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Permalink
https://escholarship.org/uc/item/1rf139h95

Journal
Proceedings of the Royal Society B: Biological Sciences, 280(1762)

ISSN
0962-8452

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Publication Date
2013-07-07

DOI
10.1098/rspb.2013.0872

Peer reviewed
## Inferring patterns of influenza transmission in swine from multiple streams of surveillance data

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Swine populations are known to be an important source of new human strains of influenza A, including those responsible for global pandemics. Yet our knowledge of the epidemiology of influenza in swine is dismayingly poor, as highlighted by the emergence of the 2009 pandemic strain and the paucity of data describing its origins. Here, we analyse a unique dataset arising from surveillance of swine influenza at a Hong Kong abattoir from 1998 to 2010. We introduce a state–space model that estimates disease exposure histories by joint inference from multiple modes of surveillance, integrating both virological and serological data. We find that an observed decrease in virus isolation rates is not due to a reduction in the regional prevalence of influenza. Instead, a more likely explanation is increased infection of swine in production farms, creating greater immunity to disease early in life. Consistent with this, we find that the weekly risk of exposure on farms equals or exceeds the exposure risk during transport to slaughter. We discuss potential causes for these patterns, including competition between influenza strains and shifts in the Chinese pork industry, and suggest opportunities to improve knowledge and reduce prevalence of influenza in the region.

### 1. Introduction

The 2009 pandemic highlighted the inadequate state of influenza surveillance in swine populations around the world. Mounting evidence indicates that pigs play a central role in the emergence of many pandemic influenza strains [1,2], and it is now known that influenza A (H1N1)pdm09 was created by reassortment of Eurasian avian-like (EA) and triple reassortant (TR) strains [3,4]. Phylogenetic reconstruction indicates that elements of the pandemic strain had been co-circulating undetected in swine for more than 10 years [3,4]. This finding underscores the need to study the epidemiology of influenza in swine populations at regional scales, and to understand the factors driving transmission dynamics. Achieving this goal requires long-term, systematic data from structured surveillance activities, but unfortunately almost no such datasets exist. One conspicuous exception is a dataset describing long-term virological and serological surveillance at a Hong Kong abattoir from 1998 to 2010, drawn from source farms across southern and southeastern China. These data have already yielded important insights about the co-circulation of influenza strains, strain replacement and viral reassortment in swine populations [5,6]. However, fundamental questions have yet to be addressed. What epidemiological processes give rise to the observed rates of virus
isolation? Where in the swine production system, from farms to abattoirs, does most transmission of influenza occur? How are transmission dynamics affected by strain interactions and the ongoing industrialization of swine production? Answering these questions would guide efforts to control influenza in swine, and would shed light on mechanisms underpinning future pandemic risks. Yet inferring these patterns of disease transmission, a challenge in the best of circumstances, is particularly difficult here, because the available dataset was collected at a single location, just prior to slaughter, preventing the direct study of transmission at different points in the swine production system.

Here, we introduce a statistical inference framework that uses a Bayesian state–space model (BSSM) to integrate simultaneously observed data streams to infer hidden patterns of transmission [7,8]. This approach analyses virological and serological data together, informed by studies of the progression of these clinical signs of infection in individual pigs [6,9,10]. Experimental infection studies demonstrate that a naïve pig infected with influenza can be expected to shed virus for 5–7 days. We define the window of recent exposure as the week prior to the sampling date; successful virus isolation indicates exposure during this period. The shedding period ends as antibodies to the infecting influenza strain increase; seropositivity reflects older exposure, which we define as infection more than one week prior to the sampling date. Virological and serological data are typically analysed separately, even when presented in the same study, but joint analysis draws additional insights from these data streams by exploiting their biological connection. Because samples are drawn simultaneously at fixed points in space and time, the differing time scales of the clinical signs provide insights into when transmission occurred, and, given knowledge of host movements, where transmission occurred.

China has long been recognized as a priority for influenza surveillance owing to the high densities of humans, swine and fowl in the region [11]. Currently, China produces and consumes almost 50% per cent of the world’s pork, requiring an enormous swine population [12,13]. Over the past decade, swine production systems in China have responded to rising pork demand from a human population growing in size and affluence, and have also been impacted by rising feed costs, natural disasters and significant disease outbreaks [12–14]. Resulting changes in the farming facilities, transport systems and patterns of trade will be reflected in the long-term influenza surveillance dataset collected at the abattoir, as will any epidemiological differences among circulating strains. In the week before slaughter, pigs are exposed to increased population mixing in trucks and holding pens, as well as ‘transport stress’ that may affect immunity to infection [15,16]. During their earlier life on the farms, infection risk will be influenced by animal density, facility design and biosecurity [17]. We investigate how these risks of exposure to influenza have increased over the past decade in Chinese swine, by using our inference framework to estimate older exposures (corresponding to time spent on the farm) and recent exposures (corresponding to transport and holding just before slaughter).

2. Model and results

(a) Framework for analysis of surveillance data

The dataset consists of regular samples from active surveillance at a Hong Kong abattoir from May 1998 through January 2010. Tracheal and nasal swabs were collected fortnightly from swine at slaughter, and monthly viral isolation rates, as well as corresponding strain typing, are available for almost all of the surveillance period. Sera were collected from swine at slaughter, and monthly seroprevalence data are available for the years 2000, 2004 and 2009 (see §4 for detailed information about the abattoir, viral isolation, strain typing and serology processing methods). Unavailable data are treated as missing values and imputed using the model.

The BSSM provides a probabilistic description of hidden process dynamics and their relation to the observed virus isolation and serological data (figure 1; see also the electronic supplementary material). Following convention [7,8], we divide the state–space model into (i) a data model, which provides the connection between model parameters and data in the form of the likelihood, (ii) a process model, which describes hidden dynamics and their relation to observed data, and (iii) a parameter model. Observed data include the number of strain-specific virus isolations per month, which we denote by \( n_i(t) \) for strain \( i \) in month \( t \). Monthly counts of seropositive samples (defined as having a titre of greater than or equal to 1:40 to any test antigen) are designated as \( n_i(t) \). Serological data were not divided into strain-specific responses owing to ambiguities in assignment. This use of serological data assumes that seropositivity reflects protection from contemporary viruses (see electronic supplementary material, figure S1; see discussion in §3b). The data model treats the virus isolation samples for each

![Figure 1. A diagram of the Bayesian state–space model consisting of data, process and parameter models (see main text for definition of all parameters). Dashed lines connect observations in the data model to relevant probabilities for each month \( t \). Solid black lines show relations between variables that make up the process model. Arrows indicate probabilities that covary positively, whereas a bar indicates suppression of one probability by another using a factor of one minus the relevant probability. Solid grey arrows indicate a deterministic relationship as in \( p_{i|v}(t) = p_i(t)p_{i|v}(t) \). The parameter model includes constant scale factors that represent the size of uncertainties by allowing different levels of variance in relations between parameters, as well as initial conditions \( p_{0|0}(0), p(0) \) and \( p(0) \) (not shown in the figure).](image-url)
month as multinomial, \( n_i(t) \sim \text{Mult}(p_{i,t}(t)) \), where the probability of isolating strain \( i \), \( p_{i,t}(t) \), is the product of the probability of virus isolation \( p_t(t) \) and the probability of finding type \( i \) given virus isolation, \( p_{i,1}(t) \). Serological data are binomially distributed, \( a_i(t) \sim \text{Bin}(p_{i,t}(t)) \); where \( p_{i,t}(t) \) is the probability of being seropositive in month \( t \).

The next layer of the process model connects the monthly data to the unobserved transmission dynamics described by the total probabilities of exposure to influenza on the farm or during transport to slaughter, denoted \( p_t(t) \) and \( p_f(t) \), respectively. We assume that previous exposure on the farm provides immunity to circulating strains of influenza, so that virus isolation is possible only when a naive animal is exposed within the week before sampling. This is reflected by the expectation \( \mathbb{E}[p_t(t)p_f(t)p_t(t)] = p_t(t)(1 - p_f(t)) \). Seropositivity is directly related to the total probability of exposure on the farm, \( \mathbb{E}[p_t(t)p_f(t)] = p_t(t) \). Using this model structure, both virus isolation and serostatus affect the inferred values of \( p(t) \) and \( p_f(t) \). The model includes scale parameters that control the variance in these statistical relations (see the electronic supplementary material). These parameters allow for uncertainty owing to unspecified sources such as assay accuracy, individual variation in animal response to exposure and month-to-month changes in the farms supplying the abattoir. All scale parameters are inferred along with the rest of the model parameters using Markov chain Monte Carlo (MCMC) sampling of the BSSM (see the electronic supplementary material, table S1).

The final element of the state–space model describes the temporal variation of the exposure parameters \( p_t(t) \), \( p_f(t) \) and \( p_{i,1}(t) \). Each of these parameters is assumed to vary as a Markov process, with expected value for the next month equal to the value in the current month. The month-to-month variance is determined by a scale parameter that is inferred from the data, as discussed earlier. This construct provides a constraint on the values of the hidden process dynamics, and potentially smooths outliers that arise from erratic sampling of a broad geographical region. It also allows us to impute missing data that make up part of our inference problem.

(b) Virus isolation and seropositivity

The posterior mean for the probability of virus isolation, without regard for strain type, follows the trends exhibited in the raw data (figure 2a). As noted in earlier work, virus isolation rates appear to decline over the past decade, with a protracted period of low isolation rates from 2005 to 2008 [6]. The posterior mean exhibits less variation than the raw frequency owing to the total probability of exposure during transport (figures 2b and 3a). Successful virus isolation at the abattoir occurs only when exposure during transport coincides with a lack of earlier exposure on the farm. A monthly estimate of this combined probability, given by \( p_t(t) \times (1 - p_f(t)) \), shows that the posterior mean for recent exposure of naive animals has decreased (see the electronic supplementary material, figure S3). Although this trend is not significant at the 95 per cent level, owing to the multiplication of uncertainties, it is consistent with the observed decrease in virus isolation rates over the decade. It is possible that pigs systematically spent longer in transport during periods of lower virus isolation than they did during periods of higher virus isolation. However, because of economic pressures this confounding variation in transportation time seems unlikely to have occurred. In any case, the inferred force of infection (FOI) during transport is insufficient to create this effect (i.e. the susceptible pool would not be exhausted within any reasonable period of transport, so new infections would continue to occur).

(d) Force of infection for swine influenza

The inference of the probabilities of exposure in transport and on the farm enables us to estimate the weekly FOI in each setting (figure 4). For a given exposure time \( T_e \) and probability \( p_xe \), the FOI can be estimated using FOI = \( -T_e \ln(1 - p_x) \). This relation is obtained by a simple rearrangement of the more familiar form: \( p_x = 1 - \exp(-\text{FOI} \times T_e) \) (see [18] for derivation). Because \( p_x(t) \) describes exposures in the week before slaughter, the weekly FOI during transport is virtually...
identical to figure 3a and has remained approximately constant at roughly 0.03 per week over the decade \( (T_e = 1 \text{ week and } \rho_e = \rho_e(t) \text{ for each month}) \). Estimation of the FOI for pigs on the farm depends on their age at slaughter and the duration of protection by maternally derived antibodies, which may prevent infection and development of haemagglutination inhibition (HI) titres [10]. Given typical ranges of 16–24 weeks of age at slaughter and 4–8 weeks of protection by maternal antibodies, we used possible exposure durations of 8, 16 and 24 weeks to estimate a range of possible FOIs \( (T_e = 8, 16, 24 \text{ weeks and } \rho_e = \rho_e(t) \text{ for each month}) \). Vaccination of swine against influenza is not reported for Chinese farms and should not affect exposure durations in these calculations [13,19].

Estimates of the weekly FOI on swine farms and during transport to slaughter are shown in figure 4. The 95 per cent HPD region is provided for the transport FOI and one farm FOI to give an indication of the scale of uncertainties; HPD regions for the farm curves are larger due to the greater uncertainty in seropositivity (figure 2c). For most assumptions about exposure duration on the farm, the weekly FOI is statistically indistinguishable between farm and transport phases over the decade. However, when the period of exposure on farms is assumed to last just eight weeks, the weekly FOI is significantly higher on farms than in transport from 2007 onwards. The robust pattern across all scenarios is that, despite the stress and increased mixing associated with transport and holding before slaughter, the weekly risk of exposure to influenza appears to be equal or higher on the production farms.

(e) Strain replacement

During the decade of surveillance, one of the major features observed was an replacement of the classic swine H1N1 (CS) with an EA strain as the dominant viral strain in southern China (previously described by Vijaykrishna et al. [6]). Experimental infection studies demonstrated a competitive advantage of EA over CS strains, as well as limited immunity in the population, suggesting a biological mechanism for this replacement [6]. The state–space model provides insights into the dynamics of strain replacement by inferring monthly,
strain-specific viral isolation rates (see the electronic supplementary material, figures S4 and S5).

The emergence of a fitter influenza strain could account for the rise in on-farm exposure revealed by our analysis. However, the EA strain appeared in 2000–2001 and became dominant in 2003–2005, whereas the rise in exposure on farms occurred mostly from 2004 to 2008 (figure 3b). This suggests that the strain replacement was not the immediate cause of the changing epidemiological pattern. Instead, we propose that large-scale changes in the swine industry are the more likely explanation, and indeed may have played an important role in the strain dynamics via introduction of novel strains and emergence of reassortant strains. The geographical sourcing of swine for the sampled abattoir shifted significantly over the decade [6].

From 2000 to 2007, 15–20 per cent of the pigs were farmed near Hong Kong, with the remainder imported from several provinces in China; by 2008, the local proportion had fallen to 5 per cent. The percentage of swine sourced from nearby Guangdong province increased from 31.1 in 2003 to 51.8 in 2010. These shifts in industry-wide patterns, along with accompanying changes in conditions at production facilities, should be considered as potential causes of the observed shifts in epidemiology and strain frequencies, alongside the biological mechanisms already proposed.

3. Discussion

The long-term, active surveillance data considered here provide a rare glimpse into the transmission dynamics of influenza in swine populations over the decade leading up to the 2009 pandemic. While the pandemic did not arise in China, the co-circulation of multiple lineages and genetic reassortment events observed in the course of this surveillance provide insight into likely virus transmission dynamics leading up to pandemic emergence in the Americas [5,6]. The joint analysis of virological and serological data allows us to infer time-varying risks of influenza exposure across production stages, despite the fact that samples were taken at a single location. To untangle this information, we used knowledge gained from experimental infection studies of influenza in swine to define recent and older time frames for infection, indicated by the potential for virus shedding and detection of antibodies, respectively. Using this connection, we separated exposure risks during transportation and holding the week before sampling from exposures occurring during earlier life on swine farms in southern and southeastern China.

The model presented here provides a statistical framework to link experimental infection studies with surveillance data streams to quantify epidemiological processes, both observed and hidden. We can infer that influenza prevalence in the region has not decreased despite an apparent reduction in virus isolation rates (see figure 2; electronic supplementary material, figure S1). Instead, we find that the FOI during transport in the week before slaughter has remained remarkably constant at approximately 3 per cent per week over the decade. By contrast, the cumulative probability of exposure on the farm has increased over this period. We conclude that the observed reduction in virus isolation rates reflects the rise in exposure early in life, which results in immunity that prevents infection in the week before sampling. This finding has two important implications: (i) a perceived reduction in
influenza prevalence at slaughter is not due to improved biosecurity or other preventative measures; and (ii) the overall prevalence of influenza in the swine population has not decreased, and hence the risk of spillover to susceptible human and bird populations is constant or growing. Our findings show that the risk, as measured by the FOI, is at least as high on swine farms as it is in transport and holding settings. Further, influenza on the farms appears to have increased over the decade, although the statistical significance of this trend would be strengthened with additional data.

(a) Epidemiology of swine influenza
The increasing scale of the pork industry in China provides the backdrop for the findings reported here. Over the past decade, a series of market-driven and natural perturbations (including rising feed costs, the 2008 Sichuan earthquake and outbreaks of non-influenza disease) have led to major disruptions in the Chinese swine population [12–14,19,20]. The resulting economic impacts motivated a push towards production in larger, more regulated facilities in an effort to stabilize supply, limit price increases and minimize disease outbreaks [13,19]. The growing size of swine farming facilities has increased the potential for sustained influenza transmission, particularly given the apparent lack of vaccination against influenza. The rise in on-farm exposure reported here could be attributed to these factors, and related changes linked to industrialization [21].

Our study reveals long-term patterns of influenza circulation under these changing conditions, inferred from data collected from a random subset of animals passing through the abattoir in Hong Kong. We estimate the rate of virus isolation to range from 0.1 to 10 per cent, with a mean of 1.6 per cent over the decade (see figure 2; electronic supplementary material, figure S3), whereas seropositivity ranges from 24 to 74 per cent, with a mean of 50 per cent (figure 2). Many of the high rates of isolation occur in the early 2000s, with relatively low seroprevalence; this pattern shifts to lower isolation rates and higher seroprevalence late in the decade. Here, we place our observations in the context of shorter-term epidemiological studies conducted elsewhere in the world.

Samples collected in 1997–1998, during fortnightly surveillance of swine at an abattoir in the United States, showed a mean virus isolation rate of 2.2 per cent, with monthly peaks as high as 16 per cent [22] (see also [23]). The mean seroprevalence, using abattoir samples as well as sera sent to the Wisconsin Animal Health Laboratory for pseudorabies virus testing, was 27.7 per cent for a strain of CS H1 influenza, with monthly values ranging from 10 to 60 per cent. These observations are consistent with our data from Hong Kong, both on average and with regard to the wide variation in monthly rates; intriguingly, the quantitative match is closer during the 1998–2004 period, when our data were also dominated by the CS strain.

A survey of Spanish swine farms during 2008–2009 found a seroprevalence of 75.4 per cent to at least one of three strains of influenza (H1N1, H1N2 and H3N2) [17]. These results were broken down by age, with seropositivity among younger fattening pigs (11–20 weeks of age, a closer match to animals included in our study) measured to be 53.1 per cent, while that among older cows was 89.9 per cent. Similar values for sows of varying ages were obtained for pig-dense European countries during a 2002–2003 survey with 85.2 per cent seroprevalence in Germany and 94 per cent in Belgium [24]. A recent survey of influenza prevalence in other Asian countries showed mean levels of virus isolation and seropositivity similar to our results for Hong Kong [21,25]. We conclude that the epidemiology of swine influenza in southern China is broadly consistent with available data from other industrialized countries worldwide.

Our findings show that efforts to limit the prevalence of influenza A in Chinese swine populations could usefully be targeted to improving practices in production facilities. Such control efforts could be informed by recently identified risk factors for influenza on Spanish swine farms: (i) high replacement rates that introduce new susceptibles or new subclinically infected animals; (ii) lack of solid separations to prevent transmission between pens; and (iii) uncontrolled entrance to farms [17]. In contrast to the situation on farms, we found that the probability of influenza exposure during the week before slaughter has not risen or fallen as a result of industry growth over the past decade. This suggests that there are unrealized opportunities to reduce spread during transport and limit potential for mixing of geographically separated strains. For example, transmission between pens holding pigs from different farms has been observed at a quarantine point in Shenzhen and provides an opportunity for improvement (see §4). However, the validity of comparisons between this system and Spanish farms remains unclear, and the economics of swine production raise challenges in motivating changes in practice for influenza control.

(b) Impact of model assumptions
The results of our analysis arise from the probabilistic relations that make up the BSSM. The strongest assumption we made is that antibodies above a certain titre are protective against challenge by any contemporary influenza strain. In our analysis, exposure probabilities are inferred by considering virus isolation and serostatus data independent of the strain type or detailed serological profile. Patterns of serological cross-reactivity and immunity in influenza are complicated, even when the test antigen is known and experimental infection is done under laboratory conditions [6,9,26]. As a result it is not possible to definitively connect a given serological profile to prior infection by one or more known strains. Under field conditions, this problem is amplified by the fact that the observed serological profiles might be caused by one or more viral strains that are not used as $n$ strains in HI panels, and have not been characterized by experimental infection studies.

A significant proportion of swine sampled in our dataset had titres greater than or equal to 1:40 to many, or all, strains tested (see the electronic supplementary material, figure S1). This pattern could arise from cross-reactivity among the strains or from multiple infections by different strains. However, the age of pigs at the time of slaughter ranges from four to six months, and it is usually assumed that only a single infection is possible during this time (though this assumption may not hold in all cases [27,28]). As a result, we argue that the assumption of immunity to contemporary strains is appropriate, and that strain-specific analyses of our data are not possible given current knowledge of influenza serology. If the assumption is too strong, and cross-immunity among these strains is not complete, then our results would overestimate the probability of recent exposure (because we would have underestimated the true proportion of swine that are susceptible to infection). To explore this possibility, we re-ran the BSSM assuming 50 per cent protection for
seropositive animals. This model, which could account for second infections by different strains or for limited protection from previous exposure, reduces the posterior mean probability of recent exposure to 2.4–2.5 per cent (from approx. 3%) over the decade. The difference in these estimates, given the related uncertainties, does not substantially change the conclusions presented earlier. Further progress on this problem will require experimental infection studies, including multiple exposures and longitudinal sampling of serum from individual animals, as well as development of methods that elucidate the connections between immunity and profiles derived from field serological data [9,29].

The model includes constant scale parameters that determine the variances in association between elements of the process model (see figure 1; electronic supplementary material). In table S1 of the electronic supplementary material, we show the inferred values, which vary inversely with the variance, along with regions of 95 per cent HPD. The scales for virus isolation and serostatus are low, and allow for considerable variance in the relation to the hidden probabilities of exposure. This uncertainty can be attributed to month-to-month variation in source farms, assay sensitivity and individual variation, as well as other unknown factors. The estimated scale parameters for hidden process dynamics are larger, allowing for less variation and resulting in smoothed dynamics, hence limiting inordinate weight to data from any single month.

(c) Broader applications of multi-stream surveillance analysis

We have introduced a mechanistic state-space model to infer unobserved transmission histories from joint analysis of virological and serological surveillance data. Our model provides robust inference of epidemiological parameters, smoothing anomalous data and imputing missing data based on mechanistic assumptions. Multiple modes of surveillance are commonly conducted in parallel, and other studies could benefit from similar joint analyses of these data streams [25,30–32]. In practice, sampling design is very important when applying this idea, because all data streams need to be sampled at the same time points, from the same population. Ideally, this sampling would occur at regular intervals from randomly selected animals, providing an unbiased picture of disease dynamics in the focal population.

This approach can be adapted to any disease for which the time course of multiple clinical outcomes, including diagnostic measures or signs of disease, have been characterized by experimental infection studies or close observation of natural infections [33]. Given different time scales associated with different clinical outcomes, such analyses would open broader opportunities to reconstruct the histories of exposure prior to sampling. Further quantitative precision could be obtained by expanding experimental infection studies to characterize individual-specific variation, and estimate the sensitivity and specificity of different assays. In principle, the kinetics of some assays could be characterized so that quantitative results could be used, instead of binary outcomes, to derive more specific information about the time since infection. Further theoretical and statistical tools to integrate these classes of data will also need to be developed. Combined with a carefully designed surveillance plan, these approaches could enable the inference of disease transmission histories in systems where host individuals can only be sampled once, an all-too-common limitation in settings ranging from livestock inspections to wildlife studies.

4. Material and methods

(a) Surveillance data

Systematic influenza surveillance was conducted between May 1998 and January 2010 at an abattoir in Hong Kong where roughly 4500 pigs per day are processed (current data are available online at http://www.fehd.gov.hk/english/sh/index.html). Tracheal or nasal swabs (109–512 per month) and serum samples (20–50 per month) were collected from a random sample of slaughtered swine [6] (see the electronic supplementary material, figure S1 for raw data). Sampled swine were between the ages of four and six months, and were sourced from provinces throughout southern and southeastern China [6]. Information that connects source farm, transport time, transport method and disease status of individual animals is unavailable. We have no information about the size of farms that provide pigs to this abattoir, but McOrist et al. [19] provide data on the size of pig farms in China and their growth between 2008 and 2011. Pigs are transported in trains and lorries, with the transport period varying depending on the starting farm location (see supplementary information in [6] for information on source farms). The pigs are held at a border quarantine station in Shenzhen, just across the border from Hong Kong, to check health status before transport to Hong Kong. Pigs from each source farm are held in separate pens, but they may be next to a pen with animals from a different farm and there is evidence of transmission among pens. After the quarantine station, each consignment is shipped separately to the abattoir in Hong Kong.

(b) Virus isolation, strain typing and seroprevalence

Swab materials were inoculated into 9- to 10-day-old embryonated chicken eggs and Madin Darby canine kidney cells; virus isolates were identified and subtyped by HI assays as previously described [6]. In addition, strains included in the study were fully sequenced and typed based on phylogenetic relationships of all gene segments [4]. Antibody prevalence against the major swine influenza lineages was characterized using the HI assays with six representative viruses—A/Sw/HK/4167/1999 (CS), A/Sw/HK/1110/2006 (TR), A/CA/4/2009 (Pand), A/SW/HK/1304/2003 (CS), A/Sw/HK/NS29/2009 (EA) and A/SW/HK/1599/2008 (EA)—starting at a dilution of 1:10. Titres greater than or equal to 1:40 were taken as seropositive [6].

(c) Bayesian state-space model and Markov chain Monte Carlo sampling

The BSSM is a hierarchical model with time-dependent associations as developed in earlier studies [7,8]. Complete details of the statistical framework are provided in the electronic supplementary material. Computational elements of the MCMC inference process use the Metropolis–Hastings algorithm with tuning during burn-in to sample from the posterior distribution, as implemented by the PyMC PYTHON package [34].

This work was funded by a grant from the National Center for Foreign Animal and Zoonotic Disease Defense, a Department of Homeland Security and Technology Center of Excellence. Data collection was supported by the National Institute of Allergy and Infectious Diseases (contract no. HHSN266200700003C). J.O.L.-S. and S.R. are supported by the RAPIDD programme of the Science and Technology Directorate of the Department of Homeland Security and NIH Fogarty International Center, J.O.L.-S. is grateful for support from the De Logi Chair in Biological Sciences. We thank the Institute for Digital Research and Education at UCLA for use of the hoffman2 cluster.
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