

Antigenic and genetic evolution of contemporary swine H1 influenza viruses in the United States



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ABSTRACT

Several lineages of influenza A viruses (IAV) currently circulate in North American pigs. Genetic diversity is further increased by transmission of IAV between swine and humans and subsequent evolution. Here, we characterized the genetic and antigenic evolution of contemporary swine H1N1 and H1N2 viruses representing clusters H1- α (1A.1), H1- β (1A.2), H1pdm (1A.3.3.2), H1- γ (1A.3.3.3), H1- δ 1 (1B.2.2), and H1- δ 2 (1B.2.1) currently circulating in pigs in the United States. The δ 1-viruses diversified into two new genetic clades, H1- δ 1a (1B.2.2.1) and H1- δ 1b (1B.2.2.2), which were also antigenically distinct from the earlier H1- δ 1-viruses. Further characterization revealed that a few key amino acid changes were associated with antigenic divergence in these groups. The continued genetic and antigenic evolution of contemporary H1 viruses might lead to loss of vaccine cross-protection that could lead to significant economic impact to the swine industry, and represents a challenge to public health initiatives that attempt to minimize swine-to-human IAV transmission.

1. Importance

The hemagglutinin (HA) protein of influenza A virus (IAV) is the primary target of protective immune responses and the major component in vaccine formulation. However, repeated introductions of non-swine IAV into swine populations and the processes of antigenic shift and drift result in virus evolution and potential mismatches between vaccines and circulating strains. Further, the increasing diversity of swine IAV represents a challenge for public health initiatives, and assessment of the risk of interspecies transmission of viruses relies on assessing antigenic diversity relative to human population immunity. In this study, we found that antigenic drift of IAV in the U.S. pig population resulted in seven distinct antigenic phenotypes of the H1 subtype currently circulating. We identified changes in the HA protein associated with the observed antigenic drift, suggesting that these amino acids may be important antigenic sites. These data demonstrate that recent evolution of H1 swine IAV resulted in novel genetic clades with distinct antigenic phenotypes that are unlikely to be protected by current vaccine formulations.

2. Introduction

Influenza A virus (IAV) is endemic in pigs globally, and three different subtypes currently co-circulate in North American swine: H1N1, H1N2, and H3N2 (Anderson et al., 2013; Lorusso et al., 2013). Swine have been indicted as a potential source of reassortant viruses that can be transmitted to humans (Brown, 2000; Vijaykrishna et al., 2011). Two major events illustrate this potential: the introduction of the swine-origin 2009 pandemic H1N1 virus (H1N1pdm09) with widespread global morbidity and mortality (Smith et al., 2009b), and the occurrence of more than 300 cases of swine-origin H3N2 IAV (H3N2v) (Jhung et al., 2013). However, the transmission of human IAV into swine populations is a frequent occurrence, and the repeated introduction and subsequent evolution of these human viral lineages in swine populations has a critical role in the genetic diversity of swine IAV (Nelson et al., 2012, 2014).

The first documented occurrence of bidirectional transmission between swine and humans dates to the 1918 H1N1 pandemic (Smith et al., 2009a), with the resulting swine strain referred to as classical H1N1 (cH1N1). The cH1N1 remained antigenically and genetically stable in the U.S. for nearly a century. In the late 1990s, a novel triple-

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reassortant H3N2 virus was detected in U.S. swine herds (Zhou et al., 1999) and reassorted with endemic cH1N1, leading to a period of rapid diversification of the cH1N1 hemagglutinin (HA) into three distinct HA clades: H1- α (1A.1), H1- β (1A.2), and H1- γ (1A.3.3.3) (Lorusso et al., 2011; Vincent et al., 2008b). Viruses containing HA and/or neuraminidase (NA) derived from human-seasonal IAV were introduced to U.S. pigs in the early 2000s (Karasin et al., 2006; Vincent et al., 2009), giving rise to two HA phylogenetic clades, H1- δ 1 (1B.2.2) and H1- δ 2 (1B.2.1) (Anderson et al., 2013; Lorusso et al., 2013). The 2009 H1N1 pandemic viruses were reintroduced from humans into U.S. pigs, resulting in reassortment with endemic swine viruses (Ducatez et al., 2011) and the establishment of a sixth swine H1 genetic clade (H1N1pdm09: 1A.3.3.2). This dynamic evolution is not restricted to the HA; N2 genes were derived from two human seasonal-lineages introduced in 1998 and 2002 or N1 genes from the classical or H1N1pdm09 lineages (Anderson et al., 2013; Lorusso et al., 2013). Moreover, the current diversity of lineages within IAV circulating in swine populations is complicated by antigenic drift of the HA (de Jong et al., 1999), and this gradual evolution may alter antigenic properties of IAV that result in vaccine failure (Both et al., 1983; Luoh et al., 1992). However, antigenic diversity is not always linearly correlated with genetic diversity, and few amino acid mutations can lead to biologically significant antigenic differences that may facilitate immune escape (Both et al., 1983; Luoh et al., 1992).

Vaccination against IAV in swine is an important tool to limit clinical disease, but it may not prevent infection and/or transmission. Products currently available in the U.S. provide good protection against homologous strains, however their efficacy against antigenically distinct viruses may diminish (Vincent et al., 2010, 2008a). Vaccine strains should be updated when there is loss in cross-reactivity against contemporary strains to improve the match between vaccine antigens and circulating antigenic diversity. Although genetic analysis of the HA gene has utility in predicting vaccine efficacy (Neher et al., 2016), antigenic cross-reactivity between field-sourced viruses and vaccine strains remains the gold standard for vaccine strain candidate selection. Thus, understanding the molecular epidemiology of IAV circulating in swine populations in association with antigenic properties are crucial to improve vaccine composition to better control IAV in pigs.

Several studies have mapped the antigenic and genetic evolution of H3N2 viruses in swine (Abente et al., 2016; de Jong et al., 2007; Lewis et al., 2014), and recently a comprehensive study characterized the high level global antigenic diversity of H1 and H3 viruses circulating in human and swine (Lewis et al., 2016). However, an extensive description of the links between genetic diversity, amino acid identity, and antigenic phenotype remains unclear for H1 viruses in the U.S. Here, we performed a comprehensive characterization of the genetic and antigenic evolution of swine H1N1 and H1N2 viruses, representing all genetic clades currently circulating in pigs in the U.S. We then conducted an in-depth antigenic characterization on contemporary strains from the H1- δ (1B.2.1 and 1B.2.2) clades due to their predominance in surveillance detections and apparent genetic expansion. Further, we examined the underlying genetic basis for significant antigenic differences among these circulating H1- δ viruses.

3. Materials and methods

3.1. Genetic evolution and amino acid sequence analysis

All available swine IAV HA and NA sequences from H1N1 and H1N2 viruses collected in the U.S. were downloaded from the Influenza Research Database (IRD) (Squires et al., 2012; Zhang et al., 2016) on December 5, 2016. To restrict our analyses to relevant field viruses, we excluded sequences with “lab” or “laboratory” host. From these data, alignments for the HA and the N1 and N2 NA genes were generated using MAFFT v7.294 (Katoh et al., 2002; Katoh and Standley, 2013). In addition, sequences with 100% identity were deleted using mothur

v1.36.0 (Schloss et al., 2009), and poor quality data was removed using two criteria: a sequence was removed if > 50% of the gene was missing or if it had more than 5 nucleotide base ambiguities. This process resulted in a set of 3211 non-identical H1 HA, 1606 N1 and 1406 N2 NA swine IAV sequences that represent the full extent of the published swine H1N1 and H1N2 genetic diversity in the U.S. (Table S1). For each alignment (i.e., the HA, the NA-N1 and the NA-N2) we inferred the best-known maximum likelihood tree using RAXML (v8.2.4; (Stamatakis, 2014)) employing the rapid bootstrap algorithm, a general time-reversible (GTR) model of nucleotide substitution, and Γ -distributed rate variation among sites. The statistical support for individual branches was estimated by bootstrap analysis with the number of replicates determined automatically using an extended majority-rule consensus tree criterion (Pattengale et al., 2010). These analyses used the computational resources of the USDA-ARS computational cluster Ceres on ARS SCINet.

To determine the temporal evolution and relative genetic diversity of U.S. swine H1 HA, we implemented a time-scaled Bayesian approach on a second dataset. We downloaded complete HA H1N1 and H1N2 swine IAV genes collected from 2000 to present from the IRD on December 5, 2016. Given deep evolutionary divergence between the two major H1 HA lineages, we separated these data into the classical lineage H1 HA including H1- α (1A.1), H1- β (1A.2), H1- γ (1A.3.3.3), and H1N1pdm09 (1A.3.3.2) ($n = 2208$), and human-seasonal H1- δ lineage H1- δ 1 (1B.2.2) and H1- δ 2 (1B.2.1) ($n = 1740$). From each of these datasets, we randomly sampled to create smaller datasets of 750 HA genes to overcome computational limitations (i.e., we randomly sampled 707 H1- δ lineage viruses and added 43 reference antigens; similarly, we randomly sampled 713 H1-classical lineage viruses, then added the 37 reference antigens). These data were then aligned using MAFFT v7.294 (Katoh et al., 2002; Katoh and Standley, 2013), and HA genes that were duplicated due to the addition of reference antigen data were removed. The resultant data were then screened using root-to-tip regression in TempEst v1.5 (Rambaut et al., 2016) and sequences with incongruent genetic divergence and sampling date were removed (Hicks and Duffy, 2012) resulting in a dataset of 730 and 726 HA genes for the H1- δ and H1-classical lineages respectively. The remaining data were analyzed using an uncorrelated relaxed lognormal molecular clock (Drummond et al., 2006), the SRD06 codon position model (Shapiro et al., 2006) that partitions codon positions (1 + 2 positions and 3 position) with an HKY85 + Γ substitution model applied to each partition. To reconstruct population dynamics we used the coalescent-based Gaussian Markov random field (GMRF) method with time-aware smoothing (Minin et al., 2008). The precision function in BEAUTi was used to sample uniformly within a one-year or one-month window for those viruses for which an exact date of collection was not available. All analyses were implemented in BEAST v1.8.4 (Drummond et al., 2012) with the BEAGLE library (Ayres et al., 2012) with two independent analyses of 100 million generations with sampling every 10,000 generations. Convergence of runs was checked in Tracer v1.6.0, runs were combined with LogCombiner v1.8.4, and evolutionary history was summarized and visualized using an annotated maximum clade credibility tree using TreeAnnotator v1.8.4 and FigTree v1.4.2. These analyses used the computational resources of the USDA-ARS computational cluster Ceres on ARS SCINet.

Hemagglutinin sequences used for antigenic characterization were translated to amino acids, trimmed to the HA1 domain, and aligned using MAFFT v7.294 (Katoh et al., 2002; Katoh and Standley, 2013). The HA1 amino acid alignments were used to identify amino acid substitutions likely to be clade-defining or to have resulted in antigenic differences between or within a genetic clade. We followed a criterion in which a substitution was considered pertinent as clade-defining if all (or all but one) viruses in one clade had the same amino acid at a specific position in comparison to a different amino acid in the same position of all (or all but one) viruses in another clade (Lewis et al., 2014, 2011; Smith et al., 2004). Antigenic outliers within the H1- δ

(1B.2) lineage were identified as strains that were mapped at an antigenic distance greater than 2 antigenic units (AU) when compared to their respective antigenic cluster representative (centroid antigen). These outliers were then compared through pairwise comparisons of antigenic distance and amino acid similarity. Pairwise amino acid distances were inferred using a JTT model of amino acid substitution with Γ -distributed rate variation among sites; pairwise antigenic distances were extracted from the antigenic map. Those strains that had a pairwise antigenic distance > 3 AU and a pairwise amino acid distance of \leq 4% were aligned and all pairwise amino acid mutations were determined.

3.2. Viruses

Seventy-two swine influenza H1N1 and H1N2 viruses were selected as hemagglutination inhibition (HI) test antigens and/or antigens for swine IAV-antisera production, representing 15 U.S. states (IA, MN, SD, NE, KS, MO, IL, WI, MI, OH, IN, KY, NC, OK, TX). Viruses were included to represent historical and contemporary circulating strains from each of the major H1 clades currently circulating in U.S. pigs isolated between 1930 and 2014 (Anderson et al., 2013). The HA phylogeny was generated as described above to identify genetic clades with contemporary data, then strains were selected with an emphasis on the oldest and youngest viruses in the clade, and a particular emphasis on H1- δ (1B.2.1 and 1B.2.2) clades. Viruses were also selected based on amino acid differences observed compared to other strains within clades. Selection was constrained by limitations posed by isolate availability. Viruses isolated from outbreaks of respiratory disease in pigs from routine diagnostic cases were obtained from the University of Minnesota Veterinary Diagnostic Laboratory (UMVDL, kindly provided by Dr. Marie Culhane) and from the USDA-National Animal Health Laboratory Network (NAHLN) voluntary swine IAV surveillance system repository held at the National Veterinary Service Laboratories. In addition, eight human H1 viruses were used as antigens: three H1N1pdm09 (1A.3.3.2), a human-seasonal virus related to the swine H1- δ 1 and H1- δ 2 clades (1B.2.2 and 1B.2.1 respectively), and four human H1 vaccine strains. Viruses were grown in Madin-Darby canine kidney (MDCK) cells or embryonated chicken's eggs. To obtain antigens for immunization, clarified virus from cell culture supernatant or allantoic fluid was concentrated by ultracentrifugation and inactivated by ultraviolet (UV) irradiation. Inactivation of the virus was confirmed by failure to replicate in two serial passages in MDCK cells. Antigens were prepared at approximately 128 HA units per 50 μ l in phosphate buffered saline (PBS) and a commercial oil-in-water adjuvant (Emulsigen D, MVP Laboratories, Inc., Ralston, NE) was added at a 1:5 ratio. Viruses used in this study are listed in Table 1.

3.3. Antisera production

Three-week-old cross-bred healthy pigs demonstrated to be free of IAV and IAV-antibodies were obtained from a high-health herd to be used for IAV antisera production. All pigs were treated with ceftiofur crystalline-free acid (Excede; Pfizer, New York, NY) and enrofloxacin (Baytril; Bayer Animal Health, Shawnee Mission, KS). Two pigs per antigen were immunized intramuscularly with adjuvanted inactivated virus. Two or three doses were given 2–3 weeks apart. When HI titers to homologous virus reached at least 1:160, pigs were humanely euthanized with pentobarbital (Fatal Plus, Vortech Pharmaceuticals, Dearborn, MI) for blood collection. Sera was obtained through centrifugation and stored at -20°C until use. Pigs were cared for in compliance with the Institutional Animal Care and Use Committee of the National Animal Disease Center, USDA-ARS.

3.4. Serological assay and antigenic characterization

Prior to HI testing, sera were heat inactivated at 56°C for 30 min,

Table 1

Swine and human H1N1 and H1N2 influenza A virus (IAV) used as test antigens in hemagglutination inhibition assays and for the generation of antisera (in bold).

Viruses	H1 clade	Subtype	Accession number
A/swine/Iowa/15/1930	H1- α	H1N1	AF091308
A/swine/Iowa/1945	H1- α	H1N1	EU139824
A/swine/Wisconsin/1/1968	H1- α	H1N1	EU139825
A/swine/Iowa/1973	H1- α	H1N1	EU139826
A/swine/Minnesota/02053/2008	H1- α	H1N1	HM461762
A/swine/Minnesota/02093/2008	H1- α	H1N1	HM461794
A/swine/North Carolina/36883/2002	H1- β	H1N1	EU139829
A/swine/Iowa/00239/2004	H1- β	H1N1	EU139832
A/swine/Iowa/02096/2008	H1- β	H1N1	HM461842
A/swine/Kentucky/02086/2008	H1- β	H1N1	HM461786
A/swine/Nebraska/02013/2008	H1- β	H1N1	HM461834
A/swine/North Carolina/02084/2008	H1- β	H1N1	HM461826
A/swine/Minnesota/A01076180/2009	H1- β	H1N1	JX042526
A/swine/Minnesota/03012/2010	H1- β	H1N1	KY290625
A/swine/Missouri/A01076190/2010	H1- β	H1N1	JQ809747
A/swine/Minnesota/37866/1999	H1- γ 2	H1N1	EU139827
A/swine/Minnesota/1192/2001	H1- γ	H1N2	CY098468
A/swine/Minnesota/00194/2003	H1- γ	H1N2	EU139830
A/swine/Kansas/00246/2004	H1- γ	H1N2	EU139831
A/swine/Ohio/511445/2007	H1- γ	H1N1	EU604689
A/swine/North Carolina/02023/2008	H1- γ	H1N1	HM461818
A/swine/Ohio/02026/2008	H1- γ	H1N1	HM461778
A/swine/Missouri/02060/2008	H1- γ	H1N1	HM461810
A/swine/Minnesota/03011/2010	H1- γ	H1N1	KY290621
A/swine/Indiana/3062/2010	H1- γ	H1N1	KY290626
A/swine/Illinois/3134/2010	H1- γ	H1N2	KY290622
A/swine/Minnesota/03003/2010	H1- γ	H1N1	KY290631
A/California/04/2009	pH1N1	H1N1	FJ966082
A/Mexico/4108/2009	pH1N1	H1N1	GQ149654
A/New York/18/2009	pH1N1	H1N1	FJ984355
A/swine/Illinois/32974/2009	pH1N1	H1N1	GU480922
A/swine/Illinois/02991/2010	pH1N1	H1N1	KY290630
A/swine/Minnesota/03019/2010	pH1N1	H1N1	KY290624
A/swine/North Carolina/03029/2010	pH1N1	H1N2	KY290623
A/swine/Indiana/03053/2010	pH1N1	H1N2	KY290627
A/swine/Illinois/5265-1/2010	pH1N1	H1N1	GU984405
A/swine/Minnesota/8761/2010	pH1N1	H1N1	GU984411
A/swine/Illinois/07003243/2007	H1- δ 1	H1N2	FJ638314
A/swine/Texas/01976/2008	H1- δ 1	H1N2	HM461850
A/swine/Minnesota/02011/2008	H1- δ 1	H1N2	HM461802
A/swine/Iowa/02039/2008	H1- δ 1a	H1N2	HM461770
A/swine/Iowa/02998/2010	H1- δ 1a	H1N2	HM193850
A/swine/Minnesota/A01134353/2011	H1- δ 1a	H1N2	JQ906881
A/swine/South Dakota/A01365260/2013	H1- δ 1a	H1N2	KF746329
A/swine/Minnesota/A01392045/2013	H1- δ 1a	H1N2	KF715130
A/swine/Nebraska/A01492366/2014	H1- δ 1a	H1N2	KJ549771
A/swine/Missouri/A01411322/2014	H1- δ 1a	H1N2	KJ437545
A/swine/Iowa/A01476931/2014	H1- δ 1a	H1N2	KP288072
A/swine/South Dakota/A01481702/2014	H1- δ 1a	H1N2	KM013804
A/swine/Illinois/A01047020/2010	H1- δ 1b	H1N2	JQ756323
A/swine/Minnesota/3128/2010	H1- δ 1b	H1N1	CY158457
A/swine/Illinois/3120/2010	H1- δ 1b	H1N2	KY290619
A/swine/Minnesota/03002/2010	H1- δ 1b	H1N2	KY290620
A/swine/Minnesota/A01301731/2012	H1- δ 1b	H1N2	KC020465
A/swine/South Dakota/A01349341/2013	H1- δ 1b	H1N2	KC844209
A/swine/Missouri/A01444664/2013	H1- δ 1b	H1N2	KC562218

(continued on next page)

Table 1 (continued)

Viruses	H1 clade	Subtype	Accession number
A/swine/Nebraska/A01290601/2013	H1- δ 1b	H1N2	KF791371
A/swine/Iowa/A01432370/2013	H1- δ 1b	H1N2	KC534990
A/swine/Indiana/A01260356/2013	H1- δ 1b	H1N2	KF680052
A/swine/Minnesota/A01366540/2014	H1- δ 1b	H1N2	KJ175145
A/swine/Indiana/A01260471/2014	H1- δ 1b	H1N2	KJ650529
A/swine/Illinois/A01410405/2014	H1- δ 1b	H1N2	KJ605069
A/swine/Iowa/A01410774/2014	H1- δ 1b	H1N2	KJ859606
A/swine/Illinois/00685/2005	H1- δ 2	H1N1	FJ638298
A/swine/North Carolina/00573/2005	H1- δ 2	H1N1	FJ638306
A/swine/Minnesota/07002083/2007	H1- δ 2	H1N1	FJ611898
A/swine/Iowa/02955/2010	H1- δ 2	H1N2	KY284571
A/swine/Missouri/03013/2010	H1- δ 2	H1N2	KY290628
A/swine/Ohio/003295/2010	H1- δ 2	H1N2	KY284570
A/swine/Minnesota/A01076626/2010	H1- δ 2	H1N1	JQ906871
A/swine/Michigan/A01259001/2012	H1- δ 2	H1N2	KC508596
A/swine/Ohio/A01351184/2013	H1- δ 2	H1N2	KF313148
A/swine/North Carolina/A01290598/2013	H1- δ 2	H1N2	KF791368
A/swine/Oklahoma/A01409770/2014	H1- δ 2	H1N2	KJ437589
A/swine/North Carolina/A01475606/2014	H1- δ 2	H1N2	KM027346
A/New Caledonia/20/1999	H1-Human	H1N2	DQ508857
A/Michigan/2/2003	H1-Human	H1N2	CY016324
A/Memphis/8/2003	H1-Human	H1N2	CY122316
A/Solomon Islands/3/2006	H1-Human	H1N2	EU124177
A/Brisbane/59/2007	H1-Human	H1N2	CY058487

then treated with a 20% suspension of kaolin (Sigma–Aldrich, St. Louis, MO) followed by adsorption with 0.5% turkey red blood cells (RBCs). HI assays were performed by testing the reference antisera panel against the 80 H1 viruses according to standard techniques (Kitikoon et al., 2014). Geometric mean titers obtained by \log_2 transformation of reciprocal titers were used for the comparison. Antigenic cartography was used for the quantitative analyses of the antigenic properties of swine H1 viruses as previously described (de Jong et al., 2007; Lewis et al., 2014; Lorusso et al., 2011; Smith et al., 2004).

4. Results

4.1. Genetic evolution of swine H1 influenza viruses currently circulating in the U.S

The H1 phylogeny revealed that 7 H1 genetic clades circulated during the period of study (Fig. 1A and B; Fig. S1); all of these clades were reported previously (Anderson et al., 2016). Of note, the H1- δ 1 (1B.2.2) clade evolved to form two independent clades (Fig. 1A) with strong statistical support. The viruses in these clades are designated as H1- δ 1a and H1- δ 1b, corresponding to 1B.2.2.1 and 1B.2.2.2, respectively, following the global swine H1 HA nomenclature (Anderson et al., 2016). The H1- δ 1a (1B.2.2.1) and H1- δ 1b (1B.2.2.2) clades, as well as all others, were supported by within (< 7%) and between (> 7%) clade nucleotide distance, statistical support greater than 70% at the clade defining node, and continual detection of viruses within each clade over the last 5 years (Anderson et al., 2016, 2013). Coalescent reconstruction of the H1- δ (1B.2) lineages (Fig. 1A) revealed yearly oscillation and a trend for increased relative genetic diversity following the emergence and reassortment of H1N1pdm09 (1A.3.3.2) with H1- δ viruses after 2009. In contrast, there was little evidence of

yearly seasonal patterns in the relative genetic diversity in the classical swine H1 lineage (Fig. 1B). Though the overall relative genetic diversity of both lineages is similar, the relative genetic diversity of the H1- δ (1B.2) lineage appears to have consistently increased from introduction in the early 2000s to present (Fig. 1A) whereas the relative diversity within the classical lineage remained more stable (Fig. 1B).

The H1 viruses circulating in pigs in the U.S. were paired with four distinct NA gene lineages: two N2 gene lineages derived from either 1998 or 2002 human-seasonal virus lineages, or an N1 of either classical swine N1 or pandemic N1 lineage. The classical N1 lineage represents the majority of the N1, whilst the pandemic N1 is becoming less frequent (Fig. S2A). Most of the H1 clades derived from the classical H1 were paired with an N1 lineage, while the majority of δ -clade viruses were paired with an N2 lineage. However, more than half of the contemporary swine H1 viruses circulating in the U.S. have an N2 gene, of which the majority is derived from the 2002 human-seasonal lineage (Fig. S2B). The δ -clades show a specific distribution pattern for either 1998 or 2002 lineage N2 genes, with H1- δ 1a (1B.2.2.1) and H1- δ 1b (1B.2.2.2) paired more frequently with a 2002 N2 lineage and the H1- δ 2 (1B.2.1) paired more frequently with a 1998 N2 lineage (Fig. S2B).

4.2. Antigenic diversity among contemporary swine H1 viruses

HI data (Table S2) were used to characterize the antigenic relationships of contemporary swine H1 viruses by antigenic cartography, focusing on the human-seasonal derived H1- δ viruses (Figs. 2 and S3). These data allow the calculation of antigenic distance between all pairs of antigens, with more than 2 AU between strains considered significant (1 AU is equivalent to a 2-fold difference in HI titer). Due to the antigenic diversity observed for the H1 viruses, antigenic clusters were separated based on genetic clades for improved visualization. For each antigenic cluster, the most centrally located strain with antigenic distances < 2 AU to the highest number of strains in that cluster was chosen as the antigenic reference strain. Overall, δ -clade viruses demonstrated limited cross-reactivity within each clade (H1- δ 1/1B.2.2 or H1- δ 2/1B.2.1), yet showed a trend to be mapped in accordance with their evolution (Figs. 2A and S3). The H1- δ 1 clade (1B.2.2) was the most antigenically diverse group, and the newly emerging H1- δ 1a (1B.2.2.1) and H1- δ 1b (1B.2.2.2) clades showed reduced cross-reactivity with older H1- δ 1 (1B.2.2) and between each other (average within clade distance of 2.9–3 AU, and between clades of 3.2 AU). Strains of the δ -clades collected after 2012 were more antigenically diverse, with antigenic distance from the representative strain ranging from 0.4 to 4.6, and most outliers were identified among these (Fig. 2A, C–E). Classical swine-lineage H1 viruses as well as H1N1pdm09 (1A.3.3.2) were clustered antigenically according to their genetic clades, with relatively stronger cross-reactivity within a clade (average within clade distance of 1–2.4 AU). The classical swine lineage and pandemic strains formed a separate antigenic group from the H1- δ viruses (Fig. S3).

4.3. Amino acid substitutions related to antigenic diversity among H1 viruses that circulate in North American pigs

We aligned the deduced HA1 amino acid sequences of the viruses used for the antigenic cartography to investigate the genetic basis for antigenic cluster differentiation. Genetic sequences from all H1 clades were compared (Fig. S4). The H1 numbering of the mature peptide was used throughout, unless indicated otherwise (Burke and Smith, 2014). The δ -clades (1B.2) maintained the deletion of one amino acid at position 130 compared to the classical-swine (1A.1, 1A.2, 1A.3.3.3) and the H1N1pdm09 (1A.3.3.2) clades as described in the first reports of these viruses in U.S. swine (Vincent et al., 2009). The classical-swine viruses showed 14 amino acid differences from the δ -viruses that fit the clade-defining criterion (L31, I57V, T72S, T133S, A135S, V152T, K160N, T184N, A195E, D196N, A224E, T241I, T245N, V249I; Fig. S4);

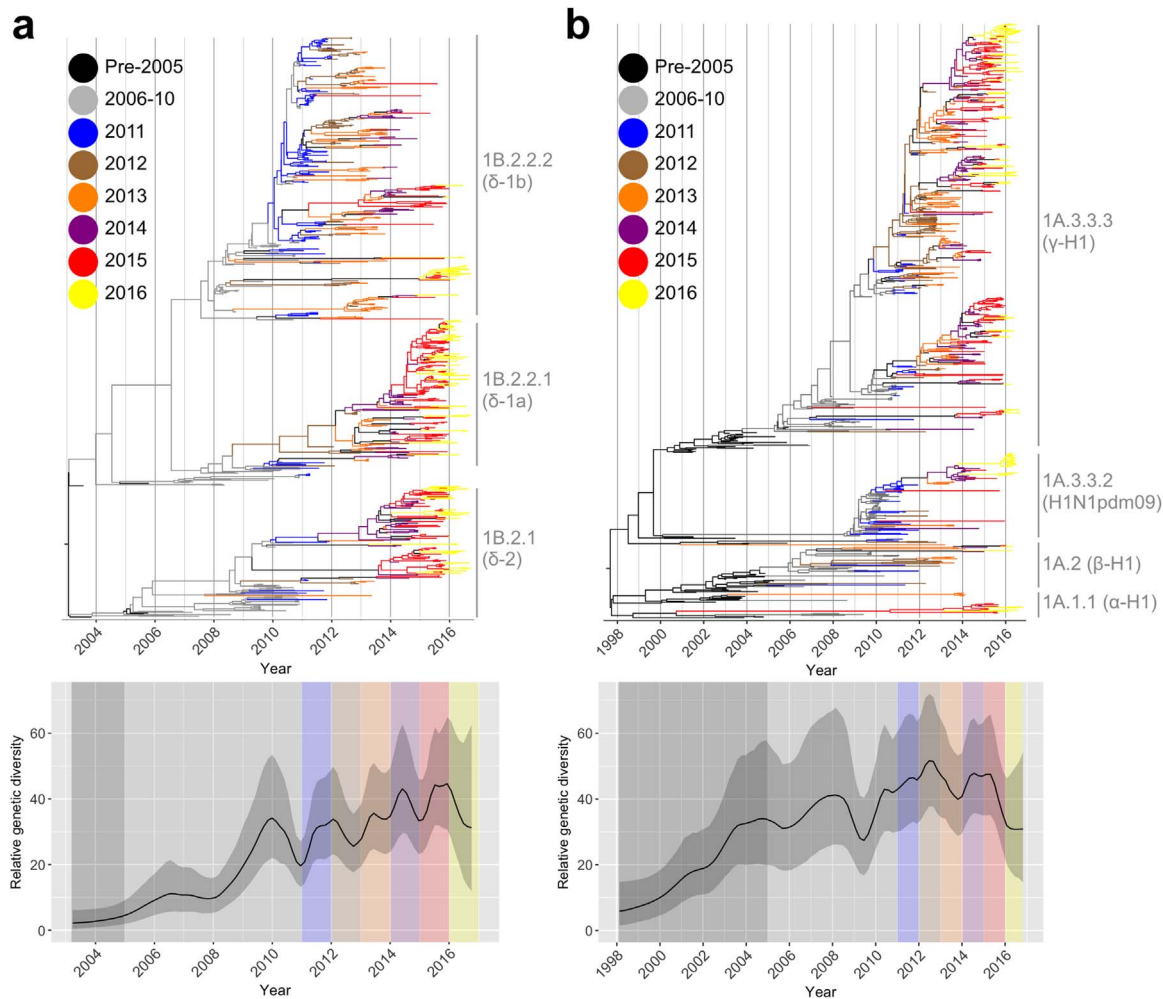


Fig. 1. Temporal phylogenies and epidemic patterns of swine influenza A H1 viruses. Evolution of the HA genes of (a) 730 H1- δ 1 and H1- δ 2 lineage and, (b) 726 classical swine H1N1 viruses from a randomly sampled subset of the 3211 HA genes in this study plus the sera and antigen reference strains. Phylogenies were inferred using the uncorrelated relaxed lognormal clock model with branches colored by year of virus isolation. Relative genetic diversity was estimated using the Gaussian Markov Random Field (GMRF) model: solid black lines in the GMRF plot represent mean relative genetic diversity, with the gray shading indicating the 95% HPD intervals.

a substitution was clade-defining if all, or all but one, viruses in one genetic clade had the same amino acid at a specific position in comparison to all, or all but one, viruses from a different genetic clade. The amino acids found in these 14 clade-defining positions were the same between δ -viruses and the tested human-seasonal H1 strains (Fig. S3). Six amino acid substitutions were identified as clade-defining between the H1- δ 1 (1B.2.2) and the H1- δ 2 (1B.2.1) viruses (A89T, I175V, T190A, W252R, D276N, I321T; Fig. S4), all of which were conserved between the swine H1- δ 1 (1B.2.2) viruses and A/Michigan/2/2003, a human seasonal virus that is genetically similar to the swine δ -clade and circulated around the time the deltas emerged in pigs (Nelson et al., 2014). Differentiation between H1- δ 1a (1B.2.2.1) and H1- δ 1b (1B.2.2.2) viruses was associated with four amino acid differences (E74K, S85P, D86E, G186E; Fig. S4).

We then analyzed the HA1 alignments to identify amino acid substitutions that could be involved with antigenic distance within an antigenic cluster. We focused this analysis on the H1- δ clade (1B.2.1 and 1B.2.2) because these viruses demonstrated a distinct genetic expansion and large antigenic diversity within clusters (Fig. 1A). Therefore, sequences of viruses in each H1- δ genetic clade were compared to the representative virus for each antigenic cluster separately to identify amino acid positions associated with the antigenic differences observed here (Fig. S5). Fourteen strains, most of which were strains detected after 2012, were mapped at a significant antigenic distance (> 2 AU) when compared to their respective antigenic cluster representatives

(arrows in Fig. 2A), and referred to as “outliers” (Fig. 2C–E). These strains showed at least 21 unique amino acid substitutions in positions in or near the receptor binding site (RBS) and/or known antigenic sites of H1 (Caton et al., 1982; Gamblin et al., 2004) in comparison to the antigenic representative for their respective clusters (Figs. 3 and S5). Positions that were observed in more than one non-outlier virus in the cluster were not considered, only positions unique to the outliers.

The three outliers for the H1- δ 1a (1B.2.2.1) cluster (A/swine/Nebraska/A01492366/2014, A/swine/South Dakota/A01481702/2014 and A/swine/Iowa/A01476931/2014) were located more than 3 AU away from the H1- δ 1a (1B.2.2.1) representative (Fig. 2C) and had similar amino acid changes among one another (T82A, K119R, E140K, R141G, P182S, V201M/T, G207R, A243T, Q282K, S288N), including an R141G mutation (Figs. 3 and S5). A mutation at position 141 was also present in A/swine/South Dakota/A01365260/2013 (R141K; 2 AU away from the cluster representative). An E140K mutation found in all three outliers was also found in A/swine/Minnesota/A01392045/2013 (1.6 AU away from the cluster representative), suggesting that positions 140 and 141 may have an additive effect on the antigenic differences observed for this cluster (Fig. 3).

All outliers of the H1- δ 1b (1B.2.2.2) cluster (A/swine/Minnesota/A01301731/2012, A/swine/Indiana/A01260356/2013, A/swine/Iowa/A01432370/2014, A/swine/Nebraska/A01290601/2013, A/swine/Illinois/A01410405/2014, and A/swine/Iowa/A01410774/2014) were positioned 3–4.6 AU away from the cluster representative

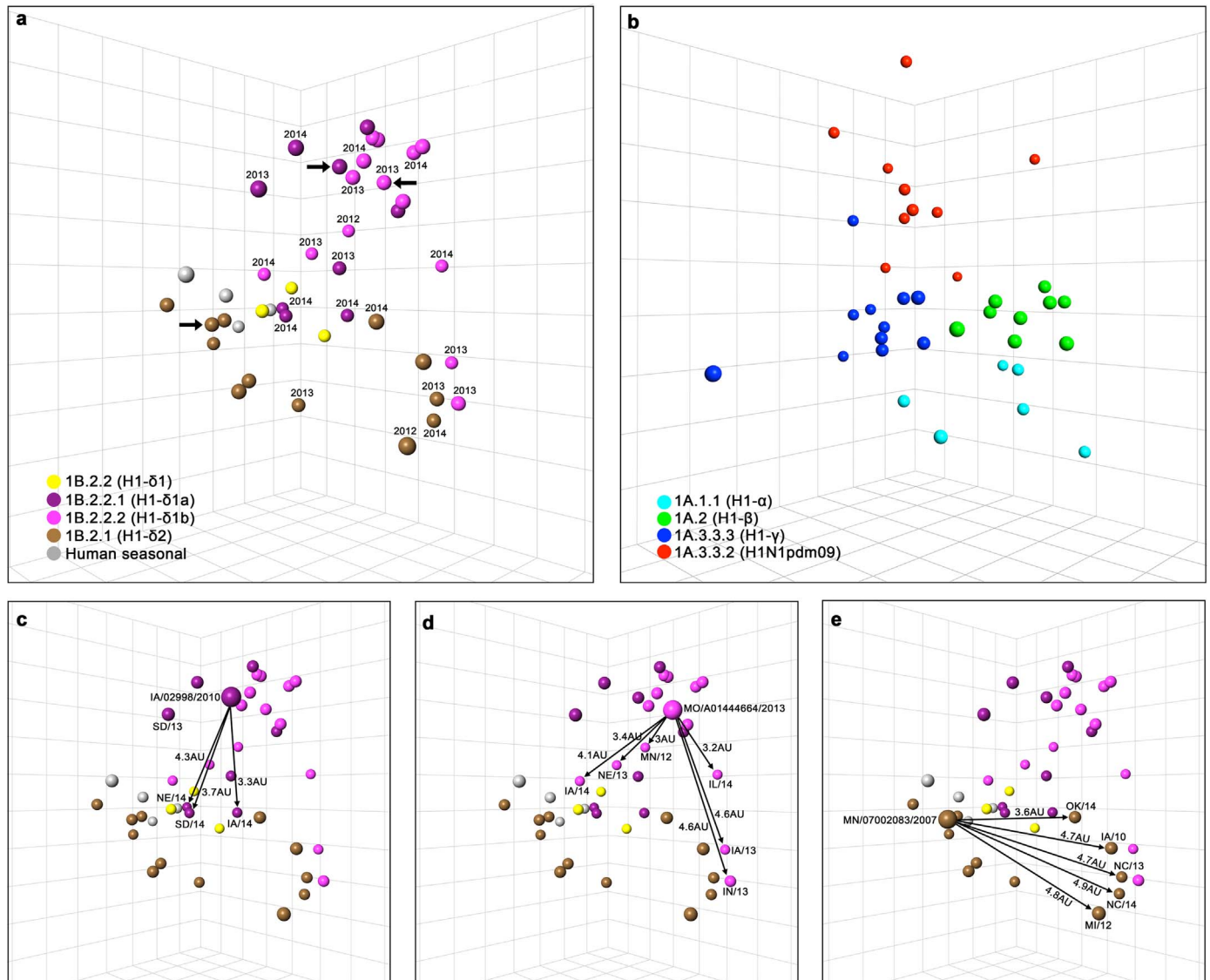


Fig. 2. 3D antigenic maps of representative historical and contemporary swine H1 influenza viruses of the 1B.2 delta (a) and 1A classical (b) lineages. Antigenic distances of outliers in comparison to antigenic cluster representative strain demonstrated for 1B.2.2.1 H1- δ 1a (c), 1B.2.2.2 H1- δ 1b (d), and 1B.2.1 H1- δ 2 (e). Isolates are shown as spheres colored by genetic clades. Each grid square represents 1 antigenic unit, corresponding to a twofold difference in HI assay titer. Year of detection is shown for contemporary delta lineage strains detected after 2012. Large arrows (a) or bigger spheres (c–e) indicate antigenic cluster representative strains A/swine/Iowa/02998/2010 (H1- δ 1a), A/swine/Missouri/A01444664/2013 (H1- δ 1b), A/swine/Minnesota/07002083/2007 (H1- δ 2). Arrows demonstrate antigenic distance from the representative strain (c–e). Outlier strains are H1- δ 1a: A/swine/Nebraska/A01492366/2014 (NE/14), A/swine/South Dakota/A01481702/2014 (SD/14), and A/swine/Iowa/A01476931/2014 (IA/14); H1- δ 1b: A/swine/Illinois/A01410405/2014 (IL/14), A/swine/Nebraska/A01290601/2013 (NE/13), A/swine/Iowa/A01410774/2014 (IA/14), A/swine/Iowa/A01432370/2013 (IA/13), A/swine/Indiana/A01260356/2013 (IN/13), and A/swine/Minnesota/A01301731/2012 (MN/12); H1- δ 2: A/swine/Oklahoma/A01409770/2014 (OK/14), A/swine/Iowa/02955/2010 (IA/10), A/swine/North Carolina/A01290598/2013 (NC/13), A/swine/Michigan/A01259001/2012 (MI/12), and A/swine/North Carolina/A01475606/2014 (NC/14).

(Fig. 2D). Although these viruses showed varied numbers of amino acid differences from the representative, a common difference among them was at position 140. Positions 131 and 141 were different from the representative for at least half of these outliers (Figs. 3 and S5).

The H1- δ 2 (1B.2.1) outlier strains (A/swine/Iowa/02955/2010, A/swine/Michigan/A01259001/2012, A/swine/North Carolina/A01290598/2013, A/swine/North Carolina/A01475606/2014, and A/swine/Oklahoma/A01409770/2014) were located 3.6–4.9 AU away from the cluster representative (Fig. 2E) and differed in 10 common amino acid positions. Seven of these 10 amino acid differences (except 140, 185, and 214) were also observed in A/swine/Ohio/A01351184/2013, a virus positioned at only 2 AU away from the cluster representative (Figs. 3 and S5), suggesting that the three other positions are likely the most significant. The A/swine/Indiana/A01260356/2013 H1- δ 1b outlier was positioned only 0.6 AU away from two outliers of the H1- δ 2 clade (NC13 and NC14); however these viruses showed less

than 90% HA1 amino acid homology, and no particular amino acid similarity was observed that could affect antigenicity (Fig. S4).

To further refine antigenically-important positions, pairs of viruses with amino acid distances less than or equal to 4% and antigenic distances greater than 3 AU were identified and specific amino acid differences between these pairs were compared (Fig. 4). This criteria identified 35 pairwise comparisons between 25 viruses (Fig. 4AB). Although there were outliers fitting this criteria in all 3 δ -clades, most pairs were H1- δ 1b (1B.2.2.2) clade viruses. The most frequently detected differences among the pairs of outliers were at position 140. Positions that occurred in more than ten of the pairwise comparisons included mutations at positions 131 ($n = 12$), 140 ($n = 30$), 141 ($n = 17$), 273 ($n = 13$), or 309 ($n = 12$) (Fig. 4C).

H1- δ 1a		82	119	140	141	182	201	207	243	273	282	288
*A/swine/lowa/02998/2010	AU	T	K	E	R	P	V	G	A	K	Q	S
A/swine/lowa/02039/2008	0.6	E	.	.
A/swine/Minnesota/A01134353/2011	1.0	E	.	.
A/swine/Missouri/A01411322/2014	1.1	G	.	.
A/swine/Minnesota/A01392045/2013	1.6	A	R	K	.	S	M	R	T	.	K	N
A/swine/South Dakota/A01365260/2013	2.0	A	R	.	K	S	M	R	T	E	K	.
A/swine/lowa/A01476931/2014	3.3	A	R	K	G	S	M	R	T	E	K	N
A/swine/Nebraska/A01492366/2014	3.7	A	R	K	G	S	T	R	T	.	K	N
A/swine/South Dakota/A01481702/2014	4.3	A	R	K	G	S	M	R	T	E	K	N

H1- δ 1b		68	94	96	127	128	131	137	138	140	141	142	145	148	165	167	169	175	183	186	193	197	219	223	251	257	260	272	273	275	276	309	314				
*A/swine/Missouri/A01444664/2013	AU	E	E	A	T	A	V	H	N	E	R	S	R	I	E	D	E	L	N	N	T	Y	V	E	W	S	L	D	E	D	S	A	V				
A/swine/Illinois/3120/2010	0.4	.	.	.	N	V	M	.	.	G	K	E	G	.	.	.	K	H		
A/swine/South Dakota/A01349341/2013	0.5	K	
A/swine/Indiana/A01260471/2014	1.0	K	
A/swine/Minnesota/A01366540/2014	1.0	.	D	I	
A/swine/Illinois/A01047020/2010	1.2
A/swine/Minnesota/03002/2010	1.2
A/swine/Minnesota/3128/2010	1.3	V
A/swine/Minnesota/A01301731/2012	2.9	E	.	.	.	K	K	
A/swine/Illinois/A01410405/2014	3.3	D	.	.	.	T	G	N	.	G	N	.	V	K	
A/swine/Nebraska/A01290601/2013	3.4	K	K	R	D	.	I	G	.	.	.	P	G	K	T	A	
A/swine/lowa/A01410774/2014	4.1	.	D	V	K	K	S	R	T	K	
A/swine/Indiana/A01260356/2013	4.6	K	K	T
A/swine/lowa/A01432370/2013	4.6	.	K	.	N	V	K	.	.	K	S	.	K	.	K	E	G	N	

H1- δ 2		69	119	121	127	129	133	140	152	167	174	185	188	208	214	221	225	258	269	272	288	307	309	
*A/swine/Minnesota/07002083/2007	AU	L	K	S	T	T	A	K	G	N	V	G	R	R	V	K	D	A	R	A	D	S	R	A
A/swine/IL/00685/2005	0.5	A
A/swine/Missouri/03013/2010	0.5	D	G	K	
A/swine/Ohio/003295/2010	1.3	D	K	
A/swine/Minnesota/A01076626/2010	1.6	D	K	
A/swine/NC/00573/2005	1.7	A	
A/swine/Ohio/A01351184/2013	2.0	S	.	V	K	N	E	K	.	G	.	.	.	
A/swine/Oklahoma/A01409770/2014	3.6	.	G	.	.	S	N	V	K	.	E	.	K	A	R	N	E	K	.	G	N	.	.	
A/swine/lowa/02955/2010	4.7	.	.	.	N	S	.	E	V	K	.	R	.	.	A	.	N	E	K	.	G	.	T	
A/swine/North Carolina/A01290598/2013	4.7	R	.	.	.	S	N	V	K	.	E	.	.	.	A	.	N	E	K	T	G	N	K	
A/swine/Michigan/A01259001/2012	4.8	.	E	.	.	S	Q	V	K	I	.	R	K	.	A	.	N	E	K	.	G	.	K	
A/swine/North Carolina/A01475606/2014	4.9	S	N	M	K	.	E	.	.	.	K	A	R	N	E	K	.	G	N	

Fig. 3. Amino acid differences identified between viruses tested in this manuscript and each respective cluster representative (*). Only amino acid positions in which outliers showed differences compared to the cluster representatives and that were located in or near the receptor binding site or antigenic sites of influenza hemagglutinin (HA) are shown, with some exceptions. Gray-scale indicates increasing antigenic distance from the representative measured as antigenic units (AU). Boxes indicate amino acid positions that were different from the cluster representatives for all outliers and that were unique to the outliers.

5. Discussion

Following the implementation of the U.S. national swine IAV surveillance system by the U.S. Department of Agriculture in 2009, the quantity of HA and NA viral sequence data has increased substantially, revealing a diversity of viruses co-circulating in U.S. pigs. Notably, a dramatic evolution has been observed for swine IAV in the past decade (Anderson et al., 2013; Gao et al., 2017; Rajao et al., 2017). Here, we characterized the genetic and antigenic diversity of currently circulating swine H1 influenza viruses in the U.S. While the classical-swine H1 viruses maintained the relative antigenic clustering characteristics that were seen for viruses circulating prior to 2009 (Lorusso et al., 2011), the δ -viruses (1B.2.1 and 1B.2.2) became more genetically and antigenically diverse. Our analysis revealed that two new genetic sub-clades emerged from the H1- δ 1 clade around 2007, H1- δ 1a (1B.2.2.1) and H1- δ 1b (1B.2.2.2), sustained transmission in pigs, and had expanded antigenic diversity.

Our phylogenetic analysis demonstrated considerable genetic diversity among H1 viruses circulating in U.S. pigs, with co-circulation of both major lineages and 7 distinct genetic clades of viruses (Anderson et al., 2013; Lorusso et al., 2011). The classical-swine H1 clades circulated consistently over the past decade, with records of H1- α (1A.1), H1- β (1A.2), H1N1pdm09 (1A.3.3.2), and H1- γ (1A.3.3.3) clade viruses detected each year (Lorusso et al., 2013). The anomaly within the classical-swine lineage is the appearance and rapid decline of the genetically and antigenically distinct clade of H1- γ 2 (1A.3.2) viruses (Anderson et al., 2015; Gao et al., 2017). Coalescent reconstruction of the population demographics of the classical-swine lineage (Fig. 1B) revealed an increase in relative genetic diversity from the early 2000s to 2010, followed by a period of more stable relative genetic diversity. This increase in diversity corresponds to the emergence and reassortment of the late 1990s novel triple-reassortant H3N2 virus with the

endemic classical-swine H1 viruses. The δ -clade viruses (1B.2.1 and 1B.2.2), however, showed a different evolutionary trajectory with a distinct increase in genetic diversity from approximately 2008 to 2016 (Fig. 1A). Our coalescent reconstruction showed an almost linear increase in relative genetic diversity, with the more recent years demonstrating oscillating patterns that correspond to the seasonal dynamics of IAV in swine (Anderson et al., 2013). This genetic diversity resulted in two new monophyletic clades descended from the H1- δ 1 (1B.2.2) clade and indicated the need for new clade designations (1B.2.2.1 and 1B.2.2.2) following criteria defined previously (Anderson et al., 2016). Antigenic characterization of representative viruses from the new clades was warranted based on the genetic distance between the two new clades. A similar pattern of increased genetic and antigenic expansion was observed for North American swine H3 viruses, in which six distinct clades evolved from the previously stable Cluster-IV (Kitikoon et al., 2013).

The molecular basis for the genetic diversity of swine H1- δ 1 (1B.2.2) viruses is a result of unique human-to-swine transmission events and within-host evolution. Amino acid positions that differed between classical-swine and H1- δ remained conserved with human-seasonal reference strains. This suggests that the HA of δ -clade viruses maintains a human-seasonal genetic signature even after more than 10 years of circulation in U.S. swine herds, despite swine H1- δ viruses having higher evolutionary rates than closely related human-seasonal viruses (Nelson et al., 2014). The combination of such human-seasonal-like characteristics, particularly at or near receptor-binding sites, and waning population immunity associated with the replacement of seasonal H1N1 human strains by the H1N1pdm09 (Pica et al., 2012) may allow novel reassortant viruses and antigenically drifted strains from the 1B genetic lineage and subsequent sub-clades to spill back to human populations, underscoring the importance of monitoring the evolution of circulating swine IAV for human pandemic preparedness.

in relative genetic diversity, we focused on testing contemporary δ -clade viruses in HI assays and analysis by antigenic cartography to evaluate potential concurrent antigenic evolution. Contemporary viruses of the H1- δ 1 (H1- δ 1a/1B.2.2.1 and H1- δ 1b/1B.2.2.2) and H1- δ 2 (1B.2.1) clades demonstrated significant antigenic distance from previously tested viruses (Lorusso et al., 2011). Antigenic drift of IAV is frequently associated with amino acid substitutions in antigenic sites on the globular head of the HA (Koel et al., 2013). Only 6–7 amino acid changes in the HA1 domain were shown to be responsible for marked antigenic differences in human and swine H3 viruses (Koel et al., 2013; Lewis et al., 2014), however H1 viruses seem to behave differently and the patterns seen for H3 viruses do not seem to be replicated for human H1 viruses (Koel et al., 2015). Here, several amino acid differences were observed between same-clade swine H1 viruses, some of which might have an antigenic effect. Positions 131, 140, 141, 273, and 309 were identified when comparing pairs of viruses with highly similar amino acid sequences (< 4%) but substantial antigenic distance (> 3 AU). With the exception of position 309, all of the remaining positions were located at or near antigenic sites or the receptor-binding site. Mutations at position 140 were present in all viruses considered outliers. Additionally, changes at positions 131 appeared in combination with position 140 in a limited number of viruses with significant antigenic distance, while position 141 appeared in combination with 140 in more than half outliers of H1- δ 1a and H1- δ 1b. Position 141 is the corresponding H1 position to position 145 in swine H3N2 viruses that was associated with significant antigenic change (Abente et al., 2016; Lewis et al., 2014). Positions 152 and 185 were associated with antigenic distance among H1- δ 2 viruses, which corresponds to positions 156 and 189 shown to have antigenic impact in swine H3N2 viruses. Although the swine H1- δ viruses are related to seasonal H1 viruses that circulated in humans prior to 2009 and maintain a relatively high degree of amino acid similarity, evolution in North American pigs led to the appearance of viruses that are now antigenically distinct from ancestral precursors (Fig. 2A).

The genetic evolution and consequent antigenic diversity among contemporary swine H1 viruses complicates the control of influenza infection in North American pigs through the use of vaccines. Current commercially available vaccines used against IAV in the U.S. swine industry are not frequently updated and, therefore, tend not to reflect the true genetic and antigenic diversity of contemporary circulating swine viruses (Rajao et al., 2014; Vincent et al., 2008b). Vaccine strain selection is a crucial process to obtain adequate and broad cross-protection against IAV infection in pigs, and antigenic cross-reactivity should be analyzed when doing so, since sequence similarity was not highly predictive of antigenic distance in many pairs of viruses (Fig. 4A). Thus, antigenic evaluation should be taken under consideration in conjunction with phylogenetic analysis for choosing efficacious vaccine strains. Nevertheless, faster and easier regulatory procedures are necessary to allow rapid modification of vaccines to keep up with the rapid evolution of circulating swine IAV lineages and possible emergence of reassorted strains.

In addition to swine being an important host in the general epidemiology and evolution of IAV, influenza illness also represents a significant economic burden to the swine industry. The current genetic evolution of the virus in swine populations and the resulting antigenic diversity contributes to vaccine failure, and these drifted viruses represent a potential zoonotic risk. We demonstrated that swine H1 influenza viruses underwent substantial antigenic diversification between 2010 and 2014, likely as a result of a few key amino acid changes. As genetic expansion was observed to continue from 2014 to 2016, these H1- δ viruses likely continued to drift, with obvious consequences for immune protection and vaccination. Characterizing the genetic evolution of swine IAV and understanding the implications to antigenic diversity and cross-protection is crucial to improve control measures that could lessen the consequences that this disease brings to pork production and public health.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found in the online version at <http://dx.doi.org/10.1016/j.virol.2018.02.006>.

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