Swine Influenza Virus (H1N2) Characterization and Transmission in Ferrets, Chile

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Phylogenetic analysis of the influenza hemagglutinin gene (HA) has suggested that commercial pigs in Chile harbor unique human seasonal H1-like influenza viruses, but further information, including characterization of these viruses, was unavailable. We isolated influenza virus (H1N2) from a swine in a backyard production farm in Central Chile and demonstrated that the HA gene was identical to that in a previous report. Its HA and neuraminidase genes were most similar to human H1 and N2 viruses from the early 1990s and internal segments were similar to influenza A(H1N1)pdm09 virus. The virus replicated efficiently in vitro and in vivo and transmitted in ferrets by respiratory droplet. Antigenically, it was distinct from other swine viruses. Hemagglutination inhibition analysis suggested that antibody titers to the swine Chilean H1N2 virus were decreased in persons born after 1990. Further studies are needed to characterize the potential risk to humans, as well as the ecology of influenza in swine in South America.

In the past decade, variant swine-origin influenza A viruses (swIAVs) of subtypes H1N2 and H3N2 have been associated with human infections, particularly in persons with close contact to swine (1–7), although evidence of human-to-human transmission is minimal. The emergence and subsequent pandemic caused by the unique influenza A(H1N1)pdm09 virus (pH1N1) from swine in 2009 highlights the public health threat posed by swIAVs, especially in populations having little preexisting immunity (8). Therefore, it is imperative to understand the circulation of swIAVs on a global level.

While our knowledge of the global diversity and evolution of swIAVs has increased, less information has been available regarding swIAVs circulating in South America. Recently, subtypes H1N1, H1N2, and H3N2 were identified in commercial pigs in Chile (9). Phylogenetic analysis from 18 swIAVs sequenced from a 2012 cross-sectional sampling of 22 production companies in Chile (95% of their commercial swine) demonstrated extensive genetic diversity in the hemagglutinin (HA) segment (9). The human seasonal H1-like viruses were unique to swine in Chile and not closely related to other H1 swIAVs. These viruses represented 2 independent introductions from humans, with Chile H1 human I clade most closely related to human viruses from the late 1980s and Chile H1 human II clade most closely related to human viruses from the early 1990s (9). No further phylogenetic or virus characterization information was available.

In addition to commercial pig farming, Chile has an extensive backyard production system, a common form of animal production throughout the world. Backyard production farms (BPFs) are a particular public health concern given poor biosecurity and close interaction between humans and multiple animal species (10). During 2014, we conducted active backyard production system surveillance in Chile’s main swine production area (11) and isolated a subtype H1N2 swIAV virus (A/swine/Chile/YA026/2014; sw/Chile). We performed phylogenetic analyses and full genomic sequencing on this virus to determine its relationship with other known influenza isolates. We propagated it in primary human bronchial and swine respiratory epithelial cells, as well as C57Bl/6 mice, to determine its infectiosity. We also characterized its transmission using ferrets. Hemagglutination inhibition (HI) assays with human serum samples were done to assess the risk this virus poses for human infection, and serological analyses were conducted with pig serum samples from Chile to determine the prevalence of the virus and its risk for emergence.

Materials and Methods

Study Area and Nasal Swab Sampling

Active surveillance was conducted as described (11) in 40 BPFs in Central Chile, representing 90% of the backyard

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production system. During this time, only 3 farms had swine (n = 3) available for sampling. We collected nasal samples using disposable sterile swabs, which were placed in 1 mL universal transport media. None of the animals were symptomatic.

**H1N2 Virus Isolation**

We screened nasal swab samples by real-time reverse transcription PCR for matrix gene as described (12). PCR-positive samples were centrifuged at 8,000 rpm for 15 min and the clarified supernatant (200 µL) inoculated on MDCK cells in 24-well plates in duplicate and incubated at 37°C in 5% CO₂ for 1 h. Following incubation, cells were washed then incubated with Eagle’s minimum essential medium (MEM; MediaTech, Manassas, VA, USA) supplemented with 2 mM glutamine, bovine serum albumin, and 1 µg/mL TPCK (L-1-tosylamide-2-phenylethyl chloromethyl ketone)–trypsin until cytopathic effect was observed. Then, cell supernatant was collected, centrifuged, and stored at −80°C as described (12). Only 1 virus was isolated: A/swine/Chile/YA026/2014 (H1N2) (sw/Chile).

**Cells**

MDCK cells were cultured in MEM supplemented with 2 mM glutamine and 10% fetal bovine serum (Gemini BioProducts, West Sacramento, CA, USA) and grown at 37°C in 5% CO₂. A549 cells were cultured in Dulbecco’s minimum essential medium (DMEM; Lonza, Walkersville, MD, USA) supplemented with 4.5 g/L glutamine and 10% fetal bovine serum and grown at 37°C in 5% CO₂. Well-differentiated (transepithelial resistance >300 Ω × cm²), primary normal human bronchial epithelial (NHBE) cells were grown as previously described (13). In brief, cells were plated on collagen-coated membranes in bronchial epithelial basal media (Lonza) supplemented with SingleQuots additives (Lonza) and media changed every 48 h. When cells became 100% confluent, basal media was replaced with DMEM:F12 (Lonza) supplemented with SingleQuots, and apical media was removed. Every 48–72 h,
Swine Influenza Virus (H1N2), Chile

the apical surface was washed with 0.9% sodium chloride solution (Sigma, St. Louis, MO, USA). Cells were incubated at the air-liquid interface at 37°C in 5% CO₂. Primary swine nasal epithelial cells (sNECs) were prepared as described (14). In brief, nasal swab specimens were collected, placed in 50 mL conical tubes, and incubated in phosphate-buffered saline (PBS) at 4°C for 10–15 min before centrifuging for 10 min at 300 × g at 4°C. The pellet was resuspended in supplemented DMEM:F12 and incubated on collagen-coated plates at 37°C in 5% CO₂. Cells were plated onto transwells and, when 100% confluent, moved to an air–liquid interface to differentiate for 1–3 wk. sNECs were moved to 33°C 1 d before infection.

Virus Sequencing
Sanger sequencing of viral stocks was performed by the St. Jude Hartwell Center (12), and deep amplicon sequencing and assembly were conducted as described previously (15). Sequences can be obtained in GenBank (accession nos. CY207245–CY207252).

Viral Propagation
A/swine/Chile/YA026/2014 (H1N2, sw/Chile), A/Memphis/3/1987 (H1N1, Mem/87), A/swine/Iowa/13-1015/2010 (H1N2, sw/IA), A/swine/Indiana/26-0818/2011 (H1N2, sw/IN), A/swine/North Carolina/18161/2002 (H1N1, sw/NC), A/swine/Italy/1310-2/1995 (H1N1, sw/IT), and

Figure 2. Phylogenetic trees comparing the internal genes of swine influenza virus (H1N2) from Chile (red) and reference viruses. We performed phylogenetic analysis for the matrix (A), nucleoprotein (B), nonstructural (C), polymerase acid (D), polymerase basic (PB) 1 (E), and PB2 (F) gene segments by using RAxML with 200-bootstraps replicates (21). Purple indicates location of control influenza A(H1N1)pdm09 CA/09 virus. Scale bars indicate number of substitutions per site. Detailed phylogenetic trees for these genes are provided in online Technical Appendix Figures 1–6 (https://wwwnc.cdc.gov/EID/article/23/2/16-1374-Techapp1.pdf).
A/California/04/2009 (pH1N1, CA/09) viruses were propagated in the allantoic cavity of 10-day-old, pathogen-free, embryonic chicken eggs at 37°C and viral titers determined by 50% tissue culture infectious dose (TCID\textsubscript{50}) analysis as previously described (16,17). These viruses were used as comparators for all studies except the ferret experiments.

**Phylogenetic Analysis**

We performed sequence assembly and editing by using BioEdit version 7.2.5 (18) and sequence alignment with MUSCLE version 3.8.3 (19). Reference sequences were obtained from the Influenza Virus Resource at NCBI (20). Phylogenetic relationships for each gene were inferred by maximum likelihood, incorporating a general time-reversible model of nucleotide substitution with a gamma-distributed rate variation among sites by using RAxML version 8.0 (27). A bootstrap resampling process of 200 replicates was implemented to provide statistical robustness to each node.

**Swine and Human Serum Samples**

During 2013–2015, swine serum samples (n = 266, 1–24 animals/BP) were collected from 80 BPFs in Central Chile, including the Valparaiso, Metropolitan, and Libertador General Bernardo O’Higgins regions. During 2009–2015, human serum samples were collected as part of an ongoing prospective observational study at the University of North Carolina Family Medicine Center (Chapel Hill, NC, USA). We chose a subset of these samples, collected 28–32 d after seasonal influenza vaccination; grouped them according to birth decade (1920–1929, 1930–1939, 1940–1949, 1950–1959, 1960–1969, 1970–1979, 1980–1989, and 1990–1999; n = 7–22/age group); and balanced for sex and race whenever possible (22).

**Hemagglutination Inhibition Assay**

Ferret antiserum was generated against sw/Chile, Mem/87, sw/IA, CA/09, sw/IN, sw/NC, and sw/IT viruses. HI assay was conducted as previously described (23). In brief, antiserum was treated with receptor-destroying enzyme (Denka Seiken, Tokyo, Japan), inactivated, and diluted 1:10. Serial dilutions of receptor-destroying enzyme–treated serum were incubated in duplicate with each virus for 15 min at room temperature, followed by 30 min incubation at 4°C with 0.5% turkey erythrocytes. HI titer was defined as the reciprocal dilution of the last well that reacted.

**In Vitro Replication**

MDCK cells were infected with a multiplicity of infection (MOI) of 0.01 for 1 h at 37°C. Cells were washed 3 times to remove unbound virus, and infected cells were cultured in MEM containing 0.075% bovine serum albumin and 1 μg/mL TPCK-treated trypsin. Aliquots of culture supernatants were collected at 6 h postinfection (hpi), 24 hpi, 48 hpi, and 72 hpi and immediately stored at −80°C. For infection of NHBE cells and sNECs, the apical surface was washed twice and incubated with serum-free DMEM containing virus for 2 h at 37°C, after which both apical and basal media were removed and fresh growth medium was added to the basal chamber as described (24). At 6 hpi, 24 hpi, 48 hpi, 72 hpi, and 96 hpi, DMEM was added to the apical surface, and cells were incubated for 30 min at 37°C (for NHBE) or 33°C (for sNEC). Apical media was collected and stored at −80°C. Viral titers were determined by TCID\textsubscript{50} on MDCK cells followed by Reed-Munch analysis (25).

**Animal Infections**

Six- to 8-week-old female BALB/c mice (Jackson Laboratory, Bar Harbor, ME, USA) (n = 11 mice/group) were lightly anesthetized with isoflurane and intranasally inoculated with PBS or 10\textsuperscript{6} TCID\textsubscript{50} units of virus in 25 μL PBS. Mice were monitored daily for signs of infection and weighed every 48 hpi (26). At 3 d postinfection (dpi) and 6 dpi, 3 control and 3 infected mice were humanely killed; lungs were harvested and homogenized in 1 mL PBS and nasal washes were collected as previously described (27). In brief, an incision was made in the trachea and 500 μL of PBS was flushed through the upper respiratory tract and collected. Viral titers were determined by TCID\textsubscript{50} (16,24).

Data are representative of 2 separate experiments. For transmission studies, 9- to 15-week-old male ferrets (n = 2, Triple F Farms, Sayre, PA, USA) were intranasally inoculated with 10\textsuperscript{6} TCID\textsubscript{50} units in 1 mL PBS. Twenty-four hours later, naive ferrets (n = 2 mice/group) were either
placed in direct contact with infected ferrets or housed in separate cages (respiratory contact) separated by ≈5 inches. Nasal washes were collected every 2 dpi for viral titration and serum samples collected at 21 dpi for HI analysis as described (28).

Ethics Statement
All sampling was approved by the ethics and biosecurity committee of Faculty of Veterinary Science, University of Chile, and collected by trained veterinarians according to Food and Agriculture Organization guidelines (http://www.fao.org/3/a-ak738e.pdf). Animal experiments were approved by the St. Jude Children’s Research Hospital institutional biosafety committee and animal care and use committee and were in compliance with the Guide for the Care and Use of Laboratory Animals. These guidelines were established by the Institute of Laboratory Animal Resources and approved by the

Figure 3. Testing for influenza in swine from backyard production farms (BPFs) in Central Chile. Swine serum samples collected from BPFs were analyzed for antibodies against influenza by nucleoprotein (NP)–specific ELISA, and hemagglutination inhibition analysis of serum samples from NP ELISA–positive farms was conducted to determine subtype. A) Results of influenza testing by BPF location. Red circles indicate location of positive farms, and gray circles indicate location of negative farms. B) Pie charts showing percentage of swine at each sampling site seropositive for each virus type: sw/Chile (red), CA/09 (yellow), sw/IA (green), and sw/NC (blue). Red star indicates location of sw/Chile H1N2 virus isolate. Insets indicate region of Central Chile covered by this study (red shading).
Governing Board of the US National Research Council. All human procedures were approved by the Biomedical Institutional Review Board at the University of North Carolina.

Statistical Analysis
We performed statistical analyses using JMP statistical software (SAS Institute, Cary, NC, USA) and GraphPad Prism software (GraphPad Software Inc., La Jolla, CA, USA). The Kruskal-Wallis test \((\alpha = 0.05)\) was used to analyze nonparametric data, while normally distributed data were analyzed by 2-way analysis of variance with virus strain and time postinfection as main effects. The Student \(t\)-test was used for posthoc comparisons between the viruses, and Tukey’s honest significant difference was used for posthoc comparisons among the times postinfection. Differences were considered significant at \(p<0.05\).

Results

Phylogenetic Similarity of Chilean H1N2 and Human Seasonal Viruses
Phylogenetic analysis demonstrated that sw/Chile H1N2 belongs to the \(\delta\)-like H1 subgroup (Figure 1, panel A) and is phylogenetically similar to the swIAVs from Chile described by Nelson et al. \((9)\). The HAs of these swIAVs are closely related to the HAs of human seasonal H1 viruses from the late 1980s and early 1990s (closest relative A/Suita/1/89, 94% nucleotide identity), although they are separated by long branch lengths and represent over 15 years of evolution \((9)\). Unfortunately, only the HA segments from the 2012 Chilean swIAVs were available in public databases, so we were unable to compare full genomes.

The sw/Chile neuraminidase (NA) segment arose from a reassortment between human seasonal H1N1 and H3N2 viruses (Figure 1, panel B). However, long branch length indicates extensive genetic divergence from human N2 sequences, with the closest sequence being an early 1990s North American H3N2 virus (A/New York/762/1993, 93% nucleotide identity). This difference can be indicative of possible temporal and species-specific adaptations, leading to genetic and antigenic differences with contemporary human N2 viruses. The sw/Chile virus internal genes are pH1N1 in origin, consistent with the high reassortment rate between pH1N1 and swine viruses (Figure 2; online Technical Appendix Figures 1–6, https://wwwnc.cdc.gov/EID/article/23/2/16-1374-Techapp1.pdf) \((9,29)\). Overall, while sw/Chile does appear to originally be of human origin, detection of these viruses in multiple BPFs suggests these viruses might have become established in the swine population in Chile.

Antigenicity and Seroprevalence of Chilean H1N2 Virus
HI assays with a panel of ferret antiserum against classical swIAV (sw/IN and sw/NC from \(\beta\) and \(\gamma\) subgroups, respectively); pandemic CA/09 virus; sw/Iowa (from \(\delta\) subgroup); and human seasonal H1 virus Mem/87 showed...
that sw/Chile H1N2 virus was antigenically distinct from other H1 viruses (Table). To determine seroprevalence in swine within the surveillance area, serum samples (n = 266, 1–24 animals/BPF) were collected from swine on 80 BPFs in Central Chile in 2013–2015. Forty-eight (60%) BPFs were positive by nucleoprotein-specific ELISA, and the prevalence of positive pigs (86/266, 32.3%) on each BPF ranged from 14.2% to 100% animals (Figure 3, panel A). HI assays performed with a panel of influenza viruses on the ELISA-positive samples demonstrated that 14/48 (29.1%) BPFs and 22/86 pigs were positive for ≥1 of the test viruses. Antibodies against sw/Chile were identified in BPFs located in 3 regions of Central Chile (Figure 3, panel B). Of concern, 11/14 (78.6%) of the positive BPFs, representing 15/22 (68.2%) of the positive samples, were positive for multiple influenza strains, suggesting that different influenza viruses are co-circulating in backyard swine in Central Chile (Figure 3, panel B). Not all nucleoprotein ELISA–positive samples could be subtyped.

**Chile H1N2 Virus in Primary Swine and Human Cells**

Swine possess both mammalian (α-2,6) and avian (α-2,3) influenza receptors (30,31). Receptor binding assays (23) revealed that all viruses used in this study had greater binding affinity for mammalian-type (α-2,6–linked sialic acid) receptors (online Technical Appendix Figure 7). These viruses also replicated effectively in MDCK cells; differentiated, primary NHBE cells; and sNECs cultured at the air-liquid interface. Cells were infected at MOI 0.01 and viral titers determined at the indicated times by TCID₅₀ analysis on MDCK cells (23). All viruses replicated to similar titers in MDCK cells (Figure 4, panel A) and sNECs (Figure 4, panel C), except for an increased replication of CA/09 virus in MDCK cells. However, significant differences in replication were observed in primary NHBE cells. The human seasonal Mem/87 and sw/Chile viruses replicated to lower titers than the other swine viruses, suggesting reduced fitness in human cells (Figure 4, panel B). Overall, these studies demonstrate that sw/Chile can productively replicate in human and swine cells.

**Chile H1N2 Virus in Mice**

Recently identified swine H1N2 viruses from China replicated efficiently in mice and pigs (32); however, no data were available on viruses from South America. Thus, we intranasally inoculated Balb/c mice with 10⁵ TCID₅₀ units of each virus and monitored weight loss for 14 dpi. Viral titers were measured 3 dpi and 6 dpi (Figure 5). CA/09 virus caused considerable weight loss, as did sw/IA (H1N2, δ₁ subclade), which was similar to CA/09, sw/IN (H1N2, β subclade), and sw/NC (H1N1, γ₂ subclade) viruses. Increased weight loss was associated with efficient replication in lungs (Figure 5, panel B). In contrast, the Mem/87 and sw/Chile viruses caused minimal illness and had lower replication in lungs (Figure 5, panels A and B). The sw/Chile virus replicated the most effectively in the upper respiratory tract compared with the other swine viruses, which reached titers similar to that of CA/09 virus (Figure 5, panel C). These data show that while sw/Chile virus can replicate in mice, it causes minimal illness. The genetic basis for differences in pathogenicity and tropism between the swine strains warrants further investigation.

**Figure 5.** Replication of influenza viruses in vivo. To evaluate pathogenicity in mice, 6- to 8-week-old BALB/c mice (n = 11 mice/group/experiment) were infected with 10⁵ 50% tissue culture infectious dose (TCID₅₀) units of the indicated viruses and weight loss was monitored for 14 days postinfection (dpi) (A). At 3 dpi and 6 dpi, lungs (B) and nasal washes (C) were collected from 3 mice/group and viral titers were determined by TCID₅₀ analysis. Data are presented as mean ± SEM. *p<0.05 versus sw/Chile virus.
Chile H1N2 Virus in Respiratory Droplets
Eurasian swine H1N2 viruses transmit in ferrets by direct contact and, in some cases, by respiratory droplet (33, 34). To test the transmissibility of sw/Chile virus, we intranasally inoculated ferrets with 10⁶ TCID₅₀ units of sw/Chile, Mem/87, or sw/IA viruses; at 1 dpi, naive ferrets were either placed in direct contact or housed in cages adjacent to the donor ferrets to monitor respiratory droplet transmission. Nasal washes were collected every 2 dpi to assess viral shed, and animals were monitored through 21 dpi. All of the donor animals shed virus, and all viruses transmitted through direct contact and respiratory droplet with some differences (Figure 6). Overall, the swine viruses replicated to a higher titer than the Mem/87 virus did in the donor and direct contact animals; swine viruses also transmitted faster by respiratory droplet. Respiratory contacts of sw/Chile virus–inoculated animals were shedding virus by 6 dpi, whereas Mem/87 virus–inoculated animals only showed detectable nasal washes after 8 dpi and 10 dpi. The sw/IA virus appeared to transmit faster and replicate to higher titer than did the sw/Chile and Mem/87 viruses (Figure 6). All animals seroconverted at 21 dpi. These studies highlight that both swine H1 δ viruses transmitted effectively in ferrets.

**Figure 6.** Evaluation of H1 virus transmission in ferrets. Donor ferrets (black lines; n = 2 ferrets/group) were inoculated with 10⁶ 50% tissue culture infectious dose (TCID₅₀) units of H1 influenza virus sw/Chile (A), Mem/87 (B), or sw/IA (C) viruses. Naive ferrets (n = 2) were either placed in the same cage with the infected group (direct contact, blue lines) or housed in separate cages (respiratory transmission, red lines) and nasal washes were collected on the indicated day postinfection for virus quantification by TCID₅₀ analysis. Lines represent individual animals. Dashed line represents limit of detection of the assay.

Decreased HI Antibody Titer in Persons Born after 1990
Because the HA and NA of sw/Chile were derived from human seasonal strains originating around 1990, we hypothesized that persons born after 1990 would have decreased HI antibody titers. To test this, we conducted HI analyses against CA/09, Mem/87, sw/IA, and sw/Chile viruses on serum samples from 137 persons in North America collected in 2010–2015; samples were grouped by decade of birth (n = 7–22 samples/group). Most persons born before 1990 had titers against all 4 viruses (Figure 7). However, few persons born after 1990 had titers against sw/Chile (9%; Figure 7, panel B); sw/IA (31.8%; Figure 7, panel C); or Mem/87 (27.2%; Figure 7, panel D) viruses. Persons with titers ranging from 1:10 to 1:160 were all born in 1990 or 1991. In contrast, most (91.7%) persons born after 1990 had titers against pH1N1 virus, ranging from 1:320 to 1:10,240 (Figure 7, panel A).

Discussion
Our data show that the swine H1N2 virus from Chile is antigenically distinct from other H1 viruses, replicates efficiently in mammalian cells, can transmit by respiratory droplet, and might pose a risk to immunologically naive persons born after 1990. Our study furthers the data recently published on prevalence and HA phylogeny of Chile swIAVs (9). We show that these viruses are circulating outside of commercial swine herds in BPFs, where animal–human interaction is high.

Since emergence of pH1N1 in 2009, reverse transmission of the virus from humans to pigs has been documented worldwide, resulting in several reassortments with classical swine and seasonal human viruses (1, 2, 4, 5, 9, 12, 35). As of July 2016, variant (v) influenza viruses have caused over 380 human infections in the United States alone. While
most of these infections have been H3N2v viruses, 8 have been H1N2v viruses. These infections are particularly concerning because most infections occurred in children (persons <18 years of age) who had direct or indirect exposure to swine (2). Census estimates indicate that 50% of BPFs in Chile have ≥1 person born after 1990, and 35% have ≥1 person born after 2000 (C. Hamilton-West, unpub. data). Our data indicate that younger age groups have reduced HI titers to sw/Chile virus; however, studies with serum samples obtained from throughout the population of Chile are warranted to better determine the risk these viruses pose to swine herds.

This study is subject to several limitations. First, these studies were limited to 1 virus isolate. A panel of swine viruses from Chile isolated from different years and regions warrant study, especially if genetic differences are found among viruses. Second, these results represent a limited number of the backyard production system within Chile. Enhanced swine surveillance throughout Chile and South America is needed. Third, studies at the animal–human interface are required to decipher the risk circulating swIAVs pose to humans. We were limited to serum samples from persons living in North America, and further work should be done on South America populations, especially with those exposed to swine. Antigenic cartography on serum samples from persons involved in backyard farming would provide invaluable new information on transmission of swine viruses to humans. Finally, further work is needed to improve understanding of swIAV genes involved in transmission in the ferret model.

Despite no evidence of sustained human-to-human transmission, increased risk for human infection with H1N2v viruses calls for further study and enhanced monitoring. Risk for possible emergence of zoonotic strains has been demonstrated worldwide. H1N2 viruses isolated from pigs in China replicated efficiently in mice and pigs (32,36,37), while viruses from South Korea and Europe transmitted to and even caused death in ferrets (33,34,38). Future work will focus on how these reassortants arose in Chile BPFs and the role of specific genetic differences in pathogenicity and transmissibility. In addition, further studies for seroprevalence in Chile BPF workers and antigenic cartography with the sw/Chile virus are necessary. Studies at the animal–human interface are needed to determine seroprevalence, identify broadly protective antibodies, and characterize T-cell responses, and active surveillance is needed to uncover infection rates to better determine the risk for zoonotic transmission of swine-origin H1N2 to human health. In summary, our findings highlight the need for continued, vigilant influenza virus surveillance in Chile and throughout South America.

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Swine Influenza Virus (H1N2) Characterization and Transmission in Ferrets, Chile

Technical Appendix

Technical Appendix Figure 1. Phylogenetic tree of the matrix (M) gene of Chilean H1N2 virus. Phylogenetic analysis was performed on complete M genome sequences using maximum likelihood (RAxML) and incorporating a general time-reversible model of nucleotide substitution with a gamma-distributed rate variation among sites. The Chilean H1N2 virus is shown in red, the A/California/04/2009 pandemic H1N1 virus is in purple and the swine viruses used as comparators throughout the studies are shown in blue. Scale bars represent the number of substitutions per site.
**Technical Appendix Figure 2.** Phylogenetic tree of the nucleoprotein (NP) gene of Chilean H1N2 virus. Phylogenetic analysis was performed on complete NP genome sequences using maximum likelihood (RAxML) and incorporating a general time-reversible model of nucleotide substitution with a gamma-distributed rate variation among sites. The Chilean H1N2 virus is shown in red, the A/California/04/2009 pandemic H1N1 virus is in purple and the swine viruses used as comparators throughout the studies are shown in blue. Scale bars represent the number of substitutions per site.
**Technical Appendix Figure 3.** Phylogenetic tree of the nonstructural (NS) gene of Chilean H1N2 virus. Phylogenetic analysis was performed on complete NS genome sequences using maximum likelihood (RAxML) and incorporating a general time-reversible model of nucleotide substitution with a gamma-distributed rate variation among sites. The Chilean H1N2 virus is shown in red, the A/California/04/2009 pandemic H1N1 virus is in purple and the swine viruses used as comparators throughout the studies are shown in blue. Scale bars represent the number of substitutions per site.
Technical Appendix Figure 4. Phylogenetic tree of the polymerase acid (PA) gene of Chilean H1N2
Phylogenetic analysis was performed on complete PA genome sequences using maximum likelihood
(RAxML) and incorporating a general time-reversible model of nucleotide substitution with a gamma-
distributed rate variation among sites. The Chilean H1N2 virus is shown in red, the A/California/04/2009
pandemic H1N1 virus is in purple and the swine viruses used as comparators throughout the studies are
shown in blue. Scale bars represent the number of substitutions per site.
Technical Appendix Figure 5. Phylogenetic tree of the polymerase basic 1 (PB1) gene of Chilean H1N2 virus. Phylogenetic analysis was performed on complete PB1 genome sequences using maximum likelihood (RAxML) and incorporating a general time-reversible model of nucleotide substitution with a gamma-distributed rate variation among sites. The Chilean H1N2 virus is shown in red, the A/California/04/2009 pandemic H1N1 virus is in purple and the swine viruses used as comparators throughout the studies are shown in blue. Scale bars represent the number of substitutions per site.
Technical Appendix Figure 6. Phylogenetic tree of the polymerase basic 2 (PB2) gene of Chilean H1N2 virus. Phylogenetic analysis was performed on complete PB2 genome sequences using maximum likelihood (RAxML) and incorporating a general time-reversible model of nucleotide substitution with a gamma-distributed rate variation among sites. The Chilean H1N2 virus is shown in red, the A/California/04/2009 pandemic H1N1 virus is in purple and the swine viruses used as comparators throughout the studies are shown in blue. Scale bars represent the number of substitutions per site.
Technical Appendix Figure 7. Sialic acid–binding affinities of H1 viruses. Both human (A-B) and swine (C-F) H1 viruses had increased binding to α-2,6–linked sialic acids suggesting increased mammalian binding efficiency. Data are representative of 3 replicates per virus. Data is presented as mean ± SEM.
Technical Appendix Figure 8. Weight and temperature during ferret transmission of H1 viruses. Ferrets (n = 2/group) were inoculated with 10^6 units of H1 influenza virus (black lines). Naive ferrets (n = 2) were placed either in the same cage with the infected group (direct contact, blue lines) or housed in separate cages (respiratory transmission, red lines) and weight and temperature data was collected every 24 hours postinfection for (A) Mem/87, (B) swine/IA, and (C) swine/Chile viruses. Lines represent individual animals. Green shading indicates normal temperature range for ferrets (http://weaselwords.com/ferret-facts/).