Global migration of influenza A viruses in swine

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The complex and unresolved evolutionary origins of the 2009 H1N1 influenza pandemic exposed major gaps in our knowledge of the global spatial ecology and evolution of influenza A viruses in swine (swIAVs). Here we undertake an expansive phylogenetic analysis of swIAV sequence data and demonstrate that the global live swine trade strongly predicts the spatial dissemination of swIAVs, with Europe and North America acting as sources of viruses in Asian countries. In contrast, China has the world’s largest swine population but is not a major exporter of live swine, and is not an important source of swIAVs in neighbouring Asian countries or globally. A meta-population simulation model incorporating trade data predicts that the global ecology of swIAVs is more complex than previously thought, and the United States and China’s large swine populations are unlikely to be representative of swIAV diversity in their respective geographic regions, requiring independent surveillance efforts throughout Latin America and Asia.

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In early 2009, a novel reassortant H1N1 influenza A virus with gene segments from two swine virus (swIAV) lineages emerged in humans, initiating the first influenza pandemic of the 21st century. The virus had a complex genetic composition that had not been previously detected in swine, with six genome segments of North American triple reassortant swine virus origin (PB2, PB1, PA, HA (H1), NP and NS) and two genome segments of Eurasian avian-like swine virus origin (NA (N1) and MP). Evolutionary analysis of this novel North American/Eurasian reassortant virus indicated that these segments had circulated undetected in swine for at least 8 years (ref. 2). The first human outbreak of the pandemic H1N1 virus (pH1N1) occurred in Mexico, and the extent of viral genetic diversity observed in Mexico supports the hypothesis that the virus first emerged there in humans. However, efforts to detect the last common ancestor of the pH1N1 virus in Mexican swine populations have not been successful to date, and the opaque evolutionary history of the pandemic virus in swine highlights the gaps in our understanding of swIAV dynamics at a global scale.

In general, influenza viruses in swine are spatially separated into distinct North American and European swIAV lineages, although viruses of North American and European origin both circulate in Asia. Multiple viral lineages co-circulate in North American swine, including (i) ‘classical’ swine viruses, which descend from the 1918 H1N1 pandemic; (ii) ‘triple reassortant’ swine viruses, which emerged in the mid-1990s with a combination of human, swine and avian segments; and (iii) ‘delta’ (δ) viruses that are closely related to human seasonal H1 viruses from the early 2000s (refs 6,7). The main European swIAV lineages include ‘avian-like’ H1N1 viruses that jumped from birds to swine in the 1970s, human-origin H1N1 viruses from the 1980s and human-origin H3N2 viruses that are antigenically described as A/Port Chalmers/1/1973-like.

Multiple North American and European-origin swIAV lineages have both been identified in Asian countries. Due to high levels of co-infection, segmental reassortment occurs frequently in swine, such that they are an important reservoir host for influenza virus genetic diversity.

Live transport is routine in swine farming, and in the United States the transport of millions of swine from Southern to Midwestern regions for end-stage production appears to drive the strongly directional dissemination of swIAVs from Southern US states with high hog production (for example, North Carolina, Texas and Oklahoma) to the traditional centre of swine farming located in the Midwestern ‘corn belt’. Large numbers of swine also enter the United States from Canada, which has been implicated in the dissemination of other important swine pathogens, including porcine reproductive and respiratory syndrome virus.

Intercontinental trade of live swine also occurs, for end-stage production or to acquire female breeding pigs for genetic improvement of swine reproduction or growth traits. Globally, the largest swine population is found in China, where over 450 million hogs reside (Fig. 1). Large swine populations also are found in the United States (>60 million hogs), Brazil (>30 million hogs), Vietnam (>20 million hogs), Germany (>20 million hogs) and Spain (>20 million hogs), among others.

Despite the global nature of both swine farming and swIAV circulation, the patterns and dynamics of the worldwide spread of this economically important virus are unknown. To characterize the phylogeography of swIAVs at a global scale, here we conduct a phylogenetic analysis of 785 whole-genome swIAV sequences collected from 10 countries/regions representing 4 continents, the largest study of its kind undertaken to date. To assess the drivers of viral migration, we compare the phylogeographic patterns with empirical data on live swine trade and swine population sizes. On

**Figure 1 | Modelled global swine distributions.** Digital layers from Gridded Livestock of the World (GLW; version 2.01; ref. 50), downloaded from the publically available Livestock Geo-Wiki database (http://www.livestock.geo-wiki.org) and manually edited in QGIS v.1.7.0. Swine densities are represented by the black shading.
the basis of these findings, we build a meta-population model to simulate the spatial dissemination of swIAVs at a global scale and to identify the regions that are at a high risk for co-invasion of divergent lineages, increased total genetic diversity and emergence of viruses with pandemic potential.

Results

Global migration of swIAVs. Phylogenetic analysis revealed that long-distance movement of influenza A viruses between countries and continents has occurred continuously in swine since the 1970s (summarized in Fig. 2). Our estimate of 18 international viral migration events is a minimum based on the currently available swIAV sequence data and certainly underestimates the true number. This lower-bound estimate is based on discrete monophyletic groups (defined by country) that are supported by high posterior probabilities (> 80), reflecting migration events that led to successful onward transmission. The estimate does not include the much higher number of international viral movements between the United States and Canada or between countries in Europe, which are each considered as meta-populations in our analysis. The estimate also does not consider viral migration events for which only one sequence is available, any viruses that do not form a well-supported clade, or which only partial sequence data were available.

Although global surveillance and sequencing of swIAVs has increased markedly in the last 5 years, our timescaled maximum clade credibility (MCC) phylogenies indicate that most intercontinental viral migration events occurred before this increase in surveillance (representative phylogeny of the NA segment is presented in Fig. 3; phylogenies for other seven segments are available in Supplementary Figs 1–7). Eight of the 18 viral migration events identified in our study were evident on the phylogenies inferred for all eight viral genome segments, indicative of the onward transmission of the full viral genome in the new location (introductions 1, 3, 4, 5, 8, 9, 10 and 15). The consistency of spatial–temporal inferences across these eight segments strengthens inferences of when and where each of these introductions occurred (Supplementary Table 1). Ten viral migration events could only be identified by a subset of genome segments, as at least one segment has been replaced in intervening years by reassortment (introductions 2, 6, 7, 11, 12, 13, 14, 16, 17 and 18). There is no evidence in these data that either of the δ-1 or δ-2 virus lineages that emerged in North American swine in the early 2000s has transmitted to swine on any other continents, despite the high rates of detection in the US swine populations of δ-1 viruses.

A consistent global spatial dynamic was observed for swIAVs during 1970–2013, based on both a conservative measure of the strongly supported monophyletic groups (Figs 2 and 3 and Supplementary Figs. 1–7) as well as ‘Markov jump’ counts of the expected number of location state transitions along the branches of the tree. ‘Markov jump’ counts provide a quantitative measure of gene flow between the regions that includes singletons and clusters that may have less phylogenetic resolution (Fig. 4, Supplementary Table 2). Overall, North America (in this case
Figure 3 | Maximum clade credibility (MCC) trees of the NA lineages in swine. Timescaled Bayesian MCC trees inferred for the NA segment for the three major swine virus lineages: (a) avian-origin Eurasian N1 swIAV lineage, the (b) classical N1 swIAV lineage and the (c) multiple human seasonal virus-origin N2 swIAV lineages circulating in swine. Branches of human seasonal H3N2 influenza virus origin are shaded grey in c, while branches associated with viruses from swine are shaded by country of origin: Argentina = brown; Canada = red; China (including Hong Kong SAR and Taiwan) = yellow; Europe = black; Japan = pink; Mexico = light blue; South Korea = green; Thailand = orange; USA = dark blue; Vietnam = purple. Posterior probabilities > 0.8 are included for key nodes, and international migration events that are supported by high posterior probabilities and long branch lengths are labelled according to Fig. 2.

referring to the United States and Canada) and Europe represent independent viral source populations for the Asian countries sampled in our study: China, Japan, South Korea, Thailand and Vietnam. In contrast, only a single swIAV migration event was observed between North America and Europe (introduction 4).

Spatial dynamics of swIAVs in North America. Bidirectional viral migration between Canada and the United States is so frequent (reflected by the extremely high number of Markov jump counts, Fig. 4) that Canada and the United States were considered as a single meta-population, similar to Europe (Fig. 2). The higher availability of swIAV sequence data from US swine makes it particularly difficult to distinguish whether an introduction was specifically of the US or Canadian swine origin, and the origin of such introductions is more appropriately characterized as ‘North American’. Using newly generated sequence data from five swIAVs of the H3N2 subtype that were collected in Mexico during 2010–2011 (A/sw/Mexico/SIG1442/2010, A/sw/Mexico/SIG1444/2011, A/sw/Mexico/SIG1447/2011, A/sw/Mexico/SIG1448/2011 and A/sw/Mexico/SIG1449/2011, accession numbers available in Supplementary Table 3), our phylogenetic analysis provides evidence of a single introduction of a H3N2 triple reassortant swIAV from the United States into Mexico that occurred between mid-2005 and mid-2006 (introduction 11, Figs 2 and 3c). At the time of sampling, all five Mexican swIAVs had acquired at least one pH1N1 segment of the human origin via reassortment, evidence that pH1N1 viruses also have circulated in Mexico’s swine population.

Spatial dynamics of swIAVs in Asia. Given China’s large swine population and long-term surveillance, it may have been expected that this country (encompassing mainland China, Hong Kong, Taiwan and Macau) would be an important source of swIAV diversity for the neighbouring Asian countries in this study. However, since 1970 there have been 16 swIAV introductions of European or North American origin into Asia, compared with only two swIAV migration events between the Asian countries, and only one definitive introduction of a swIAV from China into another country (introduction 16, Fig. 2). Overall, the genetic diversity of swIAVs in Asia derives from five swIAV introductions of European origin and 11 swIAV introductions of North American origin. Six viral introductions from Europe and North America were observed in Thailand, and five introductions were observed in China, including the earliest intercontinental swIAV migration detected to date (introduction 1, Fig. 2).

In contrast to the frequent exchange of swIAVs across European country borders and across the US–Canadian border, only two swIAV migration events were observed between any two Asian countries. A pair of H1 segments collected in South Korea in 2013 are positioned within a clade of avian-origin Eurasian viruses from China, suggesting China-to-Korea migration (introduction 16, Fig. 2 and Supplementary Fig. 4). Closely related North American triple reassortant viruses also were identified in China and Vietnam, suggestive of viral migration between these Asian countries. However, the location state probability for the node representing the common ancestor of the
potential swIAV circulation. However, the relatively long branches may not reflect a single direct transition between the origin and (final) destination location, and may conceal additional spatial movements during the elapsed time. The evolutionary history of swIAVs in Thailand is also made more complex by the frequency of reassortment involving multiple clades, poorly supported clusters and singletons. Whereas avian-like Eurasian viruses in China are monophyletic and result from a single introduction from Europe (introduction 15), the Thai viruses from this lineage are monophyletic only in the PB2, NP and N1 trees. Our estimate of the two introductions of avian-like Eurasian viruses into Thailand is therefore conservative and likely underestimates the true number (introductions 13 and 14, Figs 2 and 3, Supplementary Figs 1–7). Singletons and unsupported clusters also were observed among South Korean viruses, complicating the estimation of the number of viral migration events into South Korea as well.

Importance of live swine trade in the global dispersal of swIAVs. We used a generalized linear model (GLM) extension of phylogeographic inference to identify the putative drivers of swIAV migration events inferred from the genetic data. This approach considers single introductions and clusters that may have poor resolution, the uncertainty of which is accommodated by the analysis. The Bayesian model averaging approach found consistent and strong evidence that the amount of asymmetric live swine trade (measured in USD for the years 1996–2012) from one country to another is the dominant driver of the dispersal of SW IAVs globally. This is reflected by the maximal estimated inclusion probability of live swine trade for all six internal gene segments (probability of 1, results for PB2, PB1, PA and NP are presented in Fig. 5; results for the shorter MP and NS segments are available in Supplementary Fig. 8; the analysis could not be performed on the HA or NA due to the high frequency of human viruses in these phylogenies). Accordingly, viral migration, measured by ‘Markov jump’ counts, was positively correlated with live swine trade volume (USD; $\rho = 0.52$, $P = 1.5 \times 10^{-7}$, Spearman’s correlation, Supplementary Fig. 9).

Other potential predictors, including swine population size, contributed little to the observed global spatial patterns, except for the number of sequences of the country of destination (average probability of $\sim 0.5$ across the six internal segments). Not only is the effect of live swine trade consistently robust to the inclusion of sample size, the high inclusion probability of live swine trade in cases where the effect of sample size is particularly low (for example, the PA segment) indicates that it is also independent of the sample size. The conditional effect size (the size of the effect conditional on the predictor being included in the model) ranges between $\sim 3$ and 5 on a log scale, implying that viral lineage movement probability is several orders of magnitude higher for connections with the highest swine trade compared with connections without trade (Fig. 5).

Our GLM analysis could be affected by regional differences in the early establishment of swIAVs, with swIAVs having been potentially seeded later in Asia and decreasing the probability of viral export from Asia. To further explore this hypothesis, we conducted a sensitivity analysis focused on two more recent periods that correspond approximately to the emergence in China of classical North American swIAVs (1990–2013) and avian-origin swIAVs from Europe (2000–2013), using an ‘epoch’ extension of the diffusion model. We find that the volume of live swine is still the only well-supported predictor of viral migration for both the periods (Supplementary Table 4), including when swIAVs are thought to be endemic at high levels in Asia as well as Europe and North America. The relatively

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**Figure 4 | Heat-map of swIAV migration between locations.** Countries are listed in order of increasing geographical distance from Argentina (ARG). MEX = Mexico, USA = the United States, CAN = Canada, EUR = Europe, JPN = Japan, CHN = China (including Hong Kong SAR and Taiwan), KOR = South Korea, THA = Thailand, VNM = Vietnam. The intensity of the colour (red = high; white = low) reflects the number (no.) of ‘Markov jump’ counts inferred over the totality of phylogenies (all segments, all lineages) from one location to another (asymmetrical). Markov jump counts measure the number of inferred location state transitions, modelled by a continuous-time Markov chain process, that occur along the branches of the phylogeny. For clarity the heat-map has been divided into four sections representing (a) viral migration events within the Americas and between the Americas and Europe; (b) migrations from the Americas/Europe to Asia; (c) migrations from Asia to the Americas/Europe; and (d) migrations between the Asian countries.

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Chinese and Vietnamese clades is too low (ranging from 0.46 to 0.65 across the eight genome segments) to determine whether the North American virus was first introduced to Vietnam and disseminated to China, or vice versa (Fig. 2 and Supplementary Figs. 1-7).

Although much of the swIAV diversity in Asia appears to have emerged in the last two decades, the phylogenies suggest long-term circulation of swIAVs in Thailand and Japan, either via imports from North American or European swine in the 1970s and 1980s (Fig. 2) or direct introductions from humans as early as the 1960s (Fig. 3c). Long branch lengths and the lack of historical swIAV data from Asian countries make it difficult to infer with confidence the spatial history of older viral lineages in Asia. The relative lack of swIAV surveillance in Thailand before 2000 particularly complicates the estimates of the timing and spatial pathway of the multiple viral introductions from North America and Europe that likely occurred during the 1980s and 1990s (for example, introductions 6, 14, 17 and 18). At this time, there is little evidence of viral dissemination from Japan to Thailand to other Asian countries in our study, despite many decades of
Inclusion probability (E[β])

In coefficient (E[β])

In coefficient (E[β])

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Inclusion probability (E[β])

Figure 5 | The support and contribution of swIAV diffusion predictors among nine countries. Twelve predictors were considered: geographical distance (km), volume of live swine trade, 1996–2012 (USD), swine population size for the years 1969–2010, the total number of imports of live swine during 1969–2010, the total number of swine exports during 1969–2010, the percent change in swine population (pop) size from 1969–2010 and the number of sequences available from a given country for our analysis. ‘O’ refers to the swine population of origin, and ‘d’ refers to the swine population of destination. Support for each predictor is represented by an inclusion probability that is estimated as the posterior expectation for the indicator variable associated with each predictor (E(β)). The contribution of each predictor is represented by the mean and credible intervals of the GLM coefficients (β) on a log scale conditional on the predictor being included in the model (β|δ = 1). See Supplementary Fig. 8 for MP and NS results.

Predicted spatial dissemination of swIAVs. To explore how the global network of live swine trade may drive movements of swIAVs beyond the 10 countries for which whole-genome sequence data were available, we used data on pairwise live swine trade between the 146 countries to simulate the patterns of viral dissemination under different epidemiological scenarios. Figure 6 explores the predicted spatiotemporal spread of a new swIAV lineage that hypothetically originates in swine in one of five countries with large swine populations: Canada, China, France, Mexico and the United States. These predictions are largely consistent with the spatial movements observed in the genetic data, with a high probability of viral export from the United States and Canada into Asia, and from Europe to Asia, whereas epidemics originating from China have low probabilities of onward dissemination to other countries (Supplementary Table 5). In addition, we identified predicted connections with countries not sampled in our study, including swIAV migration from the United States and Canada to many countries in Latin America, as well as Russia, Kazakhstan, Malaysia and Singapore. Interestingly, our model suggests that a virus seeded in Mexican swine is comparatively less likely to disseminate to swine in other countries, including the United States.

We also used our model to estimate the probability of co-invasion of European and North American swIAVs lineages,
illustrating the potential for reassortment between the lineages of European and North American descents, of the kind that generated the 2009 pH1N1 virus. Overall, co-invasion is strongly regionalized, with the highest probability in East and South-East Asian countries, particularly China, South Korea and Russia (Fig. 6, Supplementary Table 5). Conversely, South Asia, the Middle East, Africa and Australia exhibited a low probability of invasion by each of these lineages. Mid-level probabilities were found in regions with a high probability of invasion by only one of the two lineages (that is, the North American lineage in the Americas and the Eurasian lineage in Europe). Interestingly, these simulations reveal a low probability of co-invasion in Mexico, where pH1N1 first emerged in humans, owing to the low probability of invasion by a European swine virus in Latin America.

Discussion
The unknown origins of the swine virus that begot the 2009 H1N1 pandemic underscores the importance of understanding how influenza A viruses evolve in swine at a global scale, including regions where swIAV surveillance is lacking. Our expansive phylogenetic analysis of global swIAV sequence data demonstrates the importance of the asymmetrical nature of the global live swine trade on the global ecology and evolution of swIAVs. Using a phylogeographic GLM approach to assess the strength of specific predictors, we determine that the size of a country’s swine population is not a major factor in the rate of viral export to other countries. As a notable case in point, China, which hosts the world’s largest swine population, does not appear to be a major source of the viral diversity observed in other Asian countries (Fig. 4d). Rather, Japan, Thailand, Vietnam and South Korea independently imported novel viruses from Europe and North America (Fig. 4b), most likely via long-distance live swine trade.

The reported pattern of swIAV dissemination is a reverse of a model proposed for the global spread of A/H3N2 seasonal influenza viruses in humans, in which a highly connected network of South-East Asian countries, including China, acts as a key source of viruses for Europe, North America and other continents, reflecting differences in the disease and mobility patterns of humans and swine. These findings have important
implications for swIAV surveillance strategies, as the relatively low levels of viral gene flow between Asian countries means that no single country in Asia can serve as a proxy for the region, including China’s large swine herds. The extent of viral genetic diversity in Thailand highlights the importance of enhancing surveillance throughout South-East Asia, including countries not sampled in our study, such as Malaysia, Indonesia, Singapore, Laos and Cambodia, and undersampled countries such as Thailand, Vietnam, Japan and South Korea. Furthermore, Russia emerged as a hotspot for invasion and co-invasion of divergent lineages in our simulations, and yet Russia has no publicly available whole-genome swIAV sequences.

The limited number of sequences from Asian countries other than China (particularly via Hong Kong, the final destination of large numbers of hogs from mainland China) reduces our ability to detect viral migration events within Asia, particularly those that do not transmit onward in swine for many years. However, the high number of viruses identified in Asia that were of North American and European origin indicates that sample bias alone cannot explain the lack of viral exchange observed between the Asian countries. Analysis of larger, less-constrained data sets including all available HA swIAV sequences from Asia identified several additional viral migration events from Europe and North American swine into Asia, but only limited evidence for one additional putative connection between two Asian countries (Supplementary Figs. 10–11). However, all inferences of spatial connections must be interpreted within the context of the many countries that are unsampled and undersampled, and long branches may conceal additional spatial movements between the origin and (final) destination location.

It is important to note that our study focused only on the international dissemination of swIAVs, and did not consider the probability of initial emergence of an epidemic within a country, which is likely to be influenced by numerous local factors related to national swine farming practices, including the size and density of farms, movements of pigs within countries and the opportunities for interspecies transmission. As demonstrated previously, the dynamics of outbreaks within a large country like the United States can be complex, with different regions acting as source and sink populations for viral diversity17. Novel IAVs of human origin have emerged repeatedly in swine in countries in North America, South America, Asia and Europe, suggesting that swine populations in these regions can sustain new epidemics24. The extent of viral export from a country of origin is a product of both the national prevalence of circulating swIAVs and the volume of live swine export. In this study, we were unable to assess whether geographic differences in the prevalence of swIAVs affect large-scale viral migration, as population-level virological and serological data indicative of swIAV prevalence are available only from a limited number of study sites and time periods that are unlikely to be sufficiently representative for a global study. We therefore recognize that there are scenarios where live swine trade alone would not be a good predictor of viral migration. For example, if the major exporters of live swine (North America and Europe) did not have large endemic swIAV populations, then live swine trade alone would not be a good predictor of viral migration. This does not appear to be the case, as North America and Europe have long histories of endemic swIAV circulation and the highest volumes of outgoing international live swine trade. The apparent association between viral endemicity and trade export may not be a coincidence, as the features that enable countries in Europe and North America to export high volumes of live swine (that is, large-scale commercial swine production) also are likely to be conducive to sustained endemic swIAV transmission. Finally, our analysis did not consider swine influenza vaccine use, which is highly heterogeneous within and between countries, and would be an important, albeit challenging, factor to integrate into future studies that consider the interaction between the national prevalence of swIAVs, the structure of swine industries, and the global spatial dynamics of trade and viral migration.

Although it is not possible at this time to incorporate empirical data on historical differences in swIAV prevalence by region, we were able to explore the interaction of influenza virus prevalence and trade volume using our simulation model. Of particular interest was the question of whether the low levels of viral export from Asia observed in our study could be an artifact of the historically lower levels of endemic swIAV activity in Asia. As a consequence, viral exports from Asian countries might be expected to rise in the future as swIAVs become endemic at higher levels throughout the region. There was no support for this hypothesis, as our simulations predicted very low rates of viral export from China to other countries even when the prevalence of swIAVs in China is set unrealistically high (for example, 58% of the Chinese swine were infected when \( R_0 = 1.5 \)). Our simulations predicted low rates of swIAV export from Japan and Thailand under similar transmissibility scenarios (Supplementary Fig. 12), with the exception of viral dissemination from Thailand to Cambodia, as these countries are trade partners. These predictions are consistent with our observation from the genetic data that swIAVs could have circulated in Japan and Thailand for many decades without substantial onward dissemination to other Asian locations sampled in our study. In addition, a sensitivity analysis of the GLM model, limited to the more recent 1990–2013 and 2000–2013 periods where swIAVs were established in Asia, lends further support for the importance of trade (Supplementary Table 4). Overall, our findings suggest that viral exchange between Asian countries with low levels of trade is unlikely to increase in the future, regardless of the potential increases in endemic swIAV activity in the region as farming practices are modernized and swine farms become larger.

Despite the importance of swine trade in the global ecology of swIAVs, it should be noted that humans may be equally, if not more, important in disseminating IAV diversity to swine herds globally25. Even in the absence of international swine trade, swIAVs of human origin would likely still circulate in the majority of countries in our study, including in Asia. Quarantine and other restrictions in international trade may have the potential to reduce the genetic diversity of swIAVs, but are not likely to prevent swIAVs from circulating in a country’s swine population (Australia is a case in point26). The frequency of human-to-swine transmission has been even more apparent since the 2009 H1N1 pandemic, and humans have disseminated pH1N1 viruses to swine in numerous countries that had not previously reported IAV activity in swine, including Australia26,27, Brazil28,29, India30, Cameroon31, Mexico32, Nigeria33, Sri Lanka34 and several countries in Europe35–37.

Unfortunately, these new data did not advance our understanding of the evolution of the pH1N1 virus during the many years of undetected circulation in swine before 2009. Given that the human pandemic likely emerged in Mexico3, the most parsimonious explanation is that the pH1N1 virus transmitted from swine to humans in Mexico or a nearby locality. However, extremely little swIAV sequence data are available from swine in Mexico and other parts of Latin America, and Eurasian viruses have not been detected in any part of the Americas to date. Our simulation model provides a quantitative indicator of where the reassortment event that produced the pH1N1 virus in swine was most likely to have occurred, based on the probability of invasion with both North American and European viruses. Given limitations in our model, including the lack of information on within-country dynamics and the likelihood of initial viral
emergence within seed countries, we consider the relative ranking of probabilities to be more important than their absolute values. To date, Asia is the only region where any reassortant viruses containing both North American and Eurasian virus segments have been detected, consistent with our simulations, which show a high probability of co-invasion by both North American and Eurasian swine lineages in China, South-East Asia and Russia. However, it remains unclear how a reassortant virus that most likely emerged in swine in Asia caused its first outbreak in humans in Mexico. Given the lack of south-to-north swine trade flows in the Americas, some swIAV lineages are likely to be exclusive to Latin America39,40, and not reach the United States or Canada. Strengthened surveillance in Latin America is needed to gain a better understanding of swIAV diversity in the region.

Finally, we have focused our study on the global dynamics of influenza A viruses in swine, but our findings invite investigation of how trade, quarantine and swine farming practices affect the spatial dynamics of other globally dispersed swine pathogens, such as porcine reproductive and respiratory syndrome virus and the porcine epidemic diarrhea virus that emerged in the United States during swine flu outbreaks in 2013. Modelling studies rooted in population sequence information, demographics and mobility data have the power to inform global surveillance and control strategies for major animal and human disease threats.

Methods

Influenza virus sample preparation. Influenza A virus samples collected from swine via routine diagnostic submissions for the years 2002–2011 were randomly selected from the existing influenza virus archive at the University of Minnesota Veterinary Diagnostic Laboratory (UMVDL). These samples were chosen to best represent this time period and the three main geographical regions of the US hog production: the Southeast region (US states of Alabama, Georgia, Kentucky, North Carolina, Tennessee and Virginia), South-central/west region (Arkansas, Colorado, New Mexico, Oklahoma and Texas) and Midwest region (Iowa, Illinois, Indiana, Kansas, Minnesota, Missouri and Nebraska). Samples from swine in Canada (2005–2011) and Mexico (2010–2011) were also selected from UMVDL, as available. Original specimen material (nasal swab, supernatant or lung tissue homogenate stored at −80 °C) were aliquoted from the archived samples and sent to the J. Craig Venter Institute (JCVI) in Rockville, MD for sequencing.

Influenza virus genome sequencing. The complete genomes of 240 influenza viruses collected from North American swine were sequenced at JCVI. Viral RNA was isolated using the ZR 96 Viral RNA kit (Zymo Research Corporation, Irvine, CA, USA). The six viral genomic RNA segments were amplified from 3 μl of purified RNA using a multi-segment reverse transcription-PCR strategy (M-RTPCR)41. The M-RTPCR amplicons were sequenced using Nextera Library construction using the MiSeq platform (Illumina Inc., San Diego, CA, USA). In addition, M-RTPCR amplicons were sheared for 7 min and Ion Torrent compatible barcodes were added to the fragments. The 200 base pair libraries were purified and sequenced using Ion Torrent (Life Technologies, Grand Island, NY, USA). All data sequences for this study were submitted to the Influenza Virus Resource at the National Center for Biotechnology Information’s GenBank, and access codes are available in Supplementary Table 3.

Phylogenetic analysis. In addition to the sequences generated for this study, whole-genome sequences from influenza A viruses collected in swine globally during 2003–2013 were downloaded from the Influenza Virus Resource at GenBank42. Viruses were removed that (a) had truncated sequences, (b) were of avian origin, (c) were of North American and Eurasian swine lineage, and (d) were from this time period and the three main geographical regions of the US hog production: the Southeast region (US states of Alabama, Georgia, Kentucky, North Carolina, Tennessee and Virginia), South-central/west region (Arkansas, Colorado, New Mexico, Oklahoma and Texas) and Midwest region (Iowa, Illinois, Indiana, Kansas, Minnesota, Missouri and Nebraska). Samples from swine in Canada (2005–2011) and Mexico (2010–2011) were also selected from UMVDL, as available. Original specimen material (nasal swab, supernatant or lung tissue homogenate stored at −80 °C) were aliquoted from the archived samples and sent to the J. Craig Venter Institute (JCVI) in Rockville, MD for sequencing.

Testing predictors of global swIAV migration. A GLM32 parameterization of the discrete phylogeographic diffusion model was employed to estimate the contribution of potential predictors to the migration pattern of swine (a representative XML file used in the analysis is provided in Supplementary Data 3; the R code used to summarize the estimates is provided in Supplementary Data 4). First, the trade value (USD) for live swine trade between countries (asymmetry) for the years 1996–2012 was obtained from the United Nations’ Commodity Trade Statistics Database (available at http://comtrade.un.org, accessed March 20, 2014) (Supplementary Data 5). For the purposes of our study, we calculated the total trade value for each country. Data from all European countries were aggregated within the category ‘Europe’, and the data from mainland China, Macao SAR and Hong Kong SAR were aggregated within the category ‘China’. Second, estimates of trading partner credits were employed to determine the total country to country exports per country were obtained for a longer time period (1969–2010) from the Food and Agriculture Organization (FAO) of the United Nations Datasets.
Meta-population model simulating the global spread of swine influenza. Next, we built a meta-population model to simulate the global spread of swiAVs and identify the potential geographical hotspots for reassortment between viruses originating from different regions, which may generate novel viruses with pandemic potentials. We employed a stochastic patch-based SIR model adapted from an earlier model for the global diffusion of human influenza 47. Each patch represents the swine population of an individual country, and patches are linked based on the live swine trade movements. To calibrate the model, we obtained swine population sizes and pairwise between-country trade information for 146 countries reporting to FAO during 1969–2010 (http://data.fao.org/datasets; Supplementary Table 7), which coincides with the study period considered for phylogenetic analysis of swiAVs. We used the averages of swine population sizes and pairwise trade volumes throughout the study period for model simulations.

The influenza simulation model is as follows. Let S, I and R denote vectors representing the number of susceptible, infected and recovered swine at any time point in each of the 146 countries studied. We let μ = 1/5 denote the daily probability that an individual recovers (so that the infectious period is 5 days88,89), and β I the daily per capita rate of infection from infectious individuals within the same country. Here β varies between countries; it is a vector scaled such that the effective reproduction number ≤R 0 is the same across all countries, where R 0 is the vector of swine population sizes. We use an R 0 of 1.5 in main analyses, consistent with limited information on swine influenza dynamics.88,89 The per capita rate of contacts with infectious swine from other countries is given by G*I, where G is a 146×146 coupling matrix representing between-countries swine fluxes.

To build G, we fitted a 146×146 matrix with off-diagonal elements based on empirical live swine trade and zonal along the diagonal. We then rescale T by the estimated trade coefficients of the phylogeographic GLM model, as the GLM model suggests that the relationship between trade and viral migration is not linear (but results are qualitatively similar with no rescaling). Following the past work27, the resulting parameter T is tuned by a free parameter that governs the amount of international versus domestic contacts between swine, while at the same time allowing the conversion of empirical swine trade data (provided in $ amount by FAO) into actual population movements. Tuning parameter μ allows obtaining realistic time course of infection, with global epidemics lasting between several months to several years, in line with [limited data available on] the global spread of past swine outbreaks. Our final coupling matrix is G = GT, where c is such that the maximum element of the coupling matrix is <10⁻⁵.

We use a spatially extended chain-binomial system to update the progression of the epidemic in each patch27:

\[ S_{i,t+1} = S_i - W_i \]
\[ I_{i,t+1} = I_i + W_i - V_i \]
\[ R_{i,t+1} = R_i + V_i \]
\[ W_i = \text{binomial}(S_i, 1 - e^{-\beta I_i (1 + G^t I_j)}) \]
\[ V_i = \text{binomial}(I_i, \mu) \]

Here, W is the daily incidence of swine influenza (that is the number of new cases), and V is the number of new ‘recovereds’. In simulations, the epidemic is initialized by infecting five swines in a predetermined seed country; we explored various scenarios with seeds in countries of the Americas, Europe and Asia. After the first infection occurs in each country, we draw from a multinomial distribution using a normalized vector G*I to

determine the source of infection. For each scenario, involving a given source country (for example, US, Canada, UK, France and China), we run 1,000 simulations, allowed to run over a 3-year time period and assess the probability of swine flu invasion in each non-source country, and its most likely source of infection. We conducted sensitivity analyses with higher and lower values of the free parameter c and R 0, which mostly affected the time course of the global epidemic and the synchronicity of epidemics across locations, but did not change markedly the identification of hotspots countries for the onward spread. To explore the probability of reassortment between viruses originating from North America and Europe, we ran 1,000 independent simulations of epidemics starting in North America (USA and Canada), and in each the 10 European countries with largest swine populations, and computed the coinvasion probability in country i following:

\[ P_{\text{coinv},i} = \left[ 1 - P_i \right] \times \left( 1 - P_{\text{coinv},i}(1 - P_i) \right) = \left[ 1 - P_i \right] \times \left( 1 - p_{\text{inv},i} \right) \]

where p_{\text{inv},i} is the probability that country i is invaded when the outbreak is seeded in country k.

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Author contributions

M.R.C., S.E.D. and D.E.W. generated the data used in the analysis. M.I.N., C.V. and P.L. conceived, designed and performed the analysis. M.I.N., C.V., A.L.V., M.R.C., S.E.D., D.E.W., A.R., M.A.S., E.C.H. and P.L. wrote the paper. The opinions expressed in this article are the author’s own and do not reflect the view of the Centers for Disease Control and Prevention, the Department of Health and Human Services or the United States government.

Additional information

Accession codes: All sequences from swine influenza A viruses that were generated and used in this study have been deposited in the GenBank database with the accession codes AHB20822 to AHB20891, AHB20919 to AHRK3256, AHB23664 to AHB23860, AHB24418 to AHB24833 and AHB20876 to AHB20868. Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications.

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