmission between ferrets.

MATERIALS AND METHODS

Ethics statement. All animal research was approved by the Beijing Association for Science and Technology (approval identifier [ID] SYXK [Beijing] 2007-0023) and performed in compliance with the Beijing Laboratory Animal Welfare and Ethics guidelines, as issued by the Beijing Administration Committee of Laboratory Animals, and in accordance with the China Agricultural University (CAU) Institutional Animal Care and Use Committee guidelines (ID SKLAB-B-2010-003) approved by the Animal Welfare Committee of CAU.

Viruses and cells. The parental rH1N1 virus was described previously and was generated by reverse genetics (7), in which the NA and M genes from A/Swine/Fujian/204/2007 (accession numbers FJI536810 to FJI536817) and the other six genes from A/Swine/Guangdong/1222/2006 (accession numbers GU086078 to GU086085) were from the same phylogenetic cluster of H1N1/2009 viruses. The pandemic H1N1/2009 virus, A/Swine/Shandong/J73/2009 (SD731), was isolated from swine in Shandong Province, China, in December 2009. Its complete genomic sequence is available in GenBank (accession numbers JF951848 to JF951855). Viruses were titrated in MDCK cells to determine the median tissue culture infectious dose (TCID_{50}) by the Reed and Muench method (21). MDCK cells and A549 human lung adenocarcinoma cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM; Gibco) containing 10% fetal bovine serum (FBS; Gibco) and 1% antibiotics (Invitrogen). In vivo and in vitro experiments involving H1N1 and H1N1/2009 viruses were conducted in biosafety level (BSL) 2+ containment, as approved by the Ministry of Agriculture of China and the China National Accreditation Service for Conformity Assessment.

Serial passages, clinical measurements, and sampling. Landrace hybrid pigs, aged 4 to 5 weeks at the outset, were sourced from a high-health-status herd (porcine reproductive and respiratory syndrome [PRRS] virus free) and were IAV and antibody (H1, H3, H5, H7, and H9) negative by M gene PCR and hemagglutination inhibition assays prior to the start of the study. Passaging was established initially in one pig by intranasal inoculation with a total dose of 10^6 TCID_{50} rH1N1 virus, delivered in a final volume of 2.5 ml per nostril, using a mucosal atomization device (MAD; Wolfe Tory Medical, Inc.) to mimic aerogenic infection. The animal was euthanized at 4 days postinoculation (dpi). The lung was removed whole and lavaged with 200 ml phosphate-buffered saline (PBS) to obtain bronchioalveolar lavage fluid (BALF), and the nasal wash was performed with 10 ml PBS (containing 1% antibiotics). Five ml BALF and two ml nasal wash were combined to infect the next pig. The remaining nasal wash, BALF, and the collected tissue samples of each passage were stored at −80°C. During each passage, veterinary assessment was made at two fixed time points daily. Clinical parameters determined were rectal temperature, appetite, mental changes, bilateral nasal and ocular discharges, wheezing, and coughing.

Gross and microscopic examination of lung lesions. During necropsy, a mean value of the percentage of gross lesions (defined as dark red consolidation) was calculated for the pulmonary lobes. The lung tissue was collected and fixed in 10% phosphate-buffered formalin for histopathological examination, which was performed as described previously (22). Tissue sections of lungs were stained with hematoxylin and eosin (H&E) and examined microscopically for bronchiolar epithelial changes and peribronchial inflammation. Lesion severity was scored by the distribution or by the extent of lesions within the sections examined (22) with the following scale: 0, no visible changes; 1+, mild focal or multifocal change; 2+, moderate multifocal change; 3+, moderate diffuse change; 4+, severe diffuse change. Two independent pathologists scored all slides from blinded experimental groups.

Deep sequencing. Viral RNA was extracted from nasal wash and BALF of the pigs after nine serial passages with the parental rH1N1 virus using the High Pure RNA isolation kit (Roche). RNA was subjected to reverse transcription-PCR (RT-PCR) using 18 primer sets that cover the entire viral genome. These primer sets were designed according to the genome sequences of the H1N1 virus and by using Primer Premier 5.0 software. The fragments, approximately 600 to 800 nucleotides in length, were sequenced using the Illumina HiSeq2000 sequencing platform in the Chinese National Human Genome Center, Shanghai. Briefly, a library was constructed with TruSeq DNA sample prep kit set A. The DNA library was diluted and hybridized to the paired-end sequencing flow cells. DNA clusters were generated on a cBot cluster generation system with the TruSeq PE cluster generation kit v2, followed by sequencing on a HiSeq 2000 system with the TruSeq SBS kit v2. The threshold for the detection of single-nucleotide polymorphisms (SNP) was set manually at 10% of the population.

Sanger sequencing. Genome RNAs of viral clones were extracted from culture supernatants using the QIAamp viral RNA kit (Qiagen). Genes were reverse transcribed and amplified using a OneStep RT-PCR kit (Qiagen). Primers were the same as those used in the deep sequencing assay. The amplified cDNA products were excised from agarose gels and purified using a QIAquick gel extraction kit (Qiagen). Full-genome DNAs were Sanger sequenced by Huada Zhongsheng Scientific Corporation, and the sequence data were analyzed using GenScan software.

Viral growth kinetics. Confluent MDCK or A549 cells were infected with L2, N9, L18, or parental rH1N1 virus at a multiplicity of infection (MOI) of 0.001 in serum-free DMEM containing 2 mg/ml polybunylphenylalanyl chloromethyl ketone (TPCK)-trypsin (Sigma-Aldrich) and were cultured in a 37°C CO2 incubator. Cell supernatants were harvested every 12 h until 72 h postinoculation (hpi) and titrated by the TCID_{50} method on MDCK cells.
Western blotting. MDCK cells infected with L2, N9, L18, or parental rH1N1 virus were collected at 12 or 24 hpi. The protein samples derived from cell lysates were heated at 100°C for 5 min and then separated on a 10% sodium dodecyl sulfate-polyacrylamide gel and transferred to a polyvinylidene difluoride (PVDF) membrane (Bio-Rad). Membranes were incubated with mouse anti-NP monoclonal antibody (Abcam) and horseradish peroxidase (HRP)-conjugated rabbit anti-goat IgG (GE Healthcare, Inc.). The membranes were developed with an enhanced chemiluminescence kit (Pierce).

Polymerase activity assay. A combination of wild-type or mutant PB2, PB1, PA, and NP expression plasmids (125 ng each) was cotransfected into 293T cells with the luciferase reporter plasmid pHY-LucI (10 ng) and internal control plasmid renilla (5 ng). At 24 h posttransfection, a luciferase assay was performed using the Dual-Luciferase reporter assay system (Promega) and read using a GloMax 96 microplate luminometer (Promega).

Generation of mutant viruses by reverse genetics. RT-PCR-derived mutant viral genes were cloned into a dual-promoter plasmid, pHW2000 (Promega). The plasmids were introduced into 293T cells using Lipofectamine 2000 (Invitrogen) into each well of a 6-well plate and transfected with 0.5 μg of each of the eight plasmids and 10 μl Lipopectamine 2000 (Invitrogen) in a total volume of 1 ml of Opti-MEM (Invitrogen) into each well. After incubation at 37°C for 6 h, the transfection mixture was removed from the cells and 2 ml of Opti-MEM containing 1 mg/ml of TPC-trypsin was added. After 48 h, the supernatant was used to inoculate MDCK cells (cultured in a T75 flask) to produce stock virus. Viral RNA was extracted and analyzed by RT-PCR, and each viral segment was sequenced to confirm the identity of the virus. Stock viruses were titrated by the TCID_{50} method on MDCK cells.

Infection and transmission in pigs. Twelve Landrace hybrid pigs, aged 4 to 5 weeks, were randomly assigned into four separate groups of three. All pigs were sourced from a high-health-status herd (PRRS virus free and IAV and antibody negative). Each pig was infected intranasally with a total dose of 10^{6} TCID_{50} test virus delivered in a final volume of 2.5 ml per nostril. Three naive animals were introduced into each cage 24 h later. Beginning at 1 dpi, nasal swabs were collected daily and titrated on MDCK cells. The clinical symptoms of each pig were recorded daily. The directly infected pigs were euthanized at 6 dpi, and the contact pigs were euthanized at 7 days postcontact (dpc). Lungs were removed for viral load assessment and histopathology.

Contact transmission in guinea pigs. Female Hartley strain SPF/VAF guinea pigs (serologically negative for IAVs) that weighed between 300 and 350 g were obtained from Vital River Laboratories. They were anesthetized by intramuscular injection of Zoletil 100 (tiletamine-zolazepam; Virbac) prior to all handling procedures, including inoculation, nasal washes, and collection of blood. Three guinea pigs each were intranasally inoculated with a dose of 10^{6} TCID_{50} of a specific virus in 200 μl PBS. Each inoculated animal was placed in a new cage with one naive guinea pig at 1 dpi. Nasal washes were collected from all six guinea pigs at 2, 4, 6, 8, and 10 dpi. The ambient conditions for this study were set at 20 to 25°C and 30% to 40% relative humidity.

Respiratory droplet transmission in ferrets. Six- to twelve-month-old male Angora ferrets (Angora Ltd.), serologically negative by the hemagglutination inhibition assay for currently circulating influenza viruses (H1, H3, H5, H7, and H9), were used. At the start of the experiment, all ferrets were greater than 1.0 kg (range, 1.12 to 1.58 kg) in weight. Ferrets that lost more than 25% of their body weight or exhibited neurological dysfunction were euthanatized and submitted to postmortem examination. Baseline rectal temperature and weight measurements were obtained prior to infection. Groups of three ferrets were intranasally inoculated with 10^{6} TCID_{50} of test virus and housed in specially designed cages inside an isolator. At 1 dpi, three naive animals were placed in an adjacent cage (5 cm away), separated by a double-layered net divider that allowed free passage of air. Nasal washes were collected at 2-day intervals beginning at 2 dpi (1 day postexposure [dpe]) and were titrated in MDCK cells. Directly infected and exposed ferrets were euthanized at 8 dpi and 9 dpe, respectively. The ambient conditions for these studies were set at 20 to 25°C and 30% to 40% relative humidity. The horizontal airflow in the isolator was at 0.1 m/s and was directed from the inoculated to exposed animals.

Statistical analyses. Statistically significant differences between experimental groups were determined using the analysis of variance (ANOVA) method. A P value of <0.05 was considered statistically significant.

RESULTS

rH1N1 virus showed enhanced pathogenicity after nine passages in pigs. To study the natural evolution of a virus that has the same gene combination as the pandemic H1N1/2009 virus, we passaged the rH1N1 virus (7) in pigs by one-on-one delivery. Passage began with the nasal inoculation of 10^{6} TCID_{50} of the rH1N1 virus. Clinical signs were monitored daily until 4 dpi, at which time the animal was euthanized to collect nasal wash and BALF. The mixture of the nasal wash (2 ml) and BALF (5 ml) subsequently was used to inoculate the next pig intranasally (passage 2 [P2]). This procedure was repeated until P9.

Overall, the clinical signs elicited from P1 to P9 virus-infected pigs increased steadily from mild to severe. The P1 and P2 virus-infected pigs did not show any overt clinical signs from 1 to 4 dpi, although a small transient elevation in body temperature was observed at 1 dpi (Fig. 1). The P3 to P6 virus-infected pigs developed pyrexia (>40.0°C) only at 1 dpi (Fig. 1). Strikingly, the P7 to P9 virus-infected pigs showed sustained pyrexia (Fig. 1) accompanied by clear symptoms of depression, anorexia, tremors, and nasal and ocular discharge. The P8 virus-infected pigs displayed severe and earlier onset of clinical signs with noticeable wheezing and coughing at 4 dpi. The P9 virus-infected pigs were the most severely affected, with the earliest onset of clinical symptoms. Three control pigs inoculated with SD731 virus (a virulent H1N1/2009 virus) at 10^{6} TCID_{50} developed clinical signs (pyrexia, wheezing, and coughing) from 2 to 4 dpi that were similar to those observed in the P9 virus-infected pigs.

To examine the pathological effects of multiply passaged rH1N1 viruses, lungs from infected pigs were examined postmortem. Gross lesions of P1 to P9 virus-infected lungs showed increasing severity (Fig. 2). P1 and P2 virus-infected lungs appeared normal, but P3 to P8 virus-infected lungs showed increasing multifocal areas of consolidation in the cardiac, diaphragmatic, and intermediate lobes. P6 virus-infected lung displayed extensive hyperemia, edema, and diffuse consolidation over the entire lung (Fig. 2). P9 virus-infected lung had the most severe lesions with...
Passaging of rH1N1 virus in pigs led to increasing pulmonary damage. Gross and histopathological examinations were performed on the respiratory tracts of infected pigs at 4 dpi. Representative gross (left) and corresponding histological (H&E staining) findings at low (middle) and high (right) magnification are presented. (A and G) P1 virus-infected lung showed mild bronchopneumonia with minor lymphocytic infiltration. (B and H) P3 virus-infected lung had moderate bronchopneumonia with inflammatory cell infiltrates in the alveoli and interstitium (boxed area magnified). (C and I) P6 virus-infected lung showed severe bronchopneumonia with hemorrhage, edema, and diffuse consolidation. (D and J) P9 virus-infected lung displayed extensive consolidation in all lobes, which resembled the lung pathology found in SD731 virus-infected pigs (Fig. 2). Microscopic lesions of the cardiac lobes (black arrow in H). (E and K) Virulent SD731 virus-infected lung showed severe bronchopneumonia similar to the changes found in the P9 lung. (F and L) Mock-infected lung showed normal morphology. Blue arrows (left) indicate pulmonary tissue consolidation. Results shown are typical of three independent examinations for each animal. Scale bar, 200 μm.

FIG 2 Passaging of rH1N1 virus in pigs led to increasing pulmonary damage. Gross and histopathological examinations were performed on the respiratory tracts of infected pigs at 4 dpi. Representative gross (left) and corresponding histological (H&E staining) findings at low (middle) and high (right) magnification are presented. (A and G) P1 virus-infected lung showed mild bronchopneumonia with minor lymphocytic infiltration. (B and H) P3 virus-infected lung had moderate bronchopneumonia with inflammatory cell infiltrates in the alveoli and interstitium (boxed area magnified). (C and I) P6 virus-infected lung showed severe bronchopneumonia with hemorrhage, edema, and diffuse consolidation. (D and J) P9 virus-infected lung displayed extensive consolidation in all lobes, which resembled the lung pathology found in SD731 virus-infected pigs (Fig. 2). Microscopic lesions of the cardiac lobes (black arrow in H). (E and K) Virulent SD731 virus-infected lung showed severe bronchopneumonia similar to the changes found in the P9 lung. (F and L) Mock-infected lung showed normal morphology. Blue arrows (left) indicate pulmonary tissue consolidation. Results shown are typical of three independent examinations for each animal. Scale bar, 200 μm.

FIG 3 Histopathological lung lesion scores from progressively passaged rH1N1 virus-infected pigs at 4 dpi. Microscopic lesions of the cardiac lobes (black bars) and diaphragmatic lobes (white bars) were evaluated and assigned a score of 0 to 4 based on evaluation criteria described in Materials and Methods. All values shown represent the mean score from three independent evaluations of each lung. A representative score from SD731 virus-infected lungs served as a positive control.

Selection of three representative viral clones from P9 quasispecies. The passaging of IAVs in animals would result in the natural selection of heterogeneous mixtures of viruses with various mutations, the so-called viral quasispecies (23, 24). To assess possible genetic diversity of viral quasispecies in the upper and lower respiratory tract of the P9 virus-infected pigs, we performed deep sequencing on viral RNA derived from nasal wash and BALF. Five consensus mutations were found in the virion populations of both the nasal wash and BALF compartments (frequencies of >91%): PB1 A469T, PA I129T, NA N329D, NS1 N205K, and NEP T48N (Table 1). Mutations detected in HA showed apparent divergence between the nasal wash and BALF; D187E, K211E, and S289N mutations were frequent in the BALF, whereas M227T, S271P, and I295V mutations were frequent in the nasal wash (Table 1). This HA finding indicates that the genetic composition of viral quasispecies in the upper and lower respiratory tract of the P9 virus-infected pigs was distinctly different. Moreover, within each compartment, the mutational frequencies of the HA protein ranged from 26.46% to 55.89% in BALF and 54.72% to 66.35% in nasal wash (Table 1), indicating relatively low mutational consensus in viral populations within each location of the airway.

To select representative P9 viral clones from the viral quasispecies, we performed plaque purification and genome sequencing to establish the sequence composition of individual clones. We isolated randomly 20 lung viral clones (L1 to L20) and 20 nasal viral clones (N1 to N20) originating from the BALF and nasal wash, respectively, of the P9 virus-infected pigs. Four of the five consensus mutations (PB1 A469T, NA N329D, NS1 N205K, and NEP T48N) were identified in all 40 clones, and PA I129T was identified in 36/40 clones. In contrast, the HA protein showed much mutational variability (see Table S1 in the supplemental material). Among all HA mutations in the 20 lung viral clones, D187E, K211E, and S289N mutations accounted for twelve, five, and seven mutations, respectively. Most P9 viruses with the D187E mutation also had the K211E and/or S289N mutation. Con-
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October 2014 Volume 88 Number 20 jvi.asm.org

versely, in the 20 nasal viral clones, mutations M227I, S271P, and I295V in the HA protein were most frequent, occurring in combinations of two (5 clones) or three (10 clones) mutations (see Table S1).

We next selected three representative viral clones harboring all five of the consensus mutations (PB1 A469T, PA 1129T, NA N329D, NS1 N205K, and NEP T48N) in combination with the D187E, K211E, and S289N mutations (as a representative lower respiratory tract virus, named L2), with the M227I, S271P, and I295V mutations (as a representative upper respiratory tract virus, named N9), or with no mutational change in the HA (named L18).

Representative P9 virus-derived viral clones exhibited enhanced replication in vitro. To evaluate the replicative abilities of the L2, N9, and L18 viral clones, they were used to infect MDCK cells and human A549 cells at an MOI of 0.001 to determine virus titers over 72 hpi. For MDCK cells, the titers of the L2, N9, and L18 viruses were significantly higher than those of the parental rH1N1 virus at almost all time points (P < 0.05) (Fig. 5C). These results clearly demonstrated that the replication of the three P9 viruses in vitro was significantly greater than that of the parental rH1N1 virus.

PB1 A469T mutation conferred enhanced polymerase activity to parental rH1N1 RNP complex. Since there is close correlation between the activity of the viral ribonucleoprotein (RNP) complex and viral adaptation, replication, and pathogenicity (10, 25), the functional effect of PB1 A469T and PA 1129T consensus mutations on viral polymerase activity was assessed. The polymerase activity of the parental rH1N1 RNP complex was determined by luciferase assay through cotransfection into 293T cells of expression plasmids encoding PB2, PB1, PA, and NP, along with

![FIG 4](https://jvi.asm.org/content/jvi/88/20/11985/F04.large.jpg)

**FIG 4** Viral titers in nasal washes, BALFs, and lungs of progressively passaged rH1N1 virus-infected pigs at 4 dpi. Viral titers of nasal washes (A) and BALF (B) were directly determined by TCID<sub>50</sub> assays on MDCK cells. (C) Collected lung tissues were weighed, and 10% (vol/vol) homogenates were prepared in cold PBS. Virus titers in cleared homogenates were determined by TCID<sub>50</sub> assays on MDCK cells. All values shown are means ± standard deviations (SD) from three independent experiments for each sample. Representative SD731 virus titers derived from corresponding sites of infected pigs served as positive controls. An asterisk indicates that the virus titer was not significantly different from that of the SD731 virus group (P > 0.05 by ANOVA). The virus detection limit was 10 TCID<sub>50</sub>.

**TABLE 1** Mutation analysis of viral quasispecies in BALF and nasal wash of P9 virus-infected pigs

<table>
<thead>
<tr>
<th>Protein</th>
<th>Nucleotide</th>
<th>Frequency* (%) in:</th>
<th>Position</th>
<th>Amino acid</th>
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<tr>
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<td>Nasal wash</td>
<td>Nucleotide</td>
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<td>G</td>
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</table>

<sup>a</sup> All mutations shown have a frequency of >10%.

<sup>b</sup> ND, not detected.

<sup>c</sup> –, silent substitution.
luciferase reporter pYH-Luci plasmid and an internal renilla control plasmid (Fig. 6). The replacement of parental PA with PA I129T plasmid had little effect on RNP activity. However, replacement of parental PB1 with PB1 A469T plasmid resulted in about a 6-fold increase in polymerase activity \( (P < 0.05) \) (Fig. 6). This result indicates that PB1 A469T is an important contributor to the enhanced replicative ability of P9 viruses.

**Representative P9-derived viruses exhibited enhanced pathogenicity and transmissibility in swine.** To assess the pathogenicity of the three representative P9 virus-derived viral clones (L2, N9, and L18 viruses) in relation to parental rH1N1 virus in vivo, each group of three pigs was infected intranasally with each virus at 10^6 TCID\(_{50}\). At 1 dpi, three naive in-contact pigs were added to each infected group. All pigs directly infected with L2, N9, and L18 viruses showed typical flu-like symptoms (i.e., pyrexia, depression, nasal discharge, wheezing, and coughing) associated with moderate to severe lung lesions, and most in-contact pigs in these three groups also displayed mild pyrexia, nasal discharge, and similar pathological damage (Table 2 and Fig. 7). In contrast, the rH1N1-inoculated and in-contact pigs showed no obvious clinical symptoms and pulmonary damage (Table 2 and Fig. 7). Pathological lung scores were 2.8 ± 0.1, 2.5 ± 0.1, and 2.3 ± 0.3 in the L2, N9, and L18 directly infected groups, respectively, which were significantly higher than the scores of the rH1N1 inoculated pigs (0.3 ± 0.1; \( P < 0.05 \)).

Virus shedding or presence was monitored from nasal swabs of directly infected pigs from 1 to 6 dpi and of in-contact pigs from 1 to 7 dpc. Viral shedding was detected in all pigs directly infected with L2, N9, L18, and rH1N1 viruses at 1 to 6 dpi. Peak titers of the three P9 viruses were approximately 10-fold higher than those of the rH1N1 virus (Fig. 8). Notably, all L2, N9, and L18 in-contact pigs shed virus, but no virus was detected in rH1N1 in-contact pigs throughout the screening period (Fig. 8). Together, these data indicate that the pathogenicity and transmissibility of P9 virus-

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**FIG 5** Three representative P9 viral clones exhibited enhanced replication in MDCK and A549 cells. Viable virus output from MDCK (A) or A549 (C) cells infected with three viruses derived from the P9 infection (L2, N9, and L18) and parental rH1N1 viruses, at an MOI of 0.001, was determined by TCID\(_{50}\) assay at 12, 24, 36, 48, 60, and 72 hpi. The values are expressed as means ± SD (\( n = 3 \)). An asterisk indicates that the value of the corresponding virus was significantly different from that of the rH1N1 virus \( (P < 0.05) \) by ANOVA. (B) Western blots for viral NP detection in MDCK cells at 12 and 24 hpi; β-actin was used as a loading control. The Western blot results are representative of three independent experiments.

**FIG 6** Activity of parental rH1N1 RNP complex reconstituted with P9 virus-derived PA I129T and PB1 A469T mutants. A combination of four plasmids expressing RNP components of parental rH1N1 virus (PB2, PB1, PA, and NP), luciferase reporter plasmid pYH-Luci, and an internal renilla control plasmid were transfected into 293T cells. Mutant PA I129T or PB1 A469T plasmid was substituted for the corresponding wild-type rH1N1 plasmids as indicated. Values shown represent means ± SD for three independent experiments, normalized to the activity of parental rH1N1 RNP complex (100%). *, \( P < 0.05 \) compared to the correspondingly transfected rH1N1 response.
Virus titers are expressed as mean log10TCID50/g ± SD.

Microscopic lung lesions were assigned a score of 0 to 4. Scores are expressed as means ± SD.

Rectal temperatures higher than 39.5°C are considered pyrexia in pigs. Peak rectal temperatures are expressed in parentheses as means ± SD.

TABLE 2 Clinical symptoms, pathological changes, and virus detection in pigs

| Virus Group | Aquarium Experiment | Directly Infected | In-Contact | Pulmonary consolidation area (%) |
|-------------|---------------------|-------------------|------------|
| L2          | 1                   | 2                 | 3          |
| N9          | 1                   | 2                 | 3          |
| L18         | 1                   | 2                 | 3          |
| G1          | 1                   | 2                 | 3          |
| G2          | 1                   | 2                 | 3          |
| G3          | 1                   | 2                 | 3          |
| G4          | 1                   | 2                 | 3          |
| G5          | 1                   | 2                 | 3          |

Representative P9 virus-derived viruses also acquired in-contact transmissibility in guinea pigs. Our previously demonstrated that the parental rH1N1 virus is not transmissible between guinea pigs. However, two of three guinea pigs infected with L2, N9, and L18 viruses developed virus shedding from in-contact guinea pigs (Fig. 9A to C), which suggests that these viruses have largely acquired the ability to transmit infection to in-contact guinea pigs.

NS1 N205K and NEP T48N in rH1N1 virus conferred in-contact transmissibility in guinea pigs. Among the three consensus amino acid mutations, the 99 virus-derived clones, L18 virus contains the NS1 N205K and NEP T48N mutation, which is a five consensus amino acid mutation contributes to the contact transmissibility of P9 virus-derived clones. L18 virus was derived from a one-site mutant viruses from rH1N1, named PB1 A469T, PA I129T, and single-site combined mutations NA N329D, and NS1 N205K-NEP T48N (derived from a one-site mutant viruses from rH1N1, named PB1 A469T-PA I129T and NA N329D). Among the three P9 virus-derived clones, L18 virus was the most transmissible in guinea pigs. Consistent with previous findings (7), there was no apparent pathological damage detected in lungs of rH1N1-infected pigs. There was no apparent pathological damage detected in lungs of H1N1-infected pigs, and the L18 virus showed moderate pulmonary consolidation (white arrows) and moderate to severe bronchopneumonia with interstitial pulmonary edema (Fig. 9D). However, two of three guinea pigs in contact with L2, N9, and L18 viruses also showed mild lung consolidation (white arrows) and moderate to severe bronchopneumonia with interstitial pulmonary edema and inflammatory cell infiltrates in alveolar space (Fig. 9D). There was no apparent pathological damage detected in lungs of L2, N9, and L18 virus-infected pigs. There was no apparent pathological damage detected in lungs of H1N1-infected pigs.
Among these six viruses, the PB1 A469T-only mutation infected one of three contact guinea pigs (Fig. 10A and E), and NS1 N205K and NEP T48N mutation infected two of three contact animals (Fig. 10D and F). However, no virus shedding was detected in any contact animals from the PA I129T and NA N329D groups (Fig. 10B and C). These results indicated that PB1 A469T, combined NS1 N205K and NEP T48N single-site mutations in the parental rH1N1 virus, conferred enhanced in-contact transmissibility in guinea pigs. Additionally, virus output from these transmissible rH1N1 mutants at 5.75 to 7.25 log10 TCID50/ml (Fig. 9B to D and 10A and D to F) was significantly higher than that of the nontransmissible rH1N1 mutants at 4.25 to 5.75 log10 TCID50/ml (Fig. 9A and 10B and C) (P < 0.05), which suggested that the transmissibility of mutant PB1 A469T and combined mutant NS1 N205K and NEP T48N in guinea pigs also was related to enhanced virus replication. No obvious clinical signs were observed in any of the infected and uninfected guinea pigs during the observation period, and all animals with viral shedding also showed seroconversion (data not shown).

Severe pathogenicity and respiratory droplet transmission of representative P9-derived viruses in ferrets. Ferrets have been widely used as an experimental model to study human infection and transmission of influenza virus (13, 26). Three ferrets were infected intranasally with each virus (L2, N9, L18, and parental rH1N1 viruses) at a dose of 106 TCID50. Directly infected ferrets were individually caged and sited in close proximity to individually kept naïve ferrets (5 cm away).

All ferrets directly infected with P9 virus-derived L2, N9, and L18 viruses developed severe clinical symptoms (pyrexia, lethargy, anorexia, sneezing, wheezing, and coughing) from 2 to 8 dpi, including one death from L2 virus infection at 7 dpi (Table 3). However, all rH1N1 virus-infected ferrets developed only mild clinical signs of slight pyrexia, anorexia, and sneezing at 2 to 4 dpi (Table 3). At postmortem at 8 dpi, gross and histopathological lesions, including severe consolidation (>40%), hemorrhage, and edema, were evident in all ferrets infected with the three P9-derived viruses. In contrast, little or no lung consolidation (0% to 5%) was found in parental rH1N1 virus-infected ferrets (Fig. 11 and Table 3). These results demonstrate that the three P9-derived viruses (L2, N9, and L18) were much more pathogenic than the parental rH1N1 virus in ferrets.

To assess the transmissibility of the P9-derived viruses in ferrets, we compared virus titers in nasal washes from directly infected and proximally exposed ferrets. High levels of virus shedding were evident for all directly infected ferrets (≥6.25 log10 TCID50/ml at 2 to 4 dpi), which was about 10- to 100-fold greater than that of the corresponding rH1N1 virus group (Fig. 12). However, for proximally exposed ferrets, only the group associated with the L2 virus was found to shed virus continuously in all three exposed ferrets and at levels comparable to that of the directly infected ferrets (Fig. 12A). Notably, the high levels of L2 virus replication in directly infected and proximally exposed ferrets were highly similar to corresponding reported findings of H1N1/2009 viruses (27, 28). Additionally, all proximally exposed L2 ferrets showed severe clinical signs and pulmonary damage (Fig. 11 and Table 3). Taken together, these results demonstrate that the three representative P9-derived viruses (L2, N9, and L18) had acquired enhanced replication and pathogenicity in ferrets, and that L2 virus additionally was highly effective in airborne transmission in ferrets. The latter finding suggests that the HA mutations...
(D187E, K211E, and S289N) in the L12 virus contribute to its airborne transmission ability.

**DISCUSSION**

The pandemic H1N1/2009 virus is thought to have been generated in swine and circulated in pig populations before its cross-species transmission to humans; however, despite considerable research, this has not been proven (2, 3, 6). Here, we serially passaged an engineered H1N1/2009-like precursor (rH1N1) virus in pigs to simulate a probable evolutionary emergence route of the H1N1/2009 virus. Dramatic changes in viral characteristics occurred after just nine serial passages, including enhanced replication and polymerase activity in vitro, increased pathogenicity in pigs and ferrets, and efficient in-contact transmission in pigs and guinea pigs. Furthermore, a P9-derived clone (L2 virus) even acquired highly efficient respiratory droplet transmissibility in ferrets, a property which was similar to that of the H1N1/2009 virus.

Adaptation is believed to be a driving force in evolution, whereby organisms, including viruses, are selected in nature because of increased fitness conferred by gene mutations (24). For IAVs, adaptive evolution does not act on a single virion but on quasispecies, which represent populations of diverse variants that are genetically similar and collectively contribute to the characteristics of the population (24, 29). Here, we used deep sequencing to examine the mutations and frequency of the viral quasispecies in the upper (nasal wash) and lower (BALF) respiratory tracts of the P9 virus-infected pigs. We found five consensus amino acid mutations in all of the quasispecies in both compartments, indicating rapid mutational selection in limited serial passages. However, mutations in the HA gene exhibited marked differences between nasal wash and BALF, revealing evolutionary divergence of HA genes between upper and lower airways within an animal. The cause for this mutational divergence could be related to the virus itself, cell surface receptors, cell types, and antibody binding (30–32). Moreover, the frequencies of HA mutations varied in the upper and lower respiratory tracts. This finding was verified by sequencing single viral clones, which reflected complex combinations of dominant and/or sporadic mutations in HA proteins within each location of the airway (see Table S1 in the supplemental material). This phenomenon shares some similarities with the findings of Murcia et al. (33) with regard to the evolutionary dynamics of a Eurasian avian-like swine influenza virus along the natural transmission chain. They observed a highly dynamic mutational spectrum with both transient and fixed mutations in the HA gene derived from consecutive daily nasal virus populations. We speculate that there are two kinds of evolutionary processes at work in the convergent evolution of influenza virions, high-frequency mutagenesis and rapid allele fixation, and such an evolutionary scheme readily facilitates IAV infection and transmission in new host species.

Most reported triple-reassortant swine H1 viruses caused mild disease, were inefficiently transmitted through the air in ferrets (34, 35), and had restricted circulation in humans (36), even though they generally possess strong binding affinity for the α2,6-linked human-like receptor (37). Our previous (7) and present studies demonstrated that the parental rH1N1 virus had viral characteristics identical to those of these reported triple-reas-
sortant viruses. Crucially, we found that by passage nine in pigs, the replication and pathogenicity of three representative mutants (L2, N9, and L18) greatly exceeded those of the parental rH1N1 virus in both pigs and ferrets. More importantly, the three viruses had enhanced contact transmissibility in both pigs and guinea pigs. Their enhanced virulence and contact transmissibility were similar to that of the H1N1/2009 virus (38–40). Since the virulence and contact transmissibility were comparable between the L2, N9, and L18 viruses, these acquired traits were determined primarily by the five consensus mutations (PB1 A469T, PA I129T, NA N329D, NS1 N205K, and NEP T48N) found in all three virus species. Furthermore, guinea pig experiments demonstrated that the PB1 A469T mutation and the combined NS1 N205K and NEP T48N mutations in rH1N1 virus conferred in-contact transmissibility in guinea pigs.

Viral polymerase may be the driving component of early evolution of an IAV in a new host, and polymerase-enhancing mutations could contribute to the increased replication and virulence of IAVs (10, 25, 41). We observed significantly enhanced polymerase activity with the PB1 A469T mutation in the parental rH1N1 RNP complex. Residue position 469 locates to a site involved in RNA polymerase activity (42), and sequence analysis revealed that PB1 469T is conserved in pandemic H1N1/2009 isolates, which could be an important pathogenicity determinant of the pandemic H1N1/2009 virus. Our previous studies have demonstrated that the NS gene of H1N1/2009 origin is critical for contact transmission of the rH1N1 virus in guinea pigs (7). Zhang et al. have reported that the NS gene of H1N1/2009 virus can confer H5N1 virus transmission by respiratory droplet between guinea pigs (43). For the combined NS1 N205K and NEP T48N mutant virus, we could not readily verify which one of the two proteins plays a major role in guinea pig transmissibility, as both changes are the result of a single-nucleotide change. We do know that NS1 protein of IAV is a multifunctional virulence factor that inhibits host cell pre-mRNA processing and counteracts host cell antiviral responses. Residue 205 is located in the C terminus (residues 201 to 230) of NS1, and this tail has been implicated in the interaction of NS1 with cleavage and polyadenylation specificity factor (CPSF), poly(A)-binding protein II (PAB II), and host importins (44–46). It is likely that the NS1 N205K mutation plays a role in the pathogenesis of IAV by influencing these interactions. NEP, formerly called NS2, has been demonstrated to mediate the export of viral RNP complexes from the nucleus (47) and to act as a quantitative switch from viral transcription during early viral replication to favor late-stage production of genomic vRNPs (48). Additionally, NEP also has been implicated in recruiting a cellular ATPase to the cell membrane to aid the efficient release of budding virions (49, 50). Accordingly, NEP is proposed to play multiple biologically important roles during the life cycle of IAVs; thus, NEP T48N mutation could alter the replication and transmission characteristics of the rH1N1 virus.

A hallmark of the pandemic H1N1/2009 virus is its efficient transmission in humans (51, 52). In the present study, we found that the P9-derived L2 virus was efficiently transmitted through the air in ferrets (a recognized model of human influenza virus infection), as observed with the pandemic H1N1/2009 virus (27, 28). This suggests that the five consensus mutations are responsible for the enhanced virulence, and that the HA mutations (D187E, K211E, and S289N) in the L2 virus are linked to effective airborne transmission in ferrets. The receptor binding specificity

**FIG 10** In-contact transmission of mutant rH1N1 viruses in guinea pigs. Guinea pigs (in six groups of three) each were intranasally inoculated with 10⁶ TCID₅₀ of PB1 A469T (A), PA I129T (B), NA N329D (C), the combined NS1 N205K and NEP T48N (NS1 N205K & NEP T48N) (D), the combined PB1 A469T and PA I129T (PB1 A469T + PA I129T) (E), or the combined NA N329D, NS1 N205K, and NEP T48N (NA N329D + NS1 N205K & NEP T48N) (F) virus. At 1 dpi, each directly infected animal (individually housed) was moved to another cage holding a naive guinea pig. Viral titers of daily nasal washes were determined by TCID₅₀ assays on MDCK cells from the directly infected (squares with dotted lines) and in-contact guinea pigs (triangles with solid lines) and are presented as a function of days postvirus inoculation. Infections with the PB1 A469T mutant and combined NS1 N205K and NEP T48N mutant viruses were transmissible to one and two in-contact guinea pigs, respectively. The virus detection limit was 10 TCID₅₀.
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TABLE 3

Infection characteristics in: Directly infected ferrets

Proximally exposed ferrets

<table>
<thead>
<tr>
<th>Virus</th>
<th>Pulmonary consolidation</th>
<th>Microscopic lung lesions</th>
<th>Weight loss</th>
<th>Pyrexia</th>
</tr>
</thead>
<tbody>
<tr>
<td>L2</td>
<td>2/3 (20)</td>
<td>1/3 (20)</td>
<td>0/3 (0)</td>
<td>2/3 (38.3)</td>
</tr>
<tr>
<td>N9</td>
<td>2/3 (20)</td>
<td>1/3 (20)</td>
<td>0/3 (0)</td>
<td>2/3 (38.3)</td>
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<tr>
<td>L18</td>
<td>2/3 (20)</td>
<td>1/3 (20)</td>
<td>0/3 (0)</td>
<td>2/3 (38.3)</td>
</tr>
<tr>
<td>rH1N1</td>
<td>2/3 (20)</td>
<td>1/3 (20)</td>
<td>0/3 (0)</td>
<td>2/3 (38.3)</td>
</tr>
</tbody>
</table>

Note: Percentages are based on the pulmonary consolidation area, and the value for each animal is listed separately.

**Microscopic lung lesions** were assigned a score of 0 to 4. Scores are expressed as means ± SD during the observation time.

Weight loss is expressed as mean maximum percentages (%) of body weight. Weight loss to C and E Lungs from pigs directly infected L2, N9, and L18 viruses, along with lungs of L2 virus-exposed pigs, showed extensive edema, consolidation, and inflammatory cell exudate in the alveolar space. (D) The directly infected rH1N1 virus was isolated from ferrets directly infected L2, N9, L18, and rH1N1 viruses at 8 d.p.i. (E to H) Images shown are representative of three ferrets from three independent experiments. Scale bar, 200 μm.
have been highly feasible for the generation of the emergent pandemic virus to have begun in swine, although we are still unable to determine whether other host species were involved in the evolution of pandemic H1N1/2009 virus prior to its jump into humans. Altogether, our findings emphasize the importance of continued monitoring of influenza viruses in pigs.

ACKNOWLEDGMENTS

We thank Lu Qi, Meng Yu, Guanlong Xu, Yandi Wei, Huijie Gao, Xiaolin Zhu, and Seng Lai Giea for excellent technical assistance.

We thank Lu Qi, Meng Yu, Guanlong Xu, Yandi Wei, Huijie Gao, Xiaolin Zhu, and Seng Lai Giea for excellent technical assistance.

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