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ABSTRACT

Sporadic activity by H5N2 influenza viruses has been observed in chickens in Taiwan from 2003 to 2012. The available information suggests that these viruses were generated by reassortment between a Mexican-like H5N2 virus and a local enzootic H6N1 virus. Yet the origin, prevalence, and pathogenicity of these H5N2 viruses have not been fully defined. Following the 2012 highly pathogenic avian influenza (HPAI) outbreaks, surveillance was conducted from December 2012 to July 2013 at a live-poultry wholesale market in Taipei. Our findings showed that H5N2 and H6N1 viruses cocirculated at low levels in chickens in Taiwan. Phylogenetic analyses revealed that all H5N2 viruses had hemagglutinin (HA) and neuraminidase (NA) genes derived from a 1994 Mexican-like virus, while their internal gene complexes were incorporated from the enzootic H6N1 virus lineage by multiple reassortment events. Pathogenicity studies demonstrated heterogeneous results even though all tested viruses had motifs (R-X-K/R-R) supportive of high pathogenicity. Serological surveys for common subtypes of avian viruses confirmed the prevalence of the H5N2 and H6N1 viruses in chickens and revealed an extraordinarily high seroconversion rate to an H9N2 virus, a subtype that is not found in Taiwan but is prevalent in mainland China. These findings suggest that reassortant H5N2 viruses, together with H6N1 viruses, have become established and enzootic in chickens throughout Taiwan and that a large-scale vaccination program might have been conducted locally that likely led to the introduction of the 1994 Mexican-like virus to Taiwan in 2003.

IMPORTANCE

H5N2 avian influenza viruses first appeared in chickens in Taiwan in 2003 and caused a series of outbreaks afterwards. Phylogenetic analyses show that the chicken H5N2 viruses have H5 and N2 genes that are closely related to those of a vaccine strain originating from Mexico in 1994, while the contemporary duck H5N2 viruses in Taiwan belong to the Eurasian gene pool. The unusually high similarity of the chicken H5N2 viruses to the Mexican vaccine strain suggests that these viruses might have been introduced to Taiwan by using inadequately inactivated or attenuated vaccines. These chicken H5N2 viruses are developing varying levels of pathogenicity that could lead to significant consequences for the local poultry industry. These findings emphasize the need for strict quality control and competent oversight in the manufacture and usage of avian influenza virus vaccines and indicate that alternatives to widespread vaccination may be desirable.

In the last decade, H5N2 influenza virus activity has been observed occasionally in chickens in Taiwan. The initial outbreak of LPAI H5N2 viruses occurred in late 2003, and a second LPAI outbreak was reported in 2008 (19). Genetic analyses of publically available sequences of the Taiwanese H5N2 viruses suggested that their surface protein genes, HA and neuraminidase (NA), were not derived from Eurasian gene pool viruses but were closely related to an H5N2 virus isolated from chickens in Mexico during an outbreak in 1994 (19). Their internal genes originated from an
H6N1 virus lineage that has been enzootic in chickens in Taiwan for the last 2 decades (19, 20).

Since January 2012, H5N2 virus activity has been observed almost every month and has covered many regions of Taiwan, resulting in significant mortality in chickens (15, 21). However, how this virus was generated and how it became prevalent in the field remain to be answered. To explore these questions, avian influenza surveillance was conducted from December 2012 to July 2013 at a wholesale live-poultry market in Taipei where birds were shipped from different areas of Taiwan. Our findings show that both H5N2 and H6N1 influenza viruses were cocirculating in chickens in Taiwan at that time. All Taiwan H5N2 viruses from chickens had multiple basic amino acids at the cleavage site of the HA connecting peptides, creating motifs that are associated with high pathogenicity in chickens. However, no additional basic amino acids were present, and the structurally adjacent glycosylation site that may influence pathogenicity was conserved in all viruses (10–13). Phylogenetic analyses revealed that these H5N2 viruses were introduced to this region from an external source but have undergone multiple reassortments with the enzootic H6N1 virus lineages.

**MATERIALS AND METHODS**

**Surveillance and virus isolation.** Influenza surveillance of live poultry was conducted in Taipei at the wholesale market. Sampling was conducted weekly from apparently healthy chickens and ducks from December 2012 to July 2013 (Fig. 1). To avoid contamination and expand representation, no more than four fecal samples were collected from each cage and only fresh droppings were taken. Samples were kept in a cool box and shipped to the laboratory within 2 h. Virus isolation was conducted using 9- to 11-day-old embryonated chicken eggs. Hemagglutinin-positive isolates were subtype by hemagglutination inhibition (HI) and neuraminidase inhibition (NI) assays using a panel of reference sera, as described previously (22, 23).

**Serological survey.** Thirty to 100 chicken blood samples were collected on each sampling occasion. Chicken sera were isolated and stored at −20°C. Serum samples collected from December 2012 to July 2013 were tested by HI for antibodies against Clk/TW/853/12 (TW853; H3N8), Dk/TW/1030/12 (TW1030; H6N8), Clk/TW/2593/12 (TW2593; H5N2), Clk/TW/2267/12 (TW2267; H6N1), and Clk/HK/NT155/08 (HK-NT155; H9N2). Prior to the tests, all chicken sera were treated by receptor-deactivating enzyme (RDE; Denka Seiken Co., Ltd., Tokyo, Japan) and absorbed using fresh chicken red blood cells to remove nonspecific inhibitors and substances responsible for nonspecific agglutination. Sera were tested starting at a 1:10 dilution in phosphate-buffered saline (PBS).

**Virus sequencing and phylogenetic analyses.** Full-genome sequences of all the H5 viruses from chickens (n = 16) and ducks (n = 12) isolated during the survey period and selected chicken H6 viruses (n = 7) from each sampling occasion were obtained as previously described (24). Sequencing was performed using a BigDye Terminator, version 3.1, cycle sequencing kit on an ABI 3730 genetic analyzer (Applied Biosystems) according to the manufacturer’s instructions. DNA sequences were compiled and edited using Lasergene, version 8.0 (DNASTAR, Madison, WI). Multiple sequence alignments were compiled using MUSCLE, version 3.8 (25). The maximum-likelihood (ML) phylogeny of each gene segment was inferred using the GTR+I+F nucleotide substitution model (general time-reversible model with a proportion of invariant sites and gamma-distributed rate variation across sites) in PhyML (version 3.0) (26). Robustness of the ML topology was evaluated by 1,000 bootstrap replicates, and the Bayesian clade posterior probability was determined from the topologies sampled in the Bayesian molecular clock analysis.

**Molecular clock analysis.** For the HA and NA genes, the genetic distance from the common ancestral node of the lineage to each viral isolate was measured from the ML tree and plotted against the sample collection dates. Linear regression was used to indicate the rate of accumulation of mutations over time. A more detailed evolutionary time scale for each virus gene phylogeny, with confidence limits, was obtained using relaxed molecular clocks under uncorrelated lognormal (UCLD) and exponential (UCED) rate distributions, implemented in a Bayesian Markov chain Monte Carlo (BMCMC) statistical framework (27), using BEAST, version 1.8 (28). The SRD06 nucleotide substitution model (29) and Bayesian Skytree demographic model (30) were used. Multiple runs were performed for each data set, giving a total of 6 × 10^7 states (with 1 × 10^7 states discarded as burn-in) that were summarized to compute statistical estimates of the parameters. Convergence of the BMCMC analysis was assessed in Tracer, version 1.6 (A. Rambaut M. Suchard, and A. J. Drummond, 2015 [http://tree.bio.ed.ac.uk/software/tracer/]).

**Pathogenicity study.** Three H5N2 isolates (from groups A and B of the H5N2 viruses, with two different motifs (R-E-K-R or R-K-K-R) at the cleavage site of the HA connecting peptide, were selected to determine the intracecal pathogenicity index (ICPI) (31). Freshly harvested allantoic fluid from embryonated chicken eggs containing viruses of a hemagglutination titer of 1.024 was diluted to 1:10 with sterile PBS, and 50 μl was intracebrally inoculated into each of 10 1-day-old chicks (24 to 36 h after hatching). Mock-infected birds were inoculated with 50 μl of PBS. All birds were examined every 24 h for 8 days and scored as 0 if normal, 1 if sick, and 2 if dead. The mean scores per bird per observation over the 8-day period were calculated. A virus with a value of ≥0.7 is considered pathogenic, and a virus with a value of ≥1.5 is considered highly pathogenic.

Animal experimental protocols were approved by the Institutional Ethical Review Board of Shantou University Medical College (reference no. SUMC2012-134).

**Nucleotide sequence accession numbers.** The nucleotide sequences obtained in the present study are available from GenBank under accession numbers KJ162585 to KJ162864.
the viruses in chickens are less diverse than those in ducks and that chicken viruses (by serological measurements against recent Taiwanese duck or of the prevalence of influenza viruses in chickens was also assessed and 3.7%, respectively. Most of these isolates were detected in the also detected in chickens and ducks with isolation rates of 0.3% viruses were isolated. Avian paramyxovirus type 1 (APMV-1) was /H11022 from the gene pool, which was very likely a cross-reaction with the 1:10 to 1:40) with an H6N8 virus (TW1030) to a 2012 H6N1 chicken virus (TW2267) from the enzootic lin- domestic ducks (influenza viruses, with 12 H5 (from the gene pool. Viruses isolation and prevalence. All of the H5 viruses isolated from ducks belonged to the Eurasian gene 6579|May 2014 Volume 88 Number 10|jvi.asm.org

TABLE 1 Isolation of influenza viruses and APMV-1 in domestic ducks and chickens in Taiwana

<table>
<thead>
<tr>
<th>Host and sampling time</th>
<th>No. of samples</th>
<th>No. of influenza virus isolatesb</th>
<th>No. of APMV-1 isolates</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>H5</td>
<td>H6</td>
</tr>
<tr>
<td>Chicken</td>
<td></td>
<td>Total</td>
<td>type</td>
</tr>
<tr>
<td>December 2012-2013</td>
<td>2,607</td>
<td>22</td>
<td>4</td>
</tr>
<tr>
<td>January 2013</td>
<td>1,771</td>
<td>18</td>
<td>5</td>
</tr>
<tr>
<td>February 2013</td>
<td>517</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>March 2013</td>
<td>1,339</td>
<td>14</td>
<td>6</td>
</tr>
<tr>
<td>April 2013</td>
<td>535</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>May 2013</td>
<td>814</td>
<td>7</td>
<td>1</td>
</tr>
<tr>
<td>June 2013</td>
<td>477</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>July 2013</td>
<td>599</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>8,659</td>
<td>67</td>
<td>16</td>
</tr>
</tbody>
</table>

Duck December 2012-2013 286 18 0 3 15 1

January 2013 106 17 10 2 5 17
February 2013 137 0 0 0 0 11
March 2013 557 49 0 3 46 52
April 2013 193 0 0 0 0 0
May 2013 539 9 0 1 8 4
June 2013 352 3 0 0 3 4
July 2013 410 36 2 26 8 7
Total 2,580 132 12 35 85 96

a APMV-1, type 1 avian paramyxovirus.
b Three subtypes of influenza viruses were detected in chickens. These were H3, H5N2, and H6N1. All of the H5 viruses isolated from ducks belonged to the Eurasian gene pool.

RESULTS

Virus isolation and prevalence. From 8,659 chicken dropping swabs, 16 H5N2 (~0.2%), 46 H6N1 (~0.5%), and a total of 5 H3 viruses were isolated. Avian paramyxovirus type 1 (APMV-1) was also detected in chickens and ducks with isolation rates of 0.3% and 3.7%, respectively. Most of these isolates were detected in the cooler months from December to March (Table 1). Samples from domestic ducks (n = 2,580) showed a slightly higher prevalence of influenza viruses, with 12 H5 (~0.5%), 35 H6 (~1.4%), and 85 (~3.3%) of other subtypes isolated. These findings suggest that the viruses in chickens are less diverse than those in ducks and that H5 viruses are prevalent at a low level in both ducks and chickens.

Serological survey of influenza viruses in chickens. Evidence of the prevalence of influenza viruses in chickens was also assessed by serological measurements against recent Taiwanese duck or chicken viruses (Table 2). Of 894 samples, none were seropositive to the chicken H3N8 virus (TW853) even though H3 viruses were isolated on three sampling occasions from December 2012 to January 2013. The seropositive rate (with HI titers of >1:20) to a recent chicken H5N2 virus (TW2593) was 8.3%, which was much higher than the virus isolation rate in this survey period. About 3.5% of serum samples were seropositive (with HI titers of >1:20) to a 2012 H6N1 chicken virus (TW2267) from the enzootic lineage in Taiwan. Only a small number (1.4%) reacted at marginal levels (HI titer of 1:10 to 1:40) with an H6N8 virus (TW1030) from the gene pool, which was very likely a cross-reaction with the chicken H6N1 virus as each of these samples had a high titer (HI of >1:40) against TW2267 (data not shown). In contrast to these results, 82.1% of the tested sera reacted with an HI titer higher than 1:20 against chicken H9N2 virus (HK-NT155) from the Ck/Bei/1/94 lineage that has been enzootic in chickens in mainland China for 2 decades even though H9N2 viruses have not previously been reported (nor were they detected in our current surveillance) in chickens in Taiwan.

Phylogenetic analyses. The HA gene of the duck H5 viruses from Taiwan clustered with Eurasian gene pool H5 viruses (Fig. 2). However, both the HA and NA of the chicken H5N2 viruses, isolated in our surveillance and earlier in Taiwan, formed a well-supported monophyletic lineage (ML bootstrap support of 98 to 100% and Bayesian clade posterior probability of 1.0) that descended from the HA and NA genes of Ck/Hidalgo/28159-232/94 (Fig. 2A and B). This H5N2 virus from the 1994 outbreak in Mexico was used as a commercial vaccine strain (32, 33). These findings suggest that the H5N2 viruses in chickens were introduced to Taiwan from Mexico either by bird movement or by inadequately inactivated or attenuated vaccines.

Within the Taiwan chicken H5N2 virus lineage, both the HA and NA genes of the viruses diverged into at least two subclades. One includes viruses isolated in 2008 and from 2012 to 2013 (group A), while the other contains viruses isolated from 2012 to 2013 (group B). After the initial outbreak in 2003, the Mexico-like H5N2 viruses might have been reintroduced to chickens on at least two independent occasions or diverged quickly after their introduction into separate subclades that have persisted independently.

Relative to the inferred most recent common ancestor (MRCA) of the Taiwanese and Mexican H5N2 viruses (circles in Fig. 2A and B), both lineages demonstrated similar accumulations of genetic substitutions over time, indicating similar evolutionary rates (Fig. 2C and D). However, the Taiwanese lineage showed an apparent shift of 10 years in its accumulation of differences from the common ancestor, consistent with the phylogenetic trees of the HA and NA genes, where the 2003 H5N2 virus from Taiwan appeared more similar to the 1994 than to the contemporary H5N2 Mexican viruses. Molecular clock models showed lower evolutionary rates in the branches of the HA and NA trees at or near the transition of viruses from Mexico to Taiwan (data not shown). These data are supportive of the unnatural introduction of a Mexican virus, originating around 1994, to Taiwan in 2003.

Phylogenetic analyses of each of the six internal genes (Fig. 3, 4, and 5) showed that all Taiwanese chicken H5N2 viruses grouped
with the H6N1 viruses that have been enzootic in chickens in Taiwan since 1997 (20). Two major groups, consistent with the groups in the HA and NA trees (Fig. 2A and B), were seen in the phylogenies of the internal gene trees of the H5N2 viruses (Fig. 3, 4, and 5). The larger group (group A) clustered with contemporary H6N1 viruses that were also isolated in our surveillance and with the recent human H6N1 virus (34), while the second group (group B) clustered with H6N1 viruses isolated from 2003 to 2005. A third subgroup of two group A H5N2 isolates with recent H6N1 viruses was found in the PB1 phylogeny but not in the phylogenies of the other genes. The internal genes did not show the evolutionary rates discrepancy observed in the HA and NA genes (Fig. 2A and B), suggesting that the H5N2 viruses might have reassorted near the time of their introduction with contemporary H6N1 chicken viruses. Overall, these findings suggest that these H5N2 viruses were generated through several independent reassortment events with enzootic H6N1 viruses.

**Molecular characterization.** As observed in the Mexican-like
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Although a full-length PB1-F2 protein is normally found in avian influenza virus isolates, the H5N2 and related H6N1 viruses from Taiwan had variable lengths for this protein. The group A viruses had a PB1-F2 protein of 57 amino acids (aa), the group B had a full-length protein of 90 aa, and the third group in the PB1 tree (Fig. 2) had a protein of 87 aa, while the 2003 isolate had a PB1-F2 of 34 aa. Truncation of PB1-F2 may moderate disease progression (40).

Three different motifs, R-E-K-R, R-K-K-R, and R-R-K-R, were found in the HA0 cleavage site were found, with both being isolated in the same sample set on only one occasion (March 2013) (35). In our survey period, only the R-E-K-R and R-K-K-R motifs at position 218 of the C terminus remained as E and D residues, consistent with what was commonly seen in avian viruses (39).

An S31N substitution was found in the M2 protein of the Taiwan H5N2 viruses also had a 20-amino-acid (aa) deletion at the stalk region of the NA protein (deletion of residues 63 to 82), indicating that these viruses were adapted to chickens (35). An S31N substitution was found in the M2 protein of the Taiwan H5N2 and most H6N1 viruses, suggesting resistance to adamantanes (36). Truncation at position 218 of the C terminus of the NS1 protein was also present in the group B chicken Taiwan H5N2 viruses, as seen in the recent H7N9 viruses and many H9N2 and H6N1 viruses from terrestrial poultry (37, 38). The host specificity markers of the PB2 protein, positions 627 and 701, remained as E and D residues, consistent with what was commonly seen in avian viruses (39).

FIG 3 Maximum-likelihood phylogenies of the PB2 and PB1 genes. Taxon names of sequences reported in this study are shown in blue (chicken H5N2 lineage), green (enzootic chicken H6N1 lineage), and purple (duck H5N2). The chicken H5N2 viruses initially isolated from Taiwan are in red. Virus subtypes are indicated in parentheses. Bootstrap support values (percent) from 1,000 pseudoreplicates are shown for selected lineages. The scale bar to the left of each tree represents 0.01 substitutions per site. Abbreviations for host species: Ck, chicken; Dk, duck; WDk, wild duck; Qa, quail; Gs, Goose; BHG, bar-headed goose. Human isolates do not use any host specification. HK, Hong Kong.
a 4-amino-acid connecting peptide from being highly pathogenic (10, 11, 41).

The residue at position −3 of HA1 had mutated from E in the Mexican and the 2003 Taiwan H5N2 viruses to K (or R) on several occasions from 2008 to 2013 (Fig. 2A and Table 3). Overall, the sequences from our study in Taiwan show seven cases of E and nine of K at position −3. The HA phylogenetic tree indicates repeated mutation from E to K (or R) at position −3 in the Taiwan H5N2 lineage, highlighting the possibility of multiple introductions of the virus into chickens in Taiwan.

Pathogenicity study. To evaluate the pathogenicity conferred by the different HA cleavage motifs, three H5N2 isolates (Ck/TW/1680/13, Ck/TW/8994/13, and Ck/TW/8996/13), which fell into groups A and B (Fig. 2A) and had either an R-K-K-R or R-E-K-R motif (Table 3), were inoculated into 1-day-old chickens to determine the ICPI of the viruses. Ck/TW/8996/13 (R-K-K-R, group A) was not pathogenic or lethal in chicks (ICPI of 0), while the highly related Ck/TW/8994/13 (R-K-K-R, group A) was isolated from the same sampling occasion, was fatal in 3 of the 10 chicks, giving an ICPI of 0.3375 (Table 4). The group B virus, Ck/TW/1680/13 (R-K-K-R), had an ICPI of 1.675 and killed all chickens by day 5 postinoculation (Table 4). Therefore, neither the R-E-K-R nor the R-K-K-R motif, in the context of a nearby glycosylation site, is sufficient to cause high pathogenicity, as measured by the ICPI. While a basic amino acid at position −3 of HA1 might be sufficient to cause some lethality to chicks, other molecular characteristics may be involved in the higher pathogenicity of Ck/TW/1680/13.

DISCUSSION

Our surveillance study has revealed that H5N2 viruses are prevalent at low levels in the chicken population of Taiwan. Phylogenetic analyses of these viruses show that they were generated by reassortment between an H5N2 virus that originated from Mexico in 1994 and a locally enzootic H6N1 virus lineage. Apparently, this
kind of reassortment has occurred on multiple occasions. How this Mexican-like H5N2 virus entered Taiwan is still not clear.

Introduction of influenza viruses from one region to another is usually associated with poultry movement, bird migration, or escape from an insufficiently inactivated or attenuated vaccine strain (22, 42, 43). Even though Taiwan is located on the East Asian migratory bird flyway and although intercontinental movement of influenza viruses related to flyways has occurred previously (44-46), influenza viruses from the North American lineage had not been documented in Taiwan before this H5N2 virus was detected in 2003. The Taiwanese Mexican-like H5N2 viruses that emerged around 2003 did not have genes of contemporary Mexican-like viruses but had surface genes more similar to a 1994 virus, Ck/Hidalgo/28159-232/94, that was used as a commercial vaccine strain (19) (Fig. 2A and B). Their internal genes were obtained by reassortment with the contemporary enzootic H6N1 viruses of Taiwan (20) (Fig. 3, 4, and 5). This time discrepancy between the emergence dates of the Mexican and Taiwanese viruses is reflected in the discordance of the dates on the phylogenetic tree and accumulation of substitutions from the lineage common ancestor (Fig. 2). A similar pattern occurred in the re-emergence of a 1950s-like seasonal human H1N1 virus in 1977 (47).

LPAI Mexican-like H5N2 viruses caused an outbreak in chickens in Japan from 2005 to 2006 (48). These LPAI viruses clustered, over all their segments, with contemporary Mexican H5N2 viruses but have not been detected in aquatic birds in Japan. These observations are highly suggestive that the H5N2 virus was artifically introduced to chickens rather than being introduced by migratory bird flyways, most probably as an escape from a vaccination program. Mexican-like H5N2 vaccines have been used previously in this region, for example, in Hong Kong from 2003 to 2013 (49), although their use in Japan or Taiwan has not been officially reported.

Our phylogenetic analyses revealed that current Taiwan H5N2 viruses diverged into groups A and B and were distinct from the
earlier isolates from 2003 and 2008, raising the possibility of multiple introductions of the viruses into Taiwan or a single introduction of a virus that persisted undetected and continuously evolved in the field. In the connecting peptide of these viruses, the residue at position 3 of HA1 varies between E and K (with one instance of R) (Table 3). Changes to basic amino acids are commonly observed at the connecting peptide when a virus changes from low to high pathogenicity (7, 13); however, the reversion to an acidic or the original residue is rare in HPAI virus lineages. While a reversion from K to the original E could have occurred on the branch leading to group A, the unusual prevalence of the ancestral state supports repeated reintroduction of the viruses (Fig. 2A).

Outbreaks resulting in high mortality in chickens were reported from Taiwan in 2012 (21). In our study, a variable level of pathogenicity was found in the viruses even though the motifs in the connecting peptides contained multiple basic amino acids. The glycosylation site at the N terminus of the HA protein may interfere with the accessibility of the cleavage enzyme to some extent (10, 11, 41). The contemporary group B viruses, which include Ck/TW/1680/13, have a truncated NS1 protein, while group A viruses have a truncated PB1-F2. This may contribute to the varied pathogenicities, as indicated by the ICPI test. Further investigation of the related molecular basis for this is warranted.

Serological studies revealed that approximately 8.3% and 3.5% of chickens surveyed were seropositive to H5N2 and H6N1 viruses. It could be possible that this reflects prior exposure to circulating viruses. However, the 82.1% seropositive rate to a Hong Kong H9N2 virus, belonging to the Ck/Beijing/94 lineage (38), provided evidence of a broad and recent chicken vaccination program in Taiwan. To date, the Ck/Beijing/94-like H9N2 viruses are highly prevalent mainly in mainland China but have not been reported in Taiwan, nor were they detected in our surveillance. This further supports the possibility that, while apparently prohibited (21), vaccination with an H5N2 virus (potentially a bi- or

<table>
<thead>
<tr>
<th>Isolation date (mo/yr)</th>
<th>Strain⁵</th>
<th>HA1 position at:</th>
<th>HA2 position +1</th>
</tr>
</thead>
<tbody>
<tr>
<td>1994 Ck/Hidalgo/28159-232/94</td>
<td>P Q R E</td>
<td>T</td>
<td>R</td>
</tr>
<tr>
<td>2005 Ck/Ibaraki/1/05</td>
<td>P Q R E</td>
<td>T</td>
<td>R</td>
</tr>
<tr>
<td>12/2003 Ck/TW/1209/03</td>
<td>P Q R E</td>
<td>K</td>
<td>K</td>
</tr>
<tr>
<td>10/2008 Ck/TW/A703-1/08</td>
<td>P Q R K</td>
<td>K</td>
<td>R</td>
</tr>
<tr>
<td>1/2012 Ck/TW/A1997/12</td>
<td>P Q R K</td>
<td>K</td>
<td>R</td>
</tr>
<tr>
<td>12/2012 Ck/TW/683/12-like (n = 3)</td>
<td>P Q R E</td>
<td>K</td>
<td>R</td>
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<td>R</td>
</tr>
<tr>
<td>3/2013 Ck/TW/2948/13</td>
<td>P Q R E</td>
<td>K</td>
<td>K</td>
</tr>
<tr>
<td>Ck/TW/7350/13</td>
<td>P Q R K</td>
<td>K</td>
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<tr>
<td>5/2013 Ck/TW/4443/13</td>
<td>P Q R E</td>
<td>K</td>
<td>R</td>
</tr>
</tbody>
</table>

⁵ The glycosylation site at +1 (H3 numbering) is conserved in all viruses in this table.

Three viruses (Ck/TW/1680/13, Ck/TW/8994/13, and Ck/TW/8996/13) were selected for determination of pathogenicity in day-old chickens using the intracerebral pathogenicity index (ICPI).

### TABLE 4 Determination of the ICPI of the chicken H5N2 viruses prevalent in Taiwan

<table>
<thead>
<tr>
<th>Virus</th>
<th>Clinical sign</th>
<th>No. of chickens with specific sign by day postinoculation</th>
<th>ICPI²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ck/TW/1680/13</td>
<td>Normal</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Sick</td>
<td>4</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td>Dead</td>
<td>1</td>
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</tr>
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² ICPI, intracerebral pathogenicity index.
trivalent vaccine including H6N1 and/or H9N2 viruses) has occurred and that this insufficiently inactivated or inadequately attenuated live-virus vaccine is the source of the H5N2 viruses found in chickens in Taiwan.

Widespread vaccination of poultry is commonly used in several countries to provide protection against avian influenza (50). The quality of the vaccine is critical to the success of such programs, while control measures such as stamping out and localized temporary vaccination upon an outbreak could be an alternative (50, 51). In the case of H5N2 in Taiwan, the reemergence of a North American vaccine strain virus in Asia after approximately 10 years is a strong indication that a vaccine may be the source of this virus. The currently emerging H5N2 variants in Taiwan are suggestive of possible reintroductions from a vaccine source, and some are developing signs of evolving into a highly pathogenic strain. This example provides a strong warning against the use of vaccination in the absence of a present threat and the need for the highest quality of vaccine and strict control of vaccination practice by competent, accountable authorities. Thorough surveillance and a localized response, if needed, may be an effective way to prevent a situation such as the H5N2 influenza virus outbreaks in Taiwan from developing.

ACKNOWLEDGMENTS
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