Imprinting of Repeated Influenza A/H3 Exposures on Antibody Quantity and Antibody Quality: Implications on Seasonal Vaccine Strain Selection and Vaccine Performance

Martina Kosikova¹#, Lei Li²#, Peter Radvak¹, Zhiping Ye¹, Xiu-Feng Wan², Hang Xie¹

¹Laboratory of Pediatric and Respiratory Viral Diseases, Division of Viral Products, Office of Vaccines Research and Review, Center for Biologics Evaluation and Research, United States Food and Drug Administration (CBER/FDA), 10903 New Hampshire Ave, Silver Spring, MD 20993; ²Department of Basic Sciences, College of Veterinary Medicine, Mississippi State University, 240 Wise Center, Mississippi State, MS 39762

#M. Kosikova and L. Li contributed equally to this manuscript.

Corresponding author: H. Xie, CBER/FDA, 10903 New Hampshire Ave, Silver Spring, MD 20993 (Hang.Xie@fda.hhs.gov).

Summary: Influenza antibodies from hosts with pre-existing immunity showed broader cross-reactivity than those without pre-existing immunity. Repeated influenza exposures imprinted both antibody quantity and antibody quality. To improve vaccine effectiveness, vaccine strain selection needs to consider the pre-existing immunity in humans.

Running title: Influenza exposure on antibody quality

Published by Oxford University Press for the Infectious Diseases Society of America 2018. This work is written by (a) US Government employee(s) and is in the public domain in the US.
ABSTRACT

**Background.** Reduced seasonal influenza vaccine effectiveness (VE) was observed in individuals who received repeated annual vaccinations. Pre-existing influenza antibody levels were also found inversely correlated with post-vaccination titers. These reports suggest that pre-existing immunity may affect contemporary seasonal vaccine performance.

**Methods.** Influenza A/H3 specific cross-reactivity of post-vaccination sera from humans with or without pre-existing immunity was assessed by hemagglutination inhibition (HAI) assay. Ferret antisera induced by repeated H3 exposures were also subjected to HAI, antibody affinity and antibody avidity analyses.

**Results.** Human post-vaccination sera derived from subjects with or without pre-existing immunity showed different cross-reactivity against H3 variant viruses. Similarly, the breadth of cross-reactive ferret antibodies induced by repeated H3 exposures was also broadened. Antigenic differences between H3 viruses characterized by ferret antisera became smaller as the number of exposures increased. Although repeated H3 exposures induced “Original Antigenic Sin” phenomena in HAI titers against later exposed viruses, resultant ferret antibodies showed gradually enhanced avidity for different H3/HA. Increased antibody avidity was found inversely correlated with decreased antigenic differences among H3 viruses characterized.

**Conclusions.** Our results suggest that repeated H3 exposures imprinted not only antibody quantity but also antibody quality. The “naïve” ferret model currently used for vaccine strain selection does not recapitulate the complexity of human pre-existing
immunity. Vaccine strains identified hereby may not provide coverage sufficient for those who were frequently infected and/or vaccinated leading to reduced VE observed.

**Keywords:** Seasonal Influenza Vaccine; Vaccine Strain Selection; Antibody Cross-reactivity; Antibody Avidity; Repeated Influenza Exposure

**INTRODUCTION**

Annual vaccination is the main preventive strategy for control of seasonal influenza, the recommendation of which was first introduced in 1960 for older adults and immunocompromised individuals at high risks for severe influenza-like illness [1]. Since 2010, the annual vaccination policy has been expanded to include all healthy persons aged ≥ 6 months in the US [2], and a similar policy has also been adopted by many countries worldwide. Thus, a child born after 2010 would expect to be vaccinated ≥70 times during an average 75-year life expectancy. The annual vaccination policy is made largely due to the virus evolution resulting in frequent antigenic drift or shift, which in turn requires seasonal vaccine strains to be updated yearly to match with circulating viruses. Even in seasons without vaccine strain changes, concerns of vaccine-induced immunity waning also necessitate revaccination [2]. This allows individuals with frequent vaccinations to develop influenza antibody repertoires that are readily recalled on later exposure to antigenically similar or related viruses. However, recent epidemiology studies have reported that lower vaccine effectiveness (VE) was observed in individuals with repeated annual vaccination than those who were not vaccinated in previous season(s) [3-10]. A potential negative effect of prior vaccination was found more pronounced for H3N2-specific VE [3, 4, 6, 9]. These reports suggest that seasonal
vaccine-induced protection may be dampened in frequent vaccinees, prompting questions about the benefits of annual vaccination policy.

Annual vaccine strain evaluation and selection organized by the World Health Organization (WHO) is key to controlling seasonal vaccine performance. In this complex process, standard antisera raised in seronegative ferrets infected with representative influenza strains are used to antigenically characterize circulating viruses and play a decisive role in identifying final vaccine strains [11-13]. However, influenza-specific immunity in humans is greatly shaped by previous exposures including natural infections and annual vaccinations, the complexity of which cannot be recapitulated by seronegative ferrets infected with single influenza strain [12, 14-21]. Antigenic distances derived by seronegative ferret model have been found to correlate poorly with both H1- and H3-specific VE in humans [20, 22, 23]. This fundamental difference is one of the major factors responsible for the 2014/15 Northern Hemisphere vaccine strain mismatch and poor vaccine performance [12]. Even in humans, pre-existing immunity is also highly variable due to age and infection/vaccination histories [14-19, 24]. In vaccine trials or VE studies, however, the pre-existing background of recruited subjects is seldom explicitly determined or analyzed. This is also a problem for annual vaccine strain selection, in which anonymous post-vaccination sera provided by courtesy of international government/industry partnerships are used for human serology [12, 18].

In this study, we demonstrated that repeated H3 exposures imprinted both antibody quantity and antibody quality, thus significantly affecting virus antigenic characterization. To improved seasonal vaccine performance, we must take into account of human pre-existing immunity during vaccine strain selection.
METHODS

Viruses

All H3N2 viruses were propagated in 9-10 days old embryonated eggs, including (1) Clade 1 A/Uruguay/716/2007X175C (175C); (2) Clade 3C.1 A/Texas/50/2012 (TX/50); (3) Clade 3C.2a A/Hong Kong/4801/2014 (HK/4801), A/Singapore/KK934/2014 (SGPKK934), A/Fiji/2/2015 (Fiji/2), A/South Australia/09/2015 (SA/09), A/South Australia/21/2015 (SA/21), A/Victoria/503/2015 (Vic/503), A/Brisbane/47/2015 (Bris/47) and A/Brisbane/82/2015 (Bris/82); (4) Clade 3C.3 A/New York/39/2012 (NY/39); (5) Clade 3C.3a A/Switzerland/9715293/2013 (SWZ/13), A/North Carolina/13/2014 (NC/13) and A/Palau/6759/2014 (PA/14); (6) Clade 3C.3b A/Victoria/511/2015 (Vic/511).

Human Sera

Archived human sera were obtained from healthy subjects administered the 2015/16 egg-based inactivated, nonadjuvanted, trivalent or quadrivalent influenza vaccines and analyzed anonymously as part of a public health, non-research, regulatory activity in support of WHO annual influenza vaccine strain selection, which is exempt from human subjects review. Post-vaccination sera from children (8–30 months of age) and adults (28–74 years of age) showing high seroconversion (≥ 16-fold rise in post-vaccination HAI titers against the 2015/16 H3 vaccine prototype virus– SWZ/13) were selected to assess the HAI cross-reactivity against a panel of previous and recent H3N2 viruses.

Ferret Infection Experiments

Seronegative male ferrets (Triple F Farm) at 15-16 weeks old were anesthetized with ketamine/xylazine mixture followed by intranasal inoculation with H3N2 viruses
individually or sequentially at 2-week intervals: (1) HK/4801 only (HK); (2) SWZ/13 followed by HK/4801 (SWZ/HK); (3) TX/50 followed by SWZ/13 and then HK/4801 (TX/SWZ/HK); (4) 175C followed by TX/50, SWZ/13 and then HK/4801 (175C/TX/SWZ/HK). Blood was collected at 14 days post-infection right before each new inoculation. All the procedures were carried out in accordance with the protocol approved by the Institutional Animal Care and Use Committee of CBER/FDA.

**HA Inhibition (HAI) Assay**

The HAI assays were performed using 8 HA units/50 μL of stock viruses and 0.75% guinea pig erythrocytes in the presence of 20 nM oseltamivir for H3N2 viruses, or 0.5% turkey erythrocytes for H1N1 and type B viruses as described previously [12, 18]. Following treatment with receptor-destroying enzyme (RDE) (Denka-Seiken), pre-vaccination and post-vaccination sera were serially 2-fold diluted for HAI testing. HAI titers represent the reciprocal of the highest serum dilution that yielded a complete hemagglutination inhibition. A titer 5 was assigned if no inhibition was observed at the starting 1:10 serum dilution. HAI geometric mean titers (GMT) were calculated.

**Microneutralization (MN) Assay**

An ELISA-based MN assay was performed as previously described [25, 26]. RDE-treated sera were incubated with 100 50% tissue culture infectious dose at 37 ºC, 5% CO₂ for one hour, and then were added to Madin-Darby Canine Kidney (MDCK)-SIAT1 cells (Sigma). After overnight incubation, infected cells were detected using influenza A
nucleoprotein specific monoclonal antibodies (Millipore). MN titers represent the reciprocal of the highest serum dilution resulting in ≥50% neutralization.

**Ferret IgG Purification**

IgG antibodies in ferret antisera were purified using NAb Protein A Plus Spin kit (ThermoFisher). Total amount of purified ferret IgG was quantitated using reducing agent-compatible Pierce™ microplate BCA protein assay kit (ThermoFisher).

**Anti-HA ELISA and Avidity Assays**

Nunc MaxiSorp™ microtiter plates (ThermoFisher) were coated with 0.2 µg/ml of recombinant HA (Protein Sciences) in pH 9.5 bicarbonate/carbonate coating buffer at 4 °C overnight followed by 2 hours room temperature blocking in blocking buffer (pH 7.4 PBS containing 1% BSA and 5% Sucrose). Ferret antisera or purified antibodies were serially diluted in assay buffer (pH 7.4 PBS containing 0.1% BSA and 0.05% Tween 20) and incubated on blocked plates at 37 °C for 90 min. For avidity assays, the plates were washed and overlaid with 100 µl/well of 4 M Urea for 15 min [27]. After thorough washing, the plates were re-blocked in blocking buffer for another hour. Bound Ab was detected using peroxidase-conjugated goat anti-ferret IgG (abcam) followed by 1-Step™ Ultra TMB-ELISA substrate (ThermoFisher). Absorbance values were measured at 405 nm in a Victor V multilabel reader (PerkinElmer). Avidity index was calculated based on the area of the entire antibody titration curve as previously reported [28].
Antigenic Cartography and Correlation Coefficient Analysis

The 2-dimensional antigenic maps with multidimensional scaling were constructed based on human or ferret HAI titers using AntigenMap (http://sysbio.cvm.msstate.edu/AntigenMap) with each horizontal or vertical gridline representing one antigenic unit distance corresponding to a 2-fold difference in HAI titers [12, 29, 30]. The correlation coefficient for any two sets of antigenic distances derived from antigenic cartography was also determined with 1 indicating a perfect positive correlation and −1 denoting a perfect negative correlation, respectively [18].

Statistical Analysis

HA-specific ferret IgG binding affinity was analyzed using nonlinear regression curve fit (Prism 6.02, GraphPad). $P < 0.05$ determined by two-way ANOVA was considered statistically significant.

RESULTS

Pre-existing Immunity Affected Human H3-specific HAI Cross-reactivity

Because of lack of infection/vaccination records, selected pediatric or adult post-vaccination sera were grouped as previously reported [12, 14, 15, 18]: (1) undetectable H3-specific pre-existing immunity (pre-vaccination HAI titer of $<40$ against SWZ/13), and (2) detectable H3-specific pre-existing immunity (pre-vaccination HAI titer of $\geq 40$ against SWZ/13). Compared to the pediatric or adult group with undetectable pre-existing immunity, the corresponding age group with detectable pre-existing immunity responded more evenly to all H3 viruses tested (Figure 1A-1B). When these same post-
vaccination HAI titers were visualized using antigenic cartography (Figure 1C-1F), interestingly the groups with detectable pre-existing immunity, regardless of ages, had difficulties in distinguishing different H3 clades as compared to the groups with undetectable pre-existing immunity (Figure 1C vs 1D and 1E vs 1F). For instance, Clade 3C.2a (green) and Clade 3C.3a (red) viruses were well separated in the antigenic map derived from pediatric post-vaccination sera with undetectable pre-existing immunity, indicative of distinct antigenicity (Figure 1C). However, in the antigenic map derived from pediatric post-vaccination sera with detectable pre-existing immunity, these two clades tended to cluster together and were not distinctly separated (Figure 1D). Similar phenomena were observed in the maps derived from adult post-vaccination sera with and without detectable pre-existing immunity (Figure 1E vs 1F). In particular, Clade 3C.2a and Clade 3C.3a viruses became completely indistinguishable in the adult map with detectable pre-existing immunity (Figure 1F). The smaller antigenic distances in pediatric or adult map with detectable pre-existing immunity indicated smaller antigenic differences among H3 viruses characterized (1.2763 vs 1.4629 and 0.8339 vs 1.0340 in pediatric and adult with undetectable pre-existing immunity respectively; Figure 1G). Correlation coefficient analysis also showed that the maps with detectable pre-existing immunity correlated poorly with those with undetectable pre-existing immunity (Figure 1G). These results indicated that the post-vaccination sera from the subjects with detectable pre-existing immunity had different cross-reactivity toward H3N2 variants from those with undetectable pre-existing immunity.
Prior H3 Exposures Affected both Quantity and Quality of Ferret Antibodies

We then investigated how prior H3 exposures may affect antibody development in a ferret model. Seronegative ferrets were either exposed to a single HK/4801 (clade 3C.2a) infection or sequentially infected with antigenically drifted 175C (clade 1), TX/50 (clade 3C.1) or SWZ/13 (clade 3C.3a) followed by HK/4801. As expected, ferret antisera raised from single HK/4801 infection showed much higher HAI titers against the homologous virus (GMT= 557) than drifted H3N2 variants (Figure 2A and Supplementary Table S1). In contrast, ferret antisera derived from sequential H3N2 infections showed better HAI responses toward first encountered H3N2 virus than later exposed HK/4801 (Figure 2A-2D). For instance, ferret 175C/TX/SWZ/HK antisera exhibited higher HAI response toward the first exposed 175C virus (GMT= 226), but much lower titers against later infected HK/4801 (GMT= 80) (Figure 2D), indicating a typical “original antigenic sin (OAS)” [31]. Yet the OAS phenomena were less pronounced in ferret MN titers against the same set of viruses (Supplementary Figure S1).

Despite lower HAI titers against later exposed viruses, ferret IgG purified from antisera with repeated H3 exposures in general exhibited enhanced binding affinity for both early and late exposed HA antigens as compared to ferret IgG elicited by single HK/4801 infection (Figure 3A-3D). Using urea as the chaotrope, we then measured antibody avidity which detects the total strength of multivalent interactions between antibody and antigen, instead of a specific interaction between one antigenic epitope and one antibody binding site detected by antibody affinity. As shown in Figure 4, antibodies induced by single HK/4801 infection had the lowest avidity index not only against the
homologous HA but also against drifted H3 HA. In contrast, repeated H3 infections gradually enhanced the avidity of the resulting ferret antibodies, e.g., ferret 175C/TX/SWZ/HK antisera had the highest avidity index against all four H3 HA antigens tested (Figure 4A-4D). Apparently, more prior H3 exposures, higher H3-specific antibody avidity.

These results suggest that prior H3 exposures not only affect the quantity but also the quality of ferret HA-specific antibodies, and the impact seems to depend upon the number of previous exposures.

**Repeated Prior H3 Exposures Expanded Ferret HAI Cross-reactivity**

As expected, ferret antisera raised from single infection mainly cross-reacted with homologous viruses (Figure 5A and Supplementary Figure S2). When ferrets were sequentially infected with SWZ/13 followed by HK/4801, the resulting SWZ/HK antisera had HAI cross-reactivity extend to Clade 3C.1, 3C.3, 3C.3a and 3C.3b viruses by showing >50% rise in HK/4801-specific HAI GMT (Figure 5B). Similarly, TX/SWZ/HK antisera induced by sequential TX/50, SWZ/13 and HK/4801 exposures broadened HAI cross-reactivity to Clade 3C.1, 3C.3, 3C.3a and 3C.3b viruses, including the two Clade 3C.2a viruses – SGP/KK943 and VIC/503 (Figure 5C) which HK and SWZ/HK antisera showed ≥50% reduction in HAI GMT (Figure 5A and 5B). When ferrets had been sequentially exposed to 175C, TX/50 and SWZ/13 followed by HK/4801, the resulting 175C/TX/SWZ/HK antisera expanded the HAI cross-reactivity to all six H3N2 clades tested including Clade 1 virus (Figure 5D).
When these same ferret HAI titers were plotted using antigenic cartography, it became obvious that different H3N2 clades were less distinguishable by antisera raised from ferrets with increased exposure histories (Figure 5E-5H). For example, Clade 3C.2a (green) and Clade 3C.3a (red) were well separated in the map derived from ferret antisera raised after single HK/4801 infection (Figure 5E). However, the antigenic differences between Clade 3C.2a and Clade 3C.3a viruses became smaller in the map derived from the TX/SWZ/HK antisera (Figure 5G). Eventually Clade 3C.2a and 3C.3a groups clustered together and became antigenically indistinguishable as determined by the 175C/TX/SWZ/HK antisera (Figure 5H). The average antigenic distances in individual ferret maps decreased when the number of previous H3N2 infections increased (Figure 5I).

The correlation analysis showed that the antigenic maps derived by ferret antisera with previous H3N2 infections correlated poorly with the one derived by ferret antisera with single HK/4801 infection: more prior H3N2 exposures, smaller correlation coefficients (Figure 5I).

**Ferret Antibody Avidities Inversely Correlated with Antigenic Distances of H3N2 Viruses Characterized**

We then correlated the avidity indexes of individual ferret antisera in Figure 4 with the average antigenic distance derived by corresponding ferret antisera in Figure 5I. It showed that reduced average antigenic distances inversely correlated with increased antibody avidities in ferrets with multiple prior H3N2 exposures ($R^2 = 0.5765$, $P = 0.0006$): higher antibody avidity and shorter antigenic distances (Figure 6). These
results suggest that repeated H3N2 exposures enhance antibody avidity thus affecting virus antigenic characterization.

**DISCUSSION**

Current vaccine strain selection uses seronegative ferret model without influenza-specific pre-existing immunity to detect epidemic viruses that are antigenically different from vaccine strains [11-13]. However, accumulated evidence indicates that early life exposure to influenza can leave an imprint on human antibody repertoires, and resulted residual protection may last lifetime [15, 16, 21, 31-35]. Intensified global connectivity helps to spread antigenically drifted influenza strains [36]. Repeated annual vaccination also contributes to widespread influenza pre-existing immunity in humans. Virtually all humans have been exposed – asymptomatically or symptomatically – to influenza, and there exist no such “influenza-naïve” persons except newborns. As shown in this study, the H3 viruses representing different genetic clades were “seen” as antigenically related by human post-vaccination sera with detectable H3 pre-existing immunity, rather than antigenically distinguishable by those with undetectable H3 pre-existing immunity. This difference is not due to poor vaccination response since all human post-vaccination sera used in this studyhad ≥ 16-fold seroconversion after seasonal vaccination. Rather, our results confirm that the background of previous exposures – whether by natural infections or annual vaccinations – can significantly change human post-vaccination antibody cross-reactivity [18].

Similarly, pre-existing immunity also can influence virus antigenic characterizations in ferrets. Clade 3C.2a and Clade 3C.3a viruses were antigenically well separated by
ferret antisera stimulated by single H3 infection; whereas their antigenic difference diminished as determined by ferret antisera with increased prior H3 exposures. This is because repeated H3 infections extended the cross-reactivity of resultant ferret antibodies from Clade 1 to Clade 3C.X, in contrast to strain-specific antibodies elicited by single H3 infection. Consequently, Clade 3C.2a and Clade 3C.3a viruses were no longer “seen” antigenically distinct by ferret antisera with repeated H3 exposures.

The broadened breath of ferret antibody cross-reactivity apparently occurred at the expense of absolute HAI titers against more recent viruses – an OAS phenomenon that has traditionally been considered detrimental to contemporary vaccine performance [31-34]. Human pre-existing influenza-specific antibody levels have been found to inversely correlate with post-vaccination titers [35-37]. In individuals with high pre-vaccination antibody titers, vaccination is found to primarily boost pre-existing influenza-specific repertoires rather than to induce de novo antibody clonotypes [37]. Many pre-existing antibodies are cross-reactive and can help to restrict subsequent infections caused by antigenic variants of influenza [37, 38]. Correspondingly, a suppressed de novo antibody response specific for later exposed virus may occur. In this study, we observed that repeated H3 infections induced more pronounced OAS phenomena in ferret HAI titers than in ferret MN titers. This could be because some pre-existing antibodies with broad cross-reactivity have no HAI activity [37].

Our current study also suggests that repeated H3 exposures resulted in improved quality of recall antibodies, which might be partially via OAS induction. Antibodies from ferrets primed with 175C may target HA antigenic sites that are also conserved by later H3 strains. When the same ferrets were repeatedly infected with evolved but still
antigenically related TX/50, SWZ/13 and HK/4801, pre-existing cross-reactive antibodies responded to limit viral infections leading to reduced HAI titers against recall viruses. This may also give opportunities to B cells specific for other antigenic sites to expand [39]. Hence, the resulting pool of ferret antibodies had diversified specificities against a wide range of H3 viruses. Since individual viruses have different antigenic sites to play a dominant role in immune system [40], repeated exposures to different H3 viruses may not necessarily increase the binding affinity of resultant antibodies for a specific antigenic determinant. However, their avidity – the multivalent bind capacity for all antigenic sites was enhanced along with increased virus encounters, thus resulting in less distinguishable antigenic differences between H3 variants characterized.

Taken together, our study shows that repeated influenza exposures imprinted not only antibody quantity but also antibody quality. The “naïve” ferret model currently used for vaccine strain selection does not reflect the complexity of human exposure history. Whether viruses appear antigenically identical or drift to “naïve” ferrets may not hold true for individuals with frequent exposures. This could lead to miss identifying a strain that indeed is antigenically drift to repeatedly vaccinated individuals [20]. Vaccine strains selected hereby may not provide coverage sufficient for those with frequent vaccinations/infections resulting in compromised VE observed. To improve seasonal vaccine performance, we should take the widespread human pre-existing immunity into consideration during vaccine strain selection, and select vaccine strains that are optimal for populations with different immune backgrounds.
**Author contributions.** H.X. conceived the ideas and designed the study. M. K., P. R. and H. X. conducted the ferret infection experiments. M. K., Z. Y. and H. X. performed the HAI assays. M. K. conducted antibody quality experiments. L. L. and X.-F. W. generated the antigenic maps and did correlation coefficient analyses. H. X. analyzed the data and performed the statistical analyses. H. X. and M. K. wrote the paper.

**Acknowledgments.** The authors sincerely appreciate Dr. Xiyan Xu of Centers for Diseases Control and Prevention for providing influenza viruses.

**Disclaimer.** The findings and conclusions in this article have not been formally disseminated by US Food and Drug Administration and should not be construed to represent any Agency determination or policy.

**Financial support.** This project was supported by the intramural research fund of CBER, US Food and Drug Administration. X.-F. W. and L. L. were supported by the National Institutes of Health (grant number R01AI116744 to X.-F. W.).

**Potential conflicts of interest.** The authors declare no competing financial interests.
References

15. Xie H, Li X, Gao J, et al. Revisiting the 1976 "swine flu" vaccine clinical trials: cross-reactive hemagglutinin and neuraminidase antibodies and their role in...


FIGURE LEGENDS

Figure 1. Different antigenic patterns of human post-vaccination sera with or without detectable H3-specific pre-existing immunity. A-B) Normalized hemagglutinin inhibition (HAI) geometric mean titers (GMT) in pediatric (A) or adult (B) subjects with or without detectable H3-specific pre-existing immunity. Post-vaccination HAI GMT were normalized to vaccine strain A/Hong Kong/4801/2014 (HK/4801) specific GMT in the same subpopulation without detectable pre-existing immunity. C-F) Antigenic maps constructed using post-vaccination HAI titers from the same pediatric (C & D) or adult (E & F) subjects with or without detectable H3-specific pre-existing immunity. Each gridline (horizontal and vertical) in the antigenic maps represents one antigenic unit distance corresponding to a 2-fold difference in HAI titers. 1) Antigenic distances of H3 viruses determined in each antigenic map and correlation analysis within the same population. Correlation coefficients of 1, −1, and 0 represent a perfect positive correlation, a perfect negative correlation, and no correlation, respectively. Both pediatric and adult participants were vaccinated with the 2015/16 egg-based inactivated, nonadjuvanted, trivalent or quadrivalent vaccines. Egg-grown H3 viruses in the testing panel included Clade 1 – A/Uruguay/716/2007X175C (175C); Clade 3C.1 – A/Texas/50/2012 (TX/50); Clade 3C.2a –HK/4801, A/Singapore/KK934/2014 (SGPKK934), A/Fiji/2/2015 (Fiji/2), A/South Australia/09/2015 (SA/09), A/South Australia/21/2015 (SA/21), A/Victoria/503/2015 (Vic/503), A/Brisbane/47/2015 (Bris/47) and A/Brisbane/82/2015 (Bris/82); Clade 3C.3 – A/New York/39/2012 (NY/39); Clade 3C.3a – A/Switzerland/9715293/2013 (SWZ/13), A/North Carolina/13/2014 (NC/13) and A/Palau/6759/2014 (PA/14); Clade 3C.3b – A/Victoria/511/2015 (Vic/511).
Figure 2. H3 virus-specific hemagglutinin inhibition (HAI) responses in ferrets with different exposure histories. Sera from ferrets infected with (1) A/Hong Kong/4801/2014 (HK/4801) only (HK), or (2) A/Switzerland/9715293/2013 (SWZ/13) followed by HK/4801 (SWZ/HK), or (3) A/Texas/50/2012 (TX/50) followed by SWZ/13 and then HK/4801 (TX/SWZ/HK), or (4) A/Uruguay/716/2007X175C (175C) followed by TX/50, SWZ/13 and then HK/4801 (175C/TX/SWZ/HK) were assessed for virus-specific HAI titers. Individual ferret titers with geometric mean titers (GMT) (bar graphs) are shown.

Figure 3. H3-specific IgG affinity in ferrets with different exposure histories. Sera were collected from ferrets infected with (1) HK, or (2) SWZ/HK, or (3) TX/SWZ/HK, or (4) 175C/TX/SWZ/HK. Purified ferret IgG was assessed and analyzed for HA-specific binding affinity using nonlinear regression (curve fit).

Figure 4. H3-specific IgG avidity in ferrets with different exposure histories. Sera were collected from ferrets infected with (1) HK, or (2) SWZ/HK, or (3) TX/SWZ/HK, or (4) 175C/TX/SWZ/HK. HA-specific ferret IgG avidity was determined in the presence of 4M Urea.

Figure 5. H3 antigenic patterns determined by ferret antisera with different exposure histories. A-D) Normalized H3-specific ferret cross-reactive hemagglutinin inhibition (HAI) geometric mean titers (GMT). Egg-grown H3 viruses in the testing panel included Clade 1 – A/Uruguay/716/2007X175C (175C); Clade 3C.1 – A/Texas/50/2012
Clade 3C.2a – A/Hong Kong/4801/2014 (HK/4801), A/Singapore/KK934/2014 (SGPKK934), A/Fiji/2/2015 (Fiji/2), A/South Australia/09/2015 (SA/09), A/South Australia/21/2015 (SA/21), A/Victoria/503/2015 (Vic/503), A/Brisbane/47/2015 (Bris/47) and A/Brisbane/82/2015 (Bris/82); Clade 3C.3 – A/New York/39/2012 (NY/39); Clade 3C.3a – A/Switzerland/9715293/2013 (SWZ/13), A/North Carolina/13/2014 (NC/13) and A/Palau/6759/2014 (PA/14); Clade 3C.3b – A/Victoria/511/2015 (Vic/511). Sera were collected from ferrets infected with (1) HK, or (2) SWZ/HK, or (3) TX/SWZ/HK, or (4) 175C/TX/SWZ/HK. Ferret HAI GMT specific for individual H3 viruses are normalized to HK/4801-specific GMT with the red dotted lines indicating 50% response. E-H)

Antigenic maps constructed using H3-specific HAI titers of ferret antisera. Each gridline (horizontal and vertical) in the antigenic maps represents one antigenic unit distance corresponding to a 2-fold difference in HAI titers. I) Antigenic distances of H3 viruses determined in each ferret antigenic map and correlation analysis relative to ferret HK antisera-derived antigenic map. Correlation coefficients of 1, −1, and 0 represent a perfect positive correlation, a perfect negative correlation, and no correlation, respectively.

**Figure 6. Correlation between IgG avidities and antigenic distances determined by ferret antisera with different exposure histories.** H3-specific IgG avidities of ferret antisera determined in Figure 4 were correlated with average antigenic distances determined by corresponding ferret antisera in Figure 5I using linear regression.
Figure 1.
Figure 2.

A. Against HK/4801 virus

B. Against SWZ/13 virus

C. Against TX/50 virus

D. Against 175C virus

Virus-specific HAI GMT (Log2)

Ferret antisera
Figure 3.
Figure 4.

A

Against HK/4801 HA

Ferret antisera

B

Against SWZ/13 HA

Ferret antisera

C

Against TX/50 HA

Ferret antisera

D

Against 175C HA

Ferret antisera
Figure 5.

Antigenic distances and correlation coefficients

<table>
<thead>
<tr>
<th>Antigenic map</th>
<th>Exposure history</th>
<th>Antigenic distance</th>
<th>Correlation coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>HK only (n=5)</td>
<td>3.6633</td>
<td>0.2847</td>
</tr>
<tr>
<td>B</td>
<td>SWZ/HK (n=4)</td>
<td>3.5227</td>
<td>0</td>
</tr>
<tr>
<td>C</td>
<td>TX/SWZ/HK (n=4)</td>
<td>3.2037</td>
<td>0</td>
</tr>
<tr>
<td>D</td>
<td>175C/TX/SWZ/HK (n=4)</td>
<td>2.0758</td>
<td>0</td>
</tr>
</tbody>
</table>
Figure 6.

$R^2 = 0.5765$

$p = 0.0006$

- Green circle: Ferret HK antisera
- Red square: Ferret SWZ/HK antisera
- Blue diamond: Ferret TX/SWZ/HK antisera
- Yellow triangle: Ferret 175CTX/SWZ/HK antisera

IgG avidity index vs. Average antigenic distance