Interspecies Transmission of Reassortant Swine Influenza A Virus Containing Genes from Swine Influenza A(H1N1)pdm09 and A(H1N2) Viruses
Helen E. Everett, Bethany Nash, Brandon Z. Londt, Michael D. Kelly, Vivien Coward, Alejandro Nunez, Pauline M. van Diemen, Ian H. Brown, Sharon M. Brookes

Author affiliation: Animal and Plant Health Agency, Weybridge, UK
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Influenza A(H1N1)pdm09 (pH1N1) virus has become established in swine in the United Kingdom and currently co-circulates with previously enzootic swine influenza A virus (IAV) strains, including avian-like H1N1 and human-like H1N2 viruses. During 2010, a swine influenza A reassortant virus, H1N2r, which caused mild clinical disease in pigs in the United Kingdom, was isolated. This reassortant virus has a novel gene constellation, incorporating the internal gene cassette of pH1N1-origin viruses and haemagglutinin and neuraminidase genes of swine IAV H1N2 origin. We investigated the pathogenesis and infection dynamics of the H1N2r isolate in pigs (the natural host) and in ferrets, which represent a human model of infection. Clinical and virologic parameters were mild in both species and both interspecies and interspecies transmission was observed when initiated from either infected pigs or infected ferrets. This novel reassortant virus has zoonotic and reverse zoonotic potential, but no apparent increased virulence or transmissibility, in comparison to pH1N1 viruses.

The ability of swine to support replication of phylogenetically diverse influenza A virus (IAV) strains from avian and mammalian origin poses a public health risk because of the potential for viral antigenic change resulting in variants with zoonotic properties (1). This risk was highlighted by emergence of the influenza A(H1N1)pdm09 (pH1N1) swine-origin influenza virus during 2009 (2). This pH1N1 line is now enzootic and has also undergone reassortment resulting in emergence of multiple genotypes and 2 main enzootic subtypes, H1N2 (H1huN2), also designated lineage IB (3), and H3N2, through acquisition of human seasonal influenza virus-origin haemagglutinin (HA) or neuraminidase (NA) gene segments (4).

Since the global dissemination of pH1N1 virus, this lineage has also undergone reassortment with swine IAV strains endemic to the corresponding geographic region (5). Within Europe, this diversification of pH1N1 virus in pigs has increased the circulating swine IAVs from 4 to ≥25 genotypes (6,7). Diversification has been rapid; pH1N1 reassortant viruses incorporating enzootic swine virus HA and NA, or NA alone, detected during 2010 in the United Kingdom (8) and Italy (9), have emerged. The NA segment was N2 subtype in both instances. Subsequently, viruses have been identified that contain the NA segment from European enzootic H3N2 strains and the remaining 7 segments from pH1N1-origin viruses (10,11) or a further reassortant containing H3 of human seasonal origin (12).
Reassortment between enzootic and pH1N1 influenza viruses was also rapidly detected elsewhere, including Asia, where a reassortant swine IAV containing pH1N1-origin NA was described during 2010 (13), and in North America, where several reassortants containing HA and NA segments from enzootic viruses have been maintained since 2011 (14). These swine IAV reassortants from North America incorporate different combinations of the pH1N1 internal gene cassette (gene segments other than HA and NA) and invariably carry the pH1N1-origin matrix (M) protein gene.

Swine IAV H1N2 virus reassortment strains circulating in Brazil acquired NA gene segments from independent introductions of human seasonal H3N2 strains similar to those circulating in the late 1990s and an HA segment derived from human H1 strains circulating in the early 2000s. These lineages have subsequently reassorted with co-circulating pH1N1 strains (15) and viruses with a pH1N1-origin internal gene cassette have been isolated from wild boar (16) and swine (17).

One such H1H2 reassortant gave rise to a human clinical case of influenza in a worker on a swine farm (18). A reassortant H1N2 virus strain was also isolated from pigs in Chile; this virus incorporated human-origin HA and NA gene segments prevalent in the 1990s and the pH1N1 internal gene cassette (19). Human infection with swine-origin H1N2 viruses (H1N2v infection) has been reported in the United States, most notable being 3 human influenza cases linked to infection with a related co-circulating H1N2 swine IAV virus incorporating a pH1N1-origin M gene segment (20).

Genetic reassortment between enzootic swine and pH1N1 strains giving rise to H1N2 virus diversification has also been reported in Asia, including lineages in South Korea containing Eurasian avian-like swine HA and Korean swine H1N2 NA gene segments and the pH1N1-like internal gene cassette (21) and another reassortant strain associated with pig respiratory disease in China incorporating the HA, basic polymerase protein 2, and M genes of swine pH1N1 virus origin; the remaining gene segments were derived from a swine H3N2 strain (22).

We characterized a prototypical reassortant swine influenza A virus, A/swine/England/1382/2010 (H1N2r) (23), which has become enzootic in the pig population in the United Kingdom (6). This virus incorporates the genes encoding the envelope glycoproteins, HA and NA, from a European swine H1N2 subtype (which themselves derive from human-origin strains) and the remaining gene cassette encoding the internal proteins from swine-origin pandemic 2009 strains (8). Serologic assessment of potential exposure of pig industry workers in the United Kingdom to swine viruses during 2009–2010 showed that antibodies to H1N2 swine IAVs were present in 24% of persons, and this prevalence was increased relative to a comparator population (24). Because the potential risks associated with a novel combination of gene segments in the H1N2r isolate were unknown, we investigated the pathogenesis and infection dynamics of this virus in pigs, the natural host, and in ferrets, which are widely established as an animal model for investigating influenza and pandemic risk in humans (25,26).

Materials and Methods

Viruses
We isolated H1N2r virus A/swine/England/1382/2010 from nasal swab specimens from pigs that had clinical signs of mild influenza-like disease (8). Virus was propagated in embryonated specific pathogen-free chicken eggs according to standard methods (27), passaged once to obtain a virus stock, and characterized by using whole-genome sequencing (8) using reference influenza A virus strains (Table).

<table>
<thead>
<tr>
<th>Virus strain</th>
<th>Abbreviation</th>
<th>Gene segment origin†</th>
<th>Use</th>
<th>Reference</th>
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<td>H1mN2‡</td>
<td>pH1N1</td>
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</tr>
<tr>
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<td>H1mN2‡</td>
<td>H1mN2</td>
<td>H1</td>
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</tr>
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</table>

*Av, avian; HA, hemagglutinin; HI, hemagglutination inhibition; hu, human; NA, neuraminidase; pH1N1, influenza A(H1N1)pdm09; Sw, swine.
†Gene segment origin refers to the virus lineage origin of gene segments encoding the HA and NA envelope proteins or the remaining gene segments of the internal gene cassette.
‡Swine gene segments previously derived from human-origin viruses.
Organ Culture

Ferret organ cultures were prepared as described (Appendix, https://wwwnc.cdc.gov/EID/article/26/2/19-0486-App1.pdf). Organ cultures were inoculated by adsorbing virus for 1 h at 37°C onto the air interface of triplicate tissue sections. The culture was then washed and replenished with supplemented Dulbecco modified Eagle medium and incubated before virus detection 24–48 hours postinoculation (hpi) in combined supernatant and tissue lysate per sample. Control organ cultures were mock inoculated. Influenza A virus nucleoprotein (NP) was detected in tissues by using immunohistochemical (IHC) analysis as described (32).

Real-Time Reverse Transcription Quantitative PCR Analysis

Virus RNA was extracted by using a QIAmp Viral RNA Biorobot Kit (QIAGEN, https://www.qiagen.com) and assayed by using a real-time quantitative reverse transcription PCR (qRT-PCR) (33). Virus RNA was quantified as relative equivalent units (REUs) (34) against a 10-fold dilution series of RNA prepared from the inoculum stock with a titer of 10^5 50% egg infectious dose (EID_{50})/mL. REU measures the amount of virus RNA present and not infectivity. However, it can be inferred from the linear relationship with the dilution series that REU values are proportional to the amount of infectious virus present.

We used Prism software (GraphPad, https://www.graphpad.com) for statistical analyses. We determined the area under the curve of the shedding profile for each animal and compared between groups by using the Tukey multiple comparisons test and 1-way analysis of variance. Positive-sense RNA encoding the virus M gene was quantified by using a positive strand-specific real-time qRT-PCR (Appendix).

In Vivo Studies

All animal studies were reviewed and approved by the Animal and Plant Health Agency Ethical Review Panel. Studies were conducted according to the Animal (Scientific Procedures) Act of 1986 and Animal Research: Reporting of In Vivo Experiments guidelines.

High health status, 6–8-week-old, male Landrace hybrid pigs and male fitch ferrets (weight range 750–1,000 g, maximum age 3 months) were bred in the United Kingdom. Before infection, all animals were confirmed free of influenza A virus RNA by performing real-time qRT-PCR analysis of nasal swab specimens (pigs) or nasal washes (ferrets) (33). These animals also showed negative results in a hemagglutination inhibition (HI) test (35) using 4 influenza A virus antigens (pH1N1, H1avN1, H3N2, and H1N2 viruses) appropriate for the pig population in the United Kingdom (Table). Animals were subcutaneously implanted with a biothermal idENTICHIP (Destron Fearing, http://destronfearing.com) for daily temperature monitoring, and a clinical scoring system was used for daily clinical monitoring (36,37).

Experimental Infection

We intranasally infected animals by using protocols appropriate for the species and nasal structure. We infected pigs by using a MAD Nasal Intranasal Mucosal Atomization Device (Wolfe-Tory Medical Inc., https://www.lmaco.com) to deliver droplets with a diameter of 30–100 μm. A total dose of 10^6 EID_{50} units diluted in phosphate-buffered saline to a final volume of 4 mL (2 mL/nostril) was administered. We anesthetized ferrets and infected them by intranasal droplet instillation of a total dose of 10^6 EID_{50} in 0.4 mL inoculum (0.2 mL/nostril).

Pathologic Analysis

For each species, postmortem examination was conducted for 4 animals at 3 days postinoculation (dpi) and 5 dpi, as well as for 2 animals at 7 dpi. Gross pathology was assessed and tissue samples were collected in 10% (vol/vol) phosphate-buffered formalin for histopathologic analysis. Lesions were scored as described for ferrets (37) and pigs (36). Influenza A virus NP was detected in tissues by using IHC analysis as described (36,37).

Experimental Infection and Transmission Studies

Six animals of each species were directly infected for study I (intraspecies transmission) and study II (interspecies transmission). In study I, immunologically naive animals (n = 6) of the same species were placed in direct contact at 1 dpi. In study II, animals of 1 species (n = 6) were infected and immediately thereafter cohoused animals of the other species (n = 6) were reintroduced. Pigs were housed on the floor and immediately adjacent to ferret cages (minimum distance 30 cm), and the floor of the ferret cage was at the height of the heads of the pigs. Virus shedding was monitored daily from 1 dpi through 12 dpi in nasal swab specimens from pigs (1/nostril) or nasal wash samples from ferrets (1 mL/animal). Serum samples were obtained before infection and at the end of the study at 21 dpi. An overdose of barbiturates was used for euthanasia of all animals.

Swab and Tissue Specimen Processing

Daily nasal swab specimens, which were pooled for each pig, were eluted in 1 mL Leibovitz L-15 medium
with l-glutamine supplemented with 100 U/mL penicillin, 1,000 μg/mL streptomycin, and 1% fetal bovine serum (all from GIBCO). Ferret nasal wash samples were obtained by using 1 mL phosphate-buffered saline (GIBCO). Tissues were analyzed by preparing a 10%–20% [wt/vol] homogenate in 1 mL supplemented Leibovitz-15 medium by using an Omni GLH Homogenizer (Omni International, https://www.omni-inc.com).

Serologic Analysis
Serum samples were analyzed by using an influenza A virus NP competitive ELISA (IDVet) according to the manufacturer’s specifications and dilutions of 1:40 for pig serum samples and 1:10 for ferret serum samples. Antibody titers to the H1N2r virus were measured by HI as described for pig (35) or ferret (38) serum samples.

Results
H1N2r Virus Ex Vivo Replication
To investigate H1N2r virus replication in the ferret animal model, we inoculated ferret organ cultures with virus strains at the same viral multiplicity of infection or mock-inoculated. We quantified influenza A virus replication by using real-time qRT-PCR for virus RNA at 24 hpi and 48 hpi. Replication of the H1N2r isolate was comparable to that of swine pH1N1 and H1N2 strains and a human-origin pH1N1 strain in tracheal cultures (Figure 1, panel A). We also detected comparable amounts of virus RNA in lung cultures at 24 hpi; these amounts decreased slightly by 48 hpi for some strains (Appendix Figure). We confirmed H1N2r virus replication in tracheal cultures by positive IHC labeling for influenza A virus NP (Figure 1, panel B). A strand-specific PCR detected replicating virus-positive sense RNA in infected tracheal cultures (Figure 1, panel C), which decreased between 24 hpi and 48 hpi, most likely reflecting virus replication kinetics. These in vitro assay results showed productive replication of the swine-origin H1N2r virus in ferret cells, but we observed no selective growth advantage linked to the novel gene constellation.

Pathologic Assessment
We then investigated the pathogenicity profile after H1N2r infection of pigs, the natural host, or ferrets, which represent a surrogate model for human infection. We assessed pathologic changes at 3, 5, and 7 dpi in both species and observed gross pathologic changes at 3 dpi in 2 of 4 directly infected pigs. In 1 of these pigs, a large portion of a single lung lobe was consolidated by lesions consistent with influenza A virus infection. We also found gross pathologic changes in the lungs in all pigs at 5 dpi, with a reduction by 7 dpi. In infected ferrets, small sporadic lesions common to influenza A virus infection occurred, and individual animals had lesions in the respiratory turbinates (5 dpi) and salivary glands (7 dpi), as well as in the left cranial lung lobe in 2 directly infected ferrets at 3 dpi and 5 dpi.

Histopathologic changes were consistent with mild inflammatory disease. For pigs (Figure 2, panel A), inflammation was evident throughout the respiratory tract at all timepoints (e.g., in respiratory turbinates [upper respiratory tract] and lung lobes [lower respiratory tract]) at 3, 5, and 7 dpi. In contrast, for ferrets, most pronounced inflammation occurred in...
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the respiratory turbinates and in the salivary glands, initially at 5 dpi and increasing by 7 dpi, which indicated a predominantly upper respiratory tract infection (Figure 2, panel B).

Influenza A virus NP labeling showed replicating virus in pig and ferret tissues, notably in pig respiratory turbinates, at 5 dpi (Figure 2, panel C). For pig lung tissues, we detected virus in the right mid lung at 3 dpi and in all lung lobes at 5 dpi (Figure 2, panel E), with evidence of virus clearing by 7 dpi. For ferret tissues, virus NP labeling was more pronounced at 7 dpi, as shown for respiratory turbinates (Figure 2, panel D) and lungs (Figure 2, panel F). We observed intense IHC virus labeling in salivary glands of ferrets. These animals showed an increase in virus labeling from 3 dpi through 7 dpi (Figure 2, panel G).

H1N2r Virus Infection

In study I (Figure 3, panel A), we directly infected 6 animals and then placed them in direct contact with 6 uninfected animals of the same species 1 day later. Clinical signs from H1N2r infection were unapparent or mild (1–2 days) in pigs and ferrets, whether directly infected or contact animals, and there was no major change in body temperature (≤1°C above baseline). We observed clinical signs in 3 of 12 pigs: 2 pigs had mild rhinitis at 1 dpi, and 1 pig had mild respiratory signs (sneezing) at 5 dpi. Clinical signs in ferrets were limited to transient rhinitis in some animals and could have been caused by infection or other factors, such as sampling.

Shedding and Transmission Profile of H1N2r Virus

We monitored virus RNA shedding daily in nasal swab specimens (pigs) or washes (ferrets) and quantified results by using real-time qRT-PCR. Nasal shedding started in directly infected animals at <1 dpi and in uninfected contacts at 1–2 days after contact (Figure 3, panels B, C). These results from study I confirmed transmission between infected and uninfected animals of the same species. Peak amounts of virus RNA were shed by directly infected animals at 2–5 dpi. In comparison with directly infected or contact pigs (Figure 3, panel B), directly infected or contact ferrets (Figure 3, panel C) shed virus for a longer duration and also shed a larger total quantity of virus. Significantly greater (p<0.0001) quantities of virus RNA were shed by directly infected ferrets in comparison with all other groups, and also by contact...
ferrets in comparison with contact pigs, as shown by area under the curve analysis (Figure 3, panel D). A greater variation in shedding occurred between individual ferrets than between pigs, particularly among contact animals.

In study II (Figure 3, panel A), directly infected pigs or ferrets (representing a human model) were cohoused with animals of the other species to assess interspecies transmission of the H1N2r virus and zoonotic potential. Direct infection of pigs (Figure 3, panel E) or ferrets (Figure 3, panel F) resulted in similar infection dynamics as for intraspecies study I. When infected ferrets were cohoused with uninfected pigs (Figure 3, panel E), the pigs became infected after a considerable 8–10-day lag, and virus shedding profiles differed between recipient pigs. These infection kinetics might indicate that some pigs became infected from their penmates, demonstrating possible onward transmission of virus and potential for the H1N2r virus to disseminate in a susceptible population. Ferrets were readily infected within 4 days when cohoused with infected pigs (Figure 3, panel F), and the infection profile was similar to that observed for intraspecies transmission, suggesting that all ferrets were infected synchronously by the infected pigs. Nasal shedding profiles observed for animals infected directly with H1N2r were comparable with profiles for human pH1N1 influenza viruses A/England/195/2009 and A/California/07/2009 (36,37,39,40).

**Serologic Analysis**

Serologic analysis by competitive ELISA for NP, which is present within virions, demonstrated that, in interspecies transmission study II, infected and contact ferrets seroconverted by 21 dpi/days after contact (Figure 4, panel A). Most infected pigs also seroconverted and showed lower final antibody titers, although 2 of 6 pigs cohoused with infected ferrets did not seroconvert within the study period. HI titers, which measure exposure to the viral envelope HA protein, were positive against the H1N2r challenge strain, indicating seroconversion of all infected or contact animals (Figure 4, panel B). Both assays showed that the antibody response was stronger in ferrets than pigs, perhaps corresponding to the higher virus load and more prolonged infection in ferrets.

**Discussion**

Our study showed that pigs and ferrets were susceptible to infection with the H1N2r virus and showed clinical signs and virologic parameters indicating mild disease. Longitudinal postmortem analysis
indicated that this virus could be more adapted to a swine host because infection disseminated more rapidly throughout the respiratory tract, and distribution of infection in the lung on all postmortem days was more evident in pigs than in ferrets. However, in contrast to pigs, ferrets shed a larger total quantity of virus for an extended duration, and seroconversion occurred in all infected and contact ferrets by 21 dpi. However, interpretation of data is limited by the different intranasal infection protocols and nasal sampling techniques for the 2 species. All animals had increased specific antibody titers postinfection. However, in comparison with ferrets, titers in pigs were lower and, in some contact pigs, were below the positive threshold. These findings are comparable with outcomes reported for pH1N1 virus infection of pigs and ferrets (36,37,39,40) and a comparison of swine H1 viruses, including H1N2v strains, when infecting ferrets (41). A North American reassortant H1N1 swine IAV incorporating pH1N1-origin polymerase acidic, NP, and M genes also reportedly showed similar replication kinetics to a swine IAV pH1N1 isolate when infecting ferrets (41). In a similar fashion, an H1N2 reassortant strain from Chile containing the pH1N1 internal gene cassette was also transmitted between ferrets by direct and indirect routes (19).

Swine are known to support replication of viruses with a wide range of HA activation pH values, whereas ferrets support replication of a narrower pH activation range for HA (40). In addition, it has been suggested that successful transmission of influenza A viruses requires a balance between the HA and NA activities (43). Therefore, the HA and NA combination of the H1N2r isolate that we studied is compatible with replication and transmission in both mammalian hosts.

In conclusion, our study demonstrates that a swine H1N2r virus can be readily transmitted from swine to ferrets. This finding could have occurred as a consequence of the room layout, the different degree of aerosolization of virus shed from ferrets or pigs, or the anatomic and physiologic differences in the pig and ferret nasal tract. Once animals became infected, whether directly or indirectly, the shedding profile was consistent within the same species. These findings indicate that the threat associated with pH1N1 reassortant viruses from swine or human origin is no greater from either donor species, although zoonotic infection is clearly plausible and might be augmented by the presence of the pH1N1 internal gene cassette.

In conclusion, our study demonstrates that a swine H1N2r virus can be readily transmitted
between mammalian species. Although this virus does not display enhanced virulence compared with other swine IAV or human pH1N1 viruses (36,39), it nevertheless shows high interspecies and intraspecies transmissibility. This virus strain represents a newly emergent reassortant virus that could enhance the genetic diversity of circulating strains and contribute to influenza A virus genotypic change at the human–animal interface, thereby increasing the potential for generating new viruses with altered disease phenotype or fitness for new host species.

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About the Author

Dr. Everett is a research scientist at the Animal and Plant Health Agency, Weybridge, UK. Her research interests are the pathogenesis of mammalian influenza viruses, mechanisms of immune-mediated protection from influenza, and strategies for assessing and controlling outbreak risk.

References


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Address for correspondence: Helen E. Everett, Animal and Plant Health Agency, New Haw, Addlestone, Surrey KT15 3NB, UK; email: helen.everett@apha.gov.uk

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Appendix

Organ Culture Preparation

Ferret tracheal cultures were prepared as follows. Petri dishes were prepared with plugs of 1% (wt/vol) low-gelling temperature agarose (Sigma, https://www.sigmaaldrich.com) in water. Plugs were immersed in Dulbecco modified Eagle medium (DMEM) medium containing 2 mol/L L-glutamine and antimicrobial drugs (200 U/mL penicillin, 2,000 μg/mL streptomycin, and 2.5 μg/mL amphotericin B (all from GIBCO, https://www.thermofisher.com), then incubated at 37°C in an atmosphere of 5% CO₂. Tracheal tissue was immersed in prewarmed, supplemented DMEM and incubated at 37°C in an atmosphere of 5% CO₂ for 20 min. The medium was replaced ≥3 times, and the tissue was dissected and placed onto sterile filter paper overlaying an agarose plug, with the filter paper forming a wick into the medium.

To culture lung tissue, a lung lobe was filled through the bronchi with phosphate-buffered saline supplemented with antimicrobial drugs and maintained at 37°C. Low-gelling temperature agarose (1% and 4% [wt/vol] in phosphate-buffered saline) was equilibrated to 38°C. The lung lobe was filled with 1% agarose and set with 4% agarose inside a syringe. Once solidified, the top of the syringe was removed, the lung extruded, and 3–4-mm sections were cut and incubated 18 h in supplemented DMEM at 37°C in an atmosphere of 5% CO₂. Sections were moved to a 6-well plate, and the outer 4% agarose layer was removed.

Positive Strand-Specific PCR

Positive-sense RNA encoding the virus matrix protein gene was quantified to assess levels of virus replication. Reverse transcription was conducted by using the M-MLV Reverse Transcription Kit (Promega, https://www.promega.com) and reverse primer NcRev1 (5’-AGTAGAAACAAAGGTAGTTTT-3’) at a final concentration of 2.5 μmol/L. Real-time PCR was
conducted by using the Quantifast Kit (QIAGEN, https://www.qiagen.com) according to the manufacturer’s specifications using the forward and reverse primers M808F (5′-GCAGATGCAGCGATTCAAG-3′) and swM1003R (5′-CACTCTGCTGTTCTGGATAT-3′) at final concentrations of 0.04 μmol/L and the probe M945RPro (FAM-5′-AGGCCCTCTTTCTAAACCTATT-3′-TAMRA at a final concentration of 0.02 μmol/L. Cycling parameters for an MxPro3000 Instrument (Agilent, https://www.agilent.com) were 1 cycle of 95°C for 3 min and 40 cycles at 94°C for 10 s, 55°C for 30 s, and 72°C for 30 s. Data readings were collected during annealing and analyzed by using MxPro software (Agilent).

**Appendix Figure.** Infection and replication of swine H1N2r virus in ferret lung organ cultures. Quantity of influenza A virus RNA in ferret lung ex vivo organ cultures at 24 h and 48 h postinoculation with swine viruses H1N2r (A/swine/England/1382/2010), Sw pdm09 (A/swine/England/1353/2009), H1N2 (A/swine/England/997/2009), and human isolate Hu pdm09 (A/England/195/2009). Results are log₁₀ REU in combined supernatants and tissue lysates for each sample. Error bars indicate SEM for triplicate cultures. hpi, hours postinoculation; Hu, human; H1N2r, reassortant swine influenza A virus; pdm, pandemic; REU, relative equivalent unit; Sw, swine.